The *Elov4* Spinocerebellar Ataxia-34 Mutation 736T>G (p.W246G) Impairs Retinal Function in the Absence of Photoreceptor Degeneration

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Abstract

Elongation of very long chain fatty acids-4 (*ELOVL4*) is essential for synthesis of very long chain polyunsaturated and saturated fatty acids (VLC-PUFA and VLC-SFA, respectively) of chain length greater than 26 carbons. Mutations in the *ELOVL4* gene cause several distinct neurodegenerative diseases including Stargardt-like macular dystrophy (STGD3), spinocerebellar ataxia 34 (SCA34), and a neuro-ichthyotic syndrome with severe seizures and spasticity, as well as erythrokeratotic variabilis (EKV), a skin disorder. However, the relationship between *ELOVL4* mutations, its VLC-PUFA and VLC-SFA products, and specific neurological symptoms remains unclear. We generated a knock-in rat line (SCA34-KI) that expresses the 736T>G (p.W246G) form of *ELOVL4* that causes human SCA34. Lipids were analyzed by gas chromatography and mass spectrometry. Retinal function was assessed using electroretinography. Retinal integrity was assessed by histology, optical coherence tomography, and immunolabeling. Analysis of retina and skin lipids showed that the W246G mutation selectively impaired synthesis of VLC-SFA, but not VLC-PUFA. Homozygous SCA34-KI rats showed reduced ERG a- and b-wave amplitudes by 90 days of age, particularly for scotopic responses. Anatomical analyses revealed no indication of neurodegeneration in heterozygote or homozygote SCA34-KI rats out to 6–7 months of age. These studies reveal a previously unrecognized role for VLC-SFA in regulating retinal function, particularly transmission from photoreceptors to the inner retina, in the absence of neurodegeneration. Furthermore, these findings suggest that the tissue specificity and symptoms associated with disease-causing *ELOVL4* mutations likely arise from selective differences in the ability of the mutant *ELOVL4* enzymes to support synthesis of VLC-PUFA and/or VLC-SFA.

Keywords Very long chain fatty acids · Neurodegeneration · Stargardt-like macular dystrophy · Retina · Cerebellum

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Introduction

Elongation of very long chain fatty acids-4 (ELOVL4) catalyzes the first, rate-limiting step in the synthesis of very long chain polyunsaturated and saturated fatty acids with fatty acid chain lengths of 28 carbons or more [VLC-PUFA and VLC-SFA, respectively [1–3]]. ELOVL4 is the only known fatty acid elongase that performs this function [1, 2]. Expression of ELOVL4 is limited to a small number of organs including the retina, brain, skin, testes, and Meibomian gland [4–6], which show tissue-specific VLC-PUFA and VLC-SFA profiles. Our meticulous recent lipidomic analysis of mouse [7], rat, and baboon [3] brains using modern analytical tools found VLC-SFA (28:0 and 30:0). These VLC-SFA were incorporated into complex sphingolipids and were enriched in synaptic vesicles prepared from baboon hippocampus, where they contribute to the regulation of presynaptic neurotransmitter release [3, 8]. Previous studies have reported phosphatidylcholine containing VLC-PUFA in early postnatal rat brain and in the brains of human patients with Zellweger’s disease, a peroxisomal disorder that disrupts lipid metabolism [9, 10], but no VLC-PUFA have been reported in the healthy mature brain. VLC-SFA are also found in the Meibomian glands and skin, where they are incorporated into ω-O-acylceramides and form the water barrier of the tear film and skin [5, 11–14]. VLC-PUFA are present in the outer segments of rod and cone photoreceptors in the retina esterified to the sn-1 position of phosphatidylcholine [15–18]. VLC-PUFA in testes [19–21] are present in an amide linkage to sphingolipids [22–24].

ELOVL4 products have emerged as important novel regulators of synaptic function and neuronal survival in the CNS [23]. Several different mutations in the ELOVL4 gene cause neurological diseases that vary according to the mutation and its inheritance pattern [22, 25–36]. Heterozygous inheritance of four different ELOVL4 mutations causes autosomal dominant spinocerebellar ataxia-34 (SCA34), a late-onset neurodegenerative disease of the cerebellum that results in a characteristic gait ataxia that may be accompanied by dysarthria and eye movement abnormalities, with or without erythrokeratitisis variabilis (EKV), a disorder of the skin [28–30, 35]. Patients with SCA34 have no reported clinical retinal phenotype [28–30, 35]. However, a recent report identified an American family with a novel ELOVL4 mutation (c.512T>C,p.I171T) that causes spinocerebellar ataxia with some family members also showing retinitis pigmentosa [37].

Heterozygous inheritance of six different ELOVL4 mutations causes autosomal dominant Stargardt-like macular dystrophy (STGD3), which typically presents with juvenile onset [26, 27, 31–33, 36]. Patients with STGD3 have no other reported central nervous system or skin phenotypes [26, 27, 31–33, 36]. Homozygous inheritance of three other recessive ELOVL4 mutations causes a devastating neuro-ichthyotic syndrome characterized by seizures, intellectual disability, spasticity, ichthyosis, and premature death [25, 34]. Homozygous inheritance of the five-base-pair STGD3 mutation in mice is neonatal lethal due to dehydration caused by the absence of ω-O-acyl-certerides, a product of ELOVL4 that provides the permeability barrier [11–14]. Conditional expression of ELOVL4 in skin rescued the neonatal lethality, but the mice developed a severe seizure syndrome at postnatal day 18 similar in many respects to the recessive human ELOVL4 neuro-ichthyotic syndrome [3]. No human patients with homozygous inheritance of STGD3 or SCA34 alleles have been reported.

Spinocerebellar ataxias comprise a large constellation of autosomal dominant hereditary disorders that are characterized by progressive discoordination of gait (ataxia) and cerebellar atrophy. Spinocerebellar ataxias arise from defects in a large number of different genes, which produce distinct types of ataxia with characteristic times of onset and rates of disease progression [38, 39]. In addition to the characteristic ataxia, SCA patients also may show additional symptoms including dysarthria (difficulty speaking), apraxia (loss of fine motor control), eye movement deficits, and abnormal spinal reflexes, depending on the specific form of the disease. Currently, there are no treatments for any form of the disease.

How the different mutations in ELOVL4 affect levels of VLC-SFA and VLC-PUFA that cause distinct tissue-specific pathologies remains unclear. A key gap in our understanding is whether these disease-causing ELOVL4 mutations also compromise the function of additional tissues. To better understand the role of ELOVL4 and its VLC-fatty acid products in the retina, we generated a novel knock-in rat model expressing the 736T>G (p.W246G) form of ELOVL4 that causes human SCA34 [35] and examined lipid composition, retinal function, and photoreceptor degeneration. Our studies reveal a previously unrecognized role for the VLC-fatty acid products of ELOVL4 in synaptic transmission from photoreceptors to the inner retina in the absence of photoreceptor degeneration. This deficit is most likely attributable to deficiency of VLC-SFA rather than VLC-PUFA.

Methods

Animals

Generation and Characterization of c.736T>G, p.W246G Knock-In Rats Using CRISPR/Cas9

The CRISPR/Cas9 genetic editing approach was used to generate a heterozygous Long-Evans rat model of human SCA34 described by Ozaki et al. [35]. To minimize potential off-target effects, human Cas9 D10A (hCas9-D10A) nickase was used to knock-in the c.736T>G mutation by targeting exon 6 of the rat Elovl4 (Ensembl: ENSRNOG00000009773) located on
rat chromosome 8 (Fig. 1a). The hCas9 mRNA, sgRNA (CCTTCCCCAAGTGGATGCAC), and single-strand oligonucleotide donor (ssODN: TTCCATGACCAT TGGGCAACACACACTGGTCTCTCTCCTACCCGACTGCC CTTCCCAAGGGGATGCACCTGGGCTCCTGATGCG CTAAGCCATAGCGCTCTTCCCTCTTCTCTCAAAGTC) were co-injected into Long-Evans rat zygotes at the laboratories of Cyagen Bioscience Inc. (Santa Clara, CA). From the injected embryos transplanted into foster mothers, three heterozygous F0 rats carried the c.736T>G, p.W246G mutant SCA34 gene. The rats were genotyped using rat Elolv4-F: 5'-GCCCTTCAGCGAGTATTCAAATGTC-3 and Elolv4-R: 5'-GGTCACGTTGGGATTGCTTTGGTCTGGG-3' PCR primers that generate a 900-base-pair (bp) product, which upon restriction digestion with StyI (New England Biolabs, Ipswich, MA), generates a 450-base-pair product that allows identification of heterozygous and homozygous rats (Fig. 1b) due to introduction of the StyI restriction site from the c.736T>G mutation. The wild-type (WT) 900-base-pair band is resistant to the Sty I digestion (Fig. 1b). We bred the F0 rats with WT Long-Evans rats over several generations to breed out any potential off-target effects and to establish the heterozygous SCA34 Long-Evans rat model. We also successfully generated homozygous SCA34 rats (MUT) by breeding heterozygous rats from the different F0 lines. Knock-in of the SCA34 mutation in the rats was further verified by Sanger sequencing at Oklahoma Medical Research Foundation, Oklahoma City, OK (Supplemental Fig. 1) and by whole-genome DNA sequencing (Novogene Corporation, Inc., Sacramento, CA) (Supplemental Fig. 2).

Subsequent studies were performed using WT, heterozygous (HET), and homozygous (MUT) SCA34-knock-in Long-Evans rats (SCA34-KI) carrying the c.736T>G (p.W246G) mutation in ELOVL4 that causes age-related SCA34 in humans [35]. The rats were maintained in a pathogen-free barrier facility on a 12-h light:12-h dark daily light cycle (~ 25–40 lx at cage level). Food and water were

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**Fig. 1** Generation and characterization of c.736T>G, p.W246G knock-in Long-Evans rat using the CRISPR/Cas9 system. **a** Schematic depiction of targeting strategy. The genomic region of the rat Elolv4 locus is diagrammed (the gene is oriented from left to right; total size is 28.21 kb). Solid bars represent open reading frame (exons). Open bars represent untranslated regions. The sgRNA cut site and the single-strand oligonucleotide donor sequencing with homology arms and the mutation site is shown by the red arrow. **b** Genotyping of WT, HET, and MUT SCA34-KI rats by StyI restriction analysis. Seven hundred four base–pair PCR products were generated from WT, HET, and MUT rats using PCR primers. The amplicons were run undigested or purified and digested with StyI restriction enzyme. Left panel: undigested amplicons ran at 704 bp for WT, HET, and MUT rats, as expected. Right panel: digestion with StyI restriction enzyme showed the WT PCR product is resistant to StyI digestion. In contrast, HET rats show two bands, one corresponding to the WT PCR product and a smaller, digested fragment arising from the mutant PCR product containing the StyI digestion site. MUT rats show only a single band corresponding to the digested, mutant PCR product with no WT PCR product, as expected.
available at all times. All animal procedures were approved by the University of Oklahoma Health Sciences Center Institutional Animal Care and Use Committee and conformed to the National Institutes of Health Guide for the Care and Use of Laboratory Animals, the Association for Research in Vision and Ophthalmology Resolution on the Use of Animals in Research, and US Public Health Service guidelines.

**Lipid Analysis**

Skin VLC-SFA were quantified using gas chromatography-mass spectrometry (GC-MS) as previously described [1]. Briefly, 50 mg of tissue was homogenized in methanol and the lipids were extracted [40] and converted to fatty acid methyl esters (FAME) as described previously [41]. The GC-MS was operated in the electron impact (EI) single-ion monitoring (SIM) mode. The 26:0, 28:0, and 30:0 response values were obtained using the m/z ratios 74.1 and 87.1 as well as 410.4, 438.4, and 466.5, respectively. Sample concentrations were determined by comparison to external standards, using 25:0 and 27:0 as internal standards.

Retinal phospholipids were measured using electrospray ionization-mass spectrometry (ESI-MS-MS) as previously described [8, 42]. Briefly, tissue was homogenized in 40% aqueous methanol and diluted 1:40 with 2-propanol/methanol/chloroform (4:2:1) containing 20 mM ammonium formate and 1.0 μM phosphatidylcholine (PC, 14:0/14:0), 1.0 μM phosphatidylethanolamine (PE, 14:0/14:0), and 0.33 μM phosphatidylserine (PS, 14:0/14:0) as internal standards. Samples were introduced into a TSQ Ultra mass spectrometer using an Advion NanoMate operating in infusion mode. PC lipids were measured using precursor ion scanning of m/z 184, PE lipids (including plasmalogens) were measured using neutral loss scanning of m/z 141, and PS lipids were measured using neutral loss scanning of m/z 185. All species detected for each group were represented as a relative percentage of the sum based on their response values.

**Electroretinography**

Flash ERGs were recorded using a Diagnosys Espion E2 ERG system (Diagnosys LLC, Lowell, MA). The rats were maintained overnight in total darkness and prepared for ERG recording under dim red light. The rats were anesthetized with an intraperitoneal injection of ketamine (100 mg/kg body weight) and xylazine (5 mg/kg body weight). A drop of 0.5% (v/v) proparacaine HCl was applied to the cornea for local anesthesia. A drop of 10% (v/v) phenylephrine was applied to the cornea to dilate the pupil. The rats were kept on a heating pad at 37 °C during recording. A gold recording electrode was placed on the cornea, with a reference electrode positioned in the mouth and a ground electrode in the tail, and the rats were placed inside a Ganzfeld illuminating sphere. Responses were differentially amplified, averaged, and stored. To assess rod photoreceptor function (scotopic ERG), five strobe flash stimuli were presented at flash intensities of 0.004, 5.635, 11.243, 22.433, and 178.92 cd·s/m². Scotopic a-wave amplitude was measured from the pre-stimulus baseline to the negative trough of the a-wave. The scotopic b-wave amplitude was measured from the trough of the a-wave to the positive peak of the b-wave. To evaluate cone function (photopic ERG), a strobe flash stimulus (3.3 log cd s/m²) was presented to dilated, light-adapted (5 min at 1.7 log cd s/m²) rats. Photopic, short-wavelength cone, middle-wavelength cone, and flicker (3, 10, 20, and 30) Hz ERG responses were recorded, and the amplitude of the cone b-wave was measured from the trough of the a-wave to the peak of the b-wave.

**Optical Coherence Tomography**

Optical coherence tomography (OCT) was used to acquire images of retinal layering in vivo. OCT analysis was performed using a Bioptigen SDOIS system (Leica Microsystems, Buffalo Grove, IL). Rats were anesthetized by intraperitoneal injection of ketamine (100 mg/kg body weight) and xylazine (5 mg/kg body weight), and a drop of 10% (v/v) phenylephrine was applied to the cornea to dilate the pupil. The animal was placed in the animal mount, the eyes were well hydrated with Genteal eye drops, the ocular lens was positioned, and the image was viewed in real time. The optic nerve head was centered on the monitor, and an imaging scan was taken. Once the scan was completed and saved, the process was repeated for the second eye. The thickness of the retinal layers was then measured from the OCT images.

**Immunolabeling and Histology**

For immunolabeling experiments, rats were anesthetized using ketamine (100 mg/kg body weight) and xylazine (5 mg/kg body weight) delivered by intraperitoneal injection. Once anesthetized, the rat was secured on a foam block with pins and the thorax was cut through the rib cage to expose the heart. A cut was made to the right atrium and a 22-gauge needle was inserted into the left ventricle, and 0.1 M phosphate-buffered saline (PBS, pH 7.2) was pumped into the body for two minutes to remove all blood. Following exsanguination, 200 ml of 4% paraformaldehyde in 0.1 M PBS was pumped into the body over a 20-min period. After perfusion, the eyeballs were enucleated, the cornea was punctured, and the eyes were immersed in 4% paraformaldehyde in 0.1 M PBS or 0.1 M cacodylate buffer. The anterior segment was then removed, and the resulting eyecups were returned to the fixative for an additional 30 min to 4 h at 4 °C. Eyecups used for frozen sections were rinsed in PBS, cryoprotected in
30% sucrose, then embedded in an optimal cutting temperature medium (Sakura Tissue Tek; VWR, West Chester, PA) and frozen by placing the tissue on an aluminum plate half submerged in liquid nitrogen. Frozen sections (10–15 μm thickness) were prepared on a cryostat, collected onto Superfrost Plus slides (Fisher Scientific, Pittsburgh, PA), and stored at −20 to −30 °C until used.

Immunolabeling was performed as described previously [3, 43]. Frozen sections of fixed eyecups were rehydrated in Hank’s buffered salt solution (HBSS, pH 7.2), and non-specific labeling was blocked using a solution containing 2–10% normal goat serum plus 5% bovine serum albumin, 1% fish gelatin, and 0.1–0.5% Triton X-100 in HBSS (“blocker”). In some experiments, antigen retrieval in 10 mM citrate buffer (pH 6.0; heated to 95 °C) was performed for 30–60 min prior to blocking. The blocker was removed and primary antibody was applied overnight at room temperature. Sections were rinsed in HBSS and an appropriate secondary antibody was applied for 60

rinsed in HBSS and an appropriate secondary antibody was applied for 60–75 min at room temperature. In some experiments, peanut agglutinin (PNA) conjugated to AlexaFluor488 was added to the secondary antibody solution to identify cone photoreceptors. Sections were rinsed and mounted using Prolong Gold plus DAPI or Prolong Diamond plus DAPI (Molecular Probes/Thermo Scientific, Cat# P36935 or P36971, respectively) to retard photobleaching. Specificity of labeling methods was confirmed by omitting primary antibody or substituting normal rabbit serum for primary antibody. Specimens labeled using a primary antibody in combination with PNA showed no bleedthrough of signals between fluorescence channels.

Fluorescence microscopy was performed using an Olympus IX70 inverted fluorescence microscope (Olympus America) fitted with a QiCAM CCD camera controlled via QCapture software (QImaging). Figures were prepared using Photoshop software after calibration of the image scale. Adjustments to brightness and contrast were applied equally to all pixels in the image, if needed, to highlight specific labeling.

For histological evaluation of the retinal structure, the rats were perfused and eyecups were prepared as described above. The eyecups were embedded in paraffin, and sections of 5-μm thickness were cut along the vertical meridian through the optic nerve and stained with hematoxylin and eosin for light microscopic evaluation of the retinal structure. To assess photoreceptor loss, the thickness of the outer nuclear layer (ONL) was measured at 1-mm distances from the optic nerve to the inferior and superior ora serrata and plotted as a “spider” diagram.

Antibodies and Lectins

Elongation of Very Long Chain Fatty Acids 4 (ELOVL4). Anti- ELOVL4 antibody was a rabbit polyclonal antiserum generated by our group [1]. Details of its production and specificity have been described previously [1, 44, 45]. This antibody recognizes a conserved C-terminal epitope and recognizes both the WT and W246G mutant forms of ELOVL4. Anti-ELOVL4 was used at a dilution of 1:300 to 1:500.

Glial Fibrillary Acidic Protein (GFAP). (Millipore, Cat# MAB360. RRID:AB_2109815; mouse monoclonal, clone GA-5). Anti-GFAP recognizes a 51-kDa band on Western blots corresponding to GFAP from human glioma cells (U33CG/343MG) [46]. This antibody does not recognize vimentin. Anti-GFAP was used at a dilution of 1:500–1:2000.

Glutamine synthetase (GS). (Millipore, Cat# MAB302, RRID:AB_2110656; mouse monoclonal, clone GS-6). Anti-GS was raised against full-length GS purified from sheep brain. Anti-GS recognizes a 45-kDa band on Western blots of sheep and rat brain [47]. Anti-GS was used at a dilution of 1:500.

Peanut agglutinin (PNA). Peanut agglutinin specifically labels the extracellular matrix surrounding cone outer segments [48]. PNA conjugated to AlexaFluor 488 (Molecular Probes/ThermoFisher Scientific Cat# L21409; RRID:AB_2315178) was used at a dilution of 1:20–1:50.

Secondary antibodies conjugated to AlexaFluor 568 were purchased from Molecular Probes/ThermoFisher Scientific and used at a dilution of 1:200–1:300 (goat anti-rabbit IgG AlexaFluor568 [Cat# A11036; RRID:AB_143011], goat anti-mouse IgG AlexaFluor568 [Cat# A11004; RRID:AB_2534072]). Normal sera were purchased from Jackson Immunoresearch (normal goat serum [Cat# 005-000-120; RRID:AB_2336990]; normal rabbit serum [Cat# 011-000-120; RRID:AB_2337123]).

Statistical Analysis

Statistical analysis was performed using one-way ANOVA with Tukey’s post hoc test using GraphPad Prism software (GraphPad, La Jolla, CA). Statistical significance was set to \( p < 0.05 \).

Results

Generation of Transgenic SCA34-KI Long-Evans Rats

We generated a novel line of Long-Evans rats using CRISPR/Cas9 by editing one copy of the wild-type (WT) Elovl4 allele in the rat genome to 736T>G (p.W246G) to generate the SCA34-KI rats (Fig. 1). Genotyping and sequencing confirmed genomic editing of one copy of the WT Elovl4 allele to the c. 736T>G SCA34 allele (Fig. 1b). Detailed Sanger and whole-genome sequencing of potential off-target sites confirmed no aberrant indels or insertions in the rat genome (Supplemental Figs. 1 and 2).
Heterozygous SCA34-KI F0 founders were crossed to WT Long-Evans rats to establish a breeding colony. SCA34-KI rats were viable and transmitted the SCA34 allele to the next generation to produce WT, heterozygote (HET), and homozygote (MUT) SCA34-KI pups in Mendelian ratios. Unlike STGD3 homozygous mice that die at birth due to dehydration [11–14], MUT SCA34-KI rats survived and were fertile.

Although the MUT rats survived birth, there were differences in their appearance compared to WT and HET rats (Fig. 2a). MUT rats showed EKV with red, scaly skin and hair loss that was absent from WT and HET animals (Fig. 2a, b). EKV was especially prominent on the eyelids, and delayed eye opening by several days and caused swelling that persisted for several weeks (Fig. 2c–e). Eyelid swelling resolved between P60 and P90, permitting complete opening of the eyelid (Fig. 2f). As would be expected, body size differed significantly between age-matched male and female SCA34-KI rats. Female rats showed no differences in weight among genotypes at either P60 or 6–7 months of age (Fig. 2g–h). In contrast, male MUT rats were smaller than WT male rats by P60 and by 6–7 months of age were smaller than both WT and HET male rats.

The W246G Mutation in ELOVL4 Selectively Affects VLC-SFA Synthesis

Glycerophospholipid levels in WT, HET, and MUT rats were analyzed to assess the effect of the W246G mutation on the ability of ELOVL4 to catalyze the formation of VLC-PUFA in the retina (Fig. 3; details in Supplemental Tables 1–3). VLC-PUFA in the retina were found only in the phosphatidylcholine (PC) fraction (Fig. 3), as reported previously [15, 16]. There were no differences in total VLC-PUFA in PC (ΣVLC-PC) across genotypes, indicating that W246G ELOVL4 was able to catalyze VLC-PUFA formation normally. Some minor differences were found in shorter chain PUFA incorporated into PC and phosphatidylethanolamine (PE), but no differences in PUFA incorporated into phosphatidyserine (PS) were observed (Supplemental Tables 1–3).

ELOVL4 also catalyzes the formation of VLC-SFA in the skin [11–14], Meibomian gland [5], and central nervous system [3]. Because VLC-SFA are expressed at very low levels in central nervous tissue, making them difficult to detect [3], the ability of W246G ELOVL4 to catalyze the formation of VLC-SFA was assessed in skin, where they are much more abundant. The levels of VLC-SFA (28:0, 30:0, and 28:0 + 30:0) were significantly reduced in the skin of MUT rats compared to WT and HET rats (Fig. 4a; Supp. Table 4). Skin VLC-SFA levels did not differ significantly between WT and HET rats. Thus, the W246G mutation in ELOVL4 selectively impaired its ability to catalyze the formation of VLC-SFA. To better understand how the W246G ELOVL4 mutation affected synthesis of SFA levels overall, we also assessed the levels of 24:0 and 26:0, long-chain-SFA precursors needed for the synthesis of VLC-SFA. Consistent with inhibition of VLC-SFA formation, levels of 24:0, a precursor to VLC-SFA formation, were elevated in the skin of HET and MUT rats (Fig. 4b; Supp. Table 4). However, there were no significant differences in levels of 26:0, the direct precursor of 28:0.

The W246G Form of ELOVL4 Does Not Induce Retinal Degeneration

To assess the presence or absence of retinal degeneration, retinal integrity was assessed using optical coherence tomography (OCT), histological, and immunolabeling approaches (Fig. 5). OCT analysis revealed no differences in total retinal thickness or in the thickness of individual retinal layers among WT, HET, and MUT rats between P30 and 6–7 months of age. Histological assessment of outer nuclear layer (ONL) thickness in the retinas of WT, HET, and MUT rats at 6–7 months of age showed no differences among genotypes, indicating that no significant loss of photoreceptors was present by this time. Analysis of eye diameter at 6–7 months of age also showed no differences among genotypes, indicating that the W246G mutation in ELOVL4 did not affect the size of the eye.

Immunolabeling for ELOVL4 and other markers for photoreceptors, Müller cells, and neurodegeneration was performed to further assess potential changes in the SCA34-KI rat retina with age (P45 to 6–7 months of age). Labeling for ELOVL4 in the WT, HET, and MUT retinas was normal (Fig. 6), with high levels of labeling in photoreceptor cell bodies in the ONL. Labeling for PNA also was normal, confirming that cones and their synapses were maintained in retinas of all genotypes (Fig. 7). Müller cells showed no elevated labeling for GFAP, a sensitive marker of reactive gliosis and retinal degeneration (Fig. 8). Labeling for glutamine synthetase, a marker for Müller cells, showed normal Müller cell morphology and distribution in retinas of all genotypes (Supplemental Fig. 3).

The W246G SCA34 Mutation in ELOVL4 Impairs Retinal Function

Mutations in ELOVL4 that cause STGD3 profoundly impair retinal light responses and cause early-onset macular degeneration in STGD3 patients [26, 27, 31–33, 36]. In contrast, human SCA34 patients typically show no retinal disease. To determine whether the W246G mutation in ELOVL4 affected retinal function, electroretinogram (ERG) recordings were made from WT, HET, and MUT rats. Rats of all three genotypes showed rod- and cone-driven ERG responses at P30 and P60, but eyelid swelling in MUT rats partially obstructed the pupil, confounding interpretation of results at these time.
points. Therefore, ERG studies focused on P90, P120, and 6–7 month time points, after eyelid swelling resolved.

Analysis of rod-driven, scotopic ERG responses revealed that the scotopic a-wave amplitude of MUT rats was reduced compared to that of WT and HET rats (Fig. 9a–e). At P90, the scotopic a-wave showed significantly reduced amplitude only at relatively high flash intensities, but significant differences in amplitude were present at all but the lowest flash intensity at P120 and 6–7 months. The scotopic b-wave of MUT rats showed a similar reduction in amplitude compared to that of WT and HET rats (Fig. 9b–f). There were no differences in scotopic a-wave or b-wave amplitudes between WT and HET rats at any flash intensity or age. To assess whether synaptic transmission of signals from rods to rod bipolar cells was...
affected, the scotopic b-wave to a-wave ratio was assessed (Fig. 10). The b-wave to a-wave ratio of MUT rats consistently was slightly reduced compared to that of age-matched WT and HET rats, suggesting that MUT rats may show a small reduction in amplification of rod signals transmitted to the inner retina. Latency of the scotopic a-wave and time to peak of the b-wave were not substantially affected, although a significant difference in a-wave latency was present at P90 at the lowest flash intensity tested (Supplemental Fig. 4).

The W246G mutation in ELOVL4 also affected cone-driven ERG responses. MUT rats showed slightly reduced amplitudes for the photopic a-wave, and S-cone- and M-
Fig. 5 Retinal layering is normal and photoreceptors are preserved in the SCA34-KI rat retinas to at least 6–7 months. a–d Optical coherence tomography (OCT) analysis of WT, HET, and MUT SCA34 rat retinas from P30 to 6–7 months of age showed no differences in thickness of any retinal layer at any time point. (Data shown as mean ± St Dev. Analysis by 1-way ANOVA with Tukey’s post hoc test. P30: 12 WT, 13 HET, 16 MUT. P90: 9 WT, 14 HET, 10 MUT. P120: 2 WT, 2 HET, 5 MUT. 6–7 months: 7 WT, 11 HET, 7 MUT). e Eye diameter in 6–7-month-old rats also is unaffected by the W246G mutation in ELOVL4. (One-way ANOVA with Tukey’s post hoc test. Data shown as mean ± St Dev. n = 3 WT, 3 HET, 3 MUT). f Histological analysis of outer nuclear layer thickness shows no differences among WT, HET, and MUT rats along the superior to inferior axis at 6–7 months of age, indicating no significant photoreceptor degeneration. (One-way ANOVA with Tukey’s post hoc test. Data shown as mean ± St Dev. n = 3 WT, 3 HET, 3 MUT). PRL, photoreceptor layer; ONL, outer nuclear layer; OPL, outer plexiform layer; INL, inner nuclear layer; IPL, inner plexiform layer; GCL, ganglion cell layer.
cone-driven a-waves compared to WT and HET at P90, P120, and 6–7 months; however, these differences were not statistically significant (Fig. 11a–i). Cone-driven a-wave latencies showed no differences across genotypes (Supplemental Fig. 5). In contrast, the flicker ERG a-wave of MUT rats differed significantly from that of WT and HET rats (Fig. 11j–l), indicating that the W246G mutation in ELOVL4 affected the cone-driven flicker responses at the level of the cones themselves. The flicker ERG a-wave showed no significant differences between WT and HET rats. Cone-driven flicker ERG a-wave latencies showed no differences across genotypes (Supplemental Fig. 6).

Cone-driven b-wave responses were more strongly affected by the W246G mutation in ELOVL4. The photopic b-wave amplitude of MUT rats was significantly reduced compared to that of WT rats at all ages tested (Fig. 12a–c). Similarly, the S-cone, M-cone, and flicker ERG b-wave amplitudes of MUT rats also were reduced significantly compared to those of WT.
rats (Fig. 12d–l). In some cases, additional differences in cone-driven b-wave amplitudes were noted among MUT, HET, and WT control rats (Fig. 12a–i). Cone-driven b-wave time-to-peak measures showed no differences across genotypes at any age (Supplemental Figs. 6 and 7).

Discussion

Several different mutations in ELOVL4 cause age-related SCA34 characterized by prominent progressive cerebellar atrophy and ataxia in patients. Here, we report successful application of the CRISPR/Cas9 approach to generate the first rat model of human SCA34 caused by the c. 736T>G (p.W246G) mutation in the human ELOVL4 as reported by Ozaki et al. [35]. The F0 heterozygous rats were viable and transmitted the 736T>G (p.W246G) mutation to the next generation as determined by genotyping and sequencing. Unlike mice in which the homozygous expression of the STGD3 mutant ELOVL4 or global deletion of ELOVL4 causes neonatal lethality [11–14], homozygous W246G MUT rats survive birth and thrive to adulthood, although the males are smaller in size compared to age-matched WT and HET rats. Both Sanger sequencing and whole gDNA sequencing confirm no off-target effects in the HET and MUT rats. Retinal ELOVL4 expression in all three genotypes is similar, suggesting that the W246G mutation did not affect retinal Elovl4 expression and protein synthesis.

The physiological, biochemical, and anatomical studies presented herein indicate that the W246G mutation in ELOVL4 that causes SCA34, a neurodegenerative disease that strikes the cerebellum and can be associated with EKV in the skin, also affects retinal function in the absence of retinal degeneration. This is the first report to characterize functional retinal deficits associated with SCA34-causing mutations in ELOVL4. Functional and anatomical studies showed that homozygous W246G ELOVL4 mutation depressed retinal light responses in the absence of neurodegeneration. Biochemical studies indicate that the W246G ELOVL4 mutation impaired production of VLC-SFA, but spared synthesis of VLC-PUFA.

The retinal and skin phenotypes of HET rats are similar to those of human SCA34 patients heterozygous for the W246G ELOVL4 mutation [35]. HET rats, like human SCA34 patients with the W246G ELOVL4 mutation, do not show severe EKV [35], although EKV is present in SCA34 patients with other ELOVL4 mutations [28–30]. Similarly, HET rats show retinal responses comparable to WT rats, consistent with the reported absence of functional retinal deficits in human SCA34 patients [35]. No human patients homozygous for any SCA34-causing ELOVL4 mutation have been reported. Speculation is that this results from these mutant alleles being very rare in the population and/or that homozygous inheritance of these alleles would be embryonically lethal. Therefore, the birth and survival of MUT rats were unexpected and presented a unique opportunity to further explore ELOVL4 function.
ELOVL4 is essential for synthesis of both VLC-PUFA and VLC-SFA [1–3]. In the retina, VLC-PUFA are the main product and are incorporated into PC and concentrated in the outer segment membranes [15–18]. VLC-PUFA levels in WT, HET, and MUT rat retinas did not differ, indicating that W246G ELOVL4 supported normal VLC-PUFA biosynthesis. Some small differences in PUFA with shorter chain lengths, which are not produced directly by ELOVL4, were detected in SCA34-KI rats. In contrast to VLC-PUFA, the homozygous W246G mutation in ELOVL4 impaired synthesis of VLC-SFA (28:0 and 30:0) by 50% or more in the skin of

![Fig. 8](Mol Neurobiol (2020) 57:4735–4753) Müller cells in the SCA34-KI rat retina are not reactive. **a–c** Müller cells in the retina of P45 WT, HET, and MUT rats show little labeling for glial fibrillary acidic protein (GFAP) as is typical of Müller cells in the healthy retina. Labeling in blood vessels (bv) is non-specific. **d–f** Müller cells in the retina of 6–7-month-old (6–7 Mo) WT, HET, and MUT rats also show little labeling for GFAP, indicating the absence of glial reactivity associated with retinal degeneration. PRL, photoreceptor layer; ONL, outer nuclear layer; OPL, outer plexiform layer; INL, inner nuclear layer; IPL, inner plexiform layer; GCL, ganglion cell layer. Scale bars = 50 μm.

![Fig. 9](Mol Neurobiol (2020) 57:4735–4753) Effects of the W246G mutant form of ELOVL4 on the scotopic ERG. Homozygous inheritance of the W246G form of ELOVL4 reduces scotopic a-wave and b-wave ERG responses. **a, c, e** Scotopic ERG a-wave amplitude for WT, HET, and MUT rats at P90, P120, and 6–7 months of age. **b, d, f** Scotopic ERG b-wave amplitude for WT, HET, and MUT rats at P90, P120, and 6–7 months of age. (Data shown as mean ± SEM. One-way ANOVA with Tukey’s post hoc test. Asterisks indicate statistical significance at $p < 0.05$ (*), $p < 0.01$ (**), and $p < 0.001$ (***)). P90: 14 WT, 14 HET, 17 MUT. P120: 13 WT, 15 HET, 15 MUT. 6–7 Mo: 9 WT, 14 HET, 7 MUT.)
Mutations in ELOVL4 that cause STGD3 all have deletion of the C-terminal ER retention motif that leads to mislocalization of the enzyme from the ER, reduced ELOVL4 protein levels, and a dominant negative effect on VLC-PUFA biosynthesis [49, 50]. In contrast, W246G ELOVL4 retains the ER retention motif in the C-terminal and would be directed to the ER membranes. However, in the homozygous state, the W246G ELOVL4 mutation seems to affect VLC-SFA biosynthesis selectively. In contrast, VLC-PUFA synthesis was spared in HET and MUT rats, while VLC-SFA synthesis was impaired in the skin of MUT rats, indicating that the W246G ELOVL4 mutation affects VLC-SFA synthesis while retinal VLC-PUFA synthesis was spared.

The molecular basis for this selective impairment of VLC-SFA synthesis is unknown, but one possibility is that the W246G mutation alters the structure in or around the active site such that access of the long-chain-SFA precursor (26:0) to the site is impaired, whereas access of the VLC-PUFA precursors (26:Xn3 and 26:Yn6) is not. Additionally, ELOVL4 could have differential protein-protein interactions in the retina that lead to differential tissue-specific effects on VLC-SFA and VLC-PUFA synthesis. Importantly, the effect of the W246G mutation on the molecular structure of the ELOVL4 protein remains unknown.

Human SCA34 patients heterozygous for the W246G ELOVL4 mutation have no reported clinical deficits in retinal function [35], although no ERG results were presented for these patients. Similarly, the rod- and cone-driven ERG responses of HET and WT rats were comparable, consistent with our lipid analyses that showed no differences in VLC-fatty acid levels between these genotypes. In contrast, MUT rats showed a clear functional retinal deficit. MUT rats showed significant reductions in scotopic ERG a-wave and b-wave amplitudes compared to WT and HET rats. The effect of the W246G ELOVL4 mutation on the photopic ERG was less pronounced. The photopic a-wave of MUT rats differed from that of WT and HET rats only for the flicker ERG, but several significant differences in photopic ERG b-wave measures were found between the MUT rats and WT and HET rats. These deficits appeared by 90 days of age and persisted until at least 6–7 months of age, the oldest age tested in this study.

Recently, a study of 10 affected members of an American family with a novel heterozygous ELOVL4 mutation (I171T) that results in spinocerebellar ataxia and retinitis pigmentosa was reported [37]. This mutation causes a late-onset spinocerebellar ataxia similar to other SCA34-causing ELOVL4 mutations [28–30, 35]. This SCA is characterized by average onset in the 5th decade, often accompanied by dysarthria and handwriting changes, but no autonomic dysfunction or dermatologic disease [37]. In contrast to other previously known SCA34-causing ELOVL4 mutations, which

**Fig. 10** Ratio of scotopic b-wave to a-wave amplitude. The scotopic b-wave to a-wave ratio in MUT rats shows a consistent reduction compared to the b-wave to a-wave ratio of age-matched HET and WT rats, suggesting that amplification of rod signals transmitted to the inner retina is reduced in MUT rats. (Data shown as mean ± SEM. One-way ANOVA with Tukey’s post hoc test. Asterisks indicate statistical significance at $p < 0.05$ (*), $p < 0.01$ (**), and $p < 0.001$ (***)). P90: 14 WT, 14 HET, 17 MUT. P120: 13 WT, 15 HET, 15 MUT. 6–7 Mo: 9 WT, 14 HET, 7 MUT.)

MUT rats compared to WT and HET rats. Synthesis of 30:0 was more severely affected than 28:0. No statistical differences in VLC-SFA levels were detected between WT and HET rats. Thus, the W246G mutation in ELOVL4 selectively impairs VLC-SFA synthesis while leaving VLC-PUFA overtly unaffected.
showed no retinal degeneration [28–30, 35], the I171T ELOVL4 mutation was associated with retinitis pigmentosa, a degenerative retinal disease, in 4 of the 10 family members examined (three males and one female).

The W246G ELOVL4 mutation affected the ERG b-wave, which is dependent on signal transmission from photoreceptors to second-order bipolar cells, more strongly than the a-wave, which reflects the light-driven responses of the photoreceptors themselves. The finding that the scotopic ERG b-wave to a-wave ratio of MUT rats was consistently lower than that of age-matched WT and HET rats also suggests a reduction of signal amplification across the photoreceptor to the bipolar cell synapse in MUT rats. A recent study in our laboratory in mice homozygous for the 5-base-pair STGD3 deletion in ELOVL4 showed that slices of MUT hippocampus had aberrant presynaptic neurotransmitter release kinetics and seizure activity [3]. Re-supply of exogenous VLC-SFA in MUT cultured hippocampal neurons restored presynaptic release kinetics to normal levels. A role for ELOVL4 and its VLC-SFA products in photoreceptor transmission is supported by the finding of VLC-SFA in retinal sphingolipids [7] and that mouse photoreceptors with the 5-base-pair STGD3 deletion in ELOVL4 have smaller synaptic vesicles and reduced b-wave amplitude compared to WT mice [44, 51].

The functional deficits in the ERG of MUT rats are unlikely to arise from neurodegeneration. Anatomical analyses of retinal layer thickness and photoreceptor numbers revealed no indications of neurodegeneration in the retina of HET or MUT rats.
out to 6–7 months of age. Immunolabeling studies confirmed that photoreceptors expressed ELOVL4 appropriately across all genotypes and appeared structurally normal. Müller glial cells, which show striking reactivity and elevated levels of GFAP in the presence of degeneration, showed no reactivity in the HET or MUT retina. Our lipid analyses showed that VLC-PUFA were present in PC at normal levels in the HET and MUT retinas, which would maintain the retrograde elovanoid feedback signaling from the retinal pigmented epithelium to the photoreceptors needed to promote photoreceptor survival [52], consistent with our studies showing an absence of neurodegeneration in the HET and MUT retinas.

Diseases arising from ELOVL4 mutations likely represent a continuum arising from differential effects of the various mutations on ELOVL4’s function in the synthesis of VLC-SFA and VLC-PUFA. Our current studies suggest that ELOVL4 mutations that selectively impair VLC-SFA synthesis compromise synaptic function in the CNS and the barrier function of the skin. The W246G ELOVL4 mutation examined here selectively impaired synthesis of VLC-SFA, with no significant effect on VLC-PUFA synthesis, leading to compromised ERG responses in MUT rats in the absence of retinal degeneration. These results are consistent with the aberrant presynaptic release and seizures induced by VLC-SFA deficiency due to homozygous knock-in of the 5-base-pair STGD3 mutation [3]. The appearance of EKV in only MUT rats is also consistent with the VLC-SFA deficiency associated with homozygous W246G ELOVL4 mutation. Heterozygous ELOVL4 mutations

Fig. 12 Effects of the W246G mutant form of ELOVL4 on the photopic ERG b-wave. Homozygous inheritance of the W246G form of ELOVL4 reduces cone-driven ERG b-wave responses. a, b, c Photopic ERG b-wave amplitude for WT, HET, and MUT rats at P90, P120, and 6–7 months of age. d, e, f S-cone-driven ERG b-wave amplitude for WT, HET, and MUT rats at P90, P120, and 6–7 months of age. g, h, i M-cone-driven ERG b-wave amplitude for WT, HET, and MUT rats at P90, 120, and 6–7 months of age. j, k, l Cone-driven flicker ERG b-wave amplitude for WT, HET, and MUT rats at P90, P120, and 6–7 months of age. (Data shown as mean ± SEM. One-way ANOVA with Tukey’s post hoc test. Asterisks indicate statistical significance at $p < 0.05$ (*), $p < 0.01$ (**), and $p < 0.001$ (***)). P90: 14 WT, 14 HET, 17 MUT. P120: 13 WT, 15 HET, 15 MUT. 6–7 Mo: 9 WT, 14 HET, 7 MUT)
that cause SCA34 with EKV [28–30] presumably compromise VLC-SFA synthesis more severely than the W246G ELOVL4 mutation. The novel I171T ELOVL4 mutation that causes SCA accompanied by retinitis pigmentosa in some patients [37] would be predicted to impair synthesis of VLC-SFA and also VLC-PUFA to some degree. Heterozygous ELOVL4 mutations that cause STGD3, a severe juvenile-onset photoreceptor degeneration [27, 32, 36], compromise VLC-PUFA biosynthesis [49, 50]. STGD3 patients have no other reported CNS or skin disease, suggesting that heterozygous STGD3 mutations may not alter VLC-SFA synthesis substantially, although complete deletion of ELOVL4 expression in mice is neonatal lethal. Homozygous inheritance of STGD3-causing ELOVL4 mutations has not been reported in humans, but mice homozygous for the 5-bp STGD3 mutation (but kept alive by transgenic expression of Elov4 in their skin) have disrupted synaptic function and severe seizures at P18 [3], indicating that homozygous inheritance of STGD3 alleles compromises synthesis of both VLC-PUFA and VLC-SFA. Finally, ELOVL4 neuro-ichihtyotic syndrome, the most severe disease associated with ELOVL4 mutations, is characterized by seizures, spasticity, intellectual disability, ichthyosis, and early death. This syndrome arises from homozygous ELOVL4 mutations that cause early truncation of the protein [25, 34], presumably eliminating all VLC-SFA and VLC-PUFA synthesis.

Based on our enzyme activity studies and a previous study showing that VLC-SFA are enriched in synaptic vesicles and regulate presynaptic release kinetics in the brain [3], the most likely explanation for impaired ERG responses in MUT rats is that reduced production of VLC-SFA in photoreceptors impairs neurotransmission from photoreceptors to the second-order neurons. The possibility that ELOVL4 is involved in the biosynthesis or regulation of other, yet unidentified, lipid products cannot be dismissed. The role of ELOVL4 in the production of VLC-SFA was not recognized until a decade after its role in VLC-PUFA biosynthesis was first identified [1]. The possibility that VLC-SFA might serve some unrecognized signaling function cannot be dismissed. Only recently has it been recognized that VLC-PUFA incorporated into phosphatidylethanolamine in photoreceptor outer segment membranes are metabolized by the retinal pigmented epithelium to form elovanoid signaling molecules that provide feedback signals that are essential for photoreceptor survival [52]. Further mechanistic studies will be needed to resolve these questions.

Taken together, these findings provide additional support for the notion that ELOVL4 and its VLC-SFA and VLC-PUFA products are critical to different aspects of neural function. The W246G mutation in ELOVL4 appears to selectively reduce, but not eliminate, production of VLC-SFA, leading to impaired transmission from photoreceptors to the inner retina, while maintaining normal VLC-PUFA biosynthesis needed for survival of photoreceptors.

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Authors’ Contributions All of the authors contributed to the drafting and editing of the manuscript. MPA, REA, and DMS were responsible for the study concept and design. MPA generated the SCA34-KI rat line. MPA and KLJ analyzed Sanger and gDNA sequencing data. Experiments and data analysis were performed by MPA, MAS, RSB, MTS, WB, AM, KLJ, DMS, and REA. All of the authors agree to be accountable for all aspects of the studies.

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Data Availability The raw data will not be deposited in a public repository due to the large size of the image files and the complexity of annotating the large volume of individual files.

Compliance with Ethical Standards

Conflict of Interest The authors declare that they have no conflicts of interest.

Ethics Approval All animal procedures were approved by the University of Oklahoma Health Sciences Center Institutional Animal Care and Use Committee and conformed to the National Institutes of Health Guide for the Care and Use of Laboratory Animals, the Association for Research in Vision and Ophthalmology Resolution on the Use of Animals in Research, and the US Public Health Service guidelines.

Consent to Participate Not applicable

Consent for Publication Not applicable

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