Toward genome editing in X-linked RP-development of a mouse model with specific treatment relevant features

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Genome editing represents a powerful tool to treat inherited disorders. Highly specific endonucleases induce a DNA double strand break near the mutant site, which is subsequently repaired by cellular DNA repair mechanisms that involve the presence of a wild type template DNA. In vivo applications of this strategy are still rare, in part due to the absence of appropriate animal models carrying human disease mutations and knowledge of the efficient targeting of endonucleases. Here we report the generation and characterization of a new mouse model for X-linked retinitis pigmentosa (XLRP) carrying a point mutation in the mutational hotspot exon ORF15 of the \textit{RPGR} gene as well as a recognition site for the homing endonuclease I-SceI. Presence of the genomic modifications was verified at the RNA and protein levels. The mutant protein was observed at low levels. Optical coherence tomography studies revealed a slowly progressive retinal degeneration with photoreceptor loss starting at 9 months of age, paralleling the onset of functional deficits as seen in the electroretinogram. Early changes to the outer retinal bands can be used as biomarker during treatment applications. We further show for the first time efficient targeting using the I-SceI enzyme at the genomic locus in a proof of concept in photoreceptors following adeno-associated virus mediated gene transfer in vivo. Taken together, our studies not only provide a human-XLRP disease model but also act as a platform to design genome editing technology for retinal degenerative diseases using the currently available endonucleases. (Translational Research 2019; 203:57 – 72)
Schlegel J, et al.

At A Glance Commentary

Background

Genome editing in the retina is currently at the height of debate with regard to clinical translation, yet important aspects of the in vivo application are still poorly understood, in part due to the lack of animal models with human-like mutations.

Translational Significance

Our mouse model contains a human-like mutation in the RPGR-ORF15 gene, in which mutations in human cause XLRP. Due to its degenerative phenotype and the presence of an I-SceI restriction site, this mouse model is well suited to study the efficacy of all classes of endonucleases and to characterize the efficacy of therapeutic applications in retinal neurons in vivo.

INTRODUCTION

Retinitis pigmentosa (RP) is a group of heterogeneous disorders affecting primarily rod photoreceptors accounting for about 3 million patients worldwide. Mutations in the gene encoding the retinitis pigmentosa GTPase regulator (RPGR) are a major cause of RP in humans. They account for 15%–20% of sporadic RP cases and up to 80% of all X-linked RP cases. Patients typically suffer from early night-blindness with reduction of the visual field and reduced visual acuity, indicating an implication of both rods and cones in the pathology. Optical coherence tomography (OCT) analyses show rapid loss of outer retinal structures that progresses over time.

The RPGR gene consists of 19 exons and encodes multiple alternatively spliced isoforms. Two major RPGR isoforms RPGR\(^\text{const}\) (encoded by 19 exons) and RPGR\(^\text{ORF15}\) (terminates within intron 15) have been reported, the latter being uniquely expressed in photoreceptors. Physiologically, both RPGR protein variants are located in the connecting cilium (CC) of the vertebrate photoreceptor, where they take part in a protein complex associated with the intraflagellar transport machinery that facilitates cilary protein trafficking.

In addition, the RPGR\(^\text{const}\) protein is also important for the transport of cargo toward the cilium via prenylation of the C-terminus by PDE6delta.

Most of the disease-causing mutations in humans are found in RPGR\(^\text{ORF15}\). The terminal exon ORF15 (exon 15+ part of intron 15) of the RPGR\(^\text{ORF15}\) isoform is a mutational hot spot, accounting for >60% of RPGR mutations. This exon consists of extensive purine-rich repeats and encodes for glycine (G) and glutamic acid (E) residues. The majority of the exon ORF15 mutations result in a frameshift due to deletions or duplications. Although the mechanism of action of these mutations is not completely understood, an effect on the isoelectric point or alterations in the post-translational modifications of RPGR\(^\text{ORF15}\) have been proposed.

Recently, several groups have shown the therapeutic effect of delivering the full-length RPGR\(^\text{ORF15}\) with either wild type exon ORF15 or as a codon optimized version via adeno-associated virus (AAV) vector mediated gene transfer. While production of a recombinant protein has been shown to restore function in the dog model of the disease (X linked progressive retinal atrophy, XLPRA2) and in a knock out (KO) mouse model, potential drawbacks with this form of treatment are always related to the artificial control of transgene expression or the interaction of residual endogenous protein with the recombinant protein.

In vivo genome editing represents a powerful tool to circumvent this issue, as it enables the correction of the disease-causing mutation in the genome, which would result in restoration of endogenous protein production. It is based on the cell’s capacity to repair DNA double strand breaks (DSBs) by either error-prone nonhomologous end-joining (NHEJ), or in the presence of a template DNA by homology-directed repair or microhomology-mediated end-joining. Induction of DSB at the target locus can be performed by specific endonucleases, of which a comprehensive toolbox exists, that is, homing endonucleases (HE) such as I-SceI, zinc-finger nucleases, TALE nucleases, or the CRISPR-Cas system. However, size limitations in AAV vectors generate problems when TALE nucleases or CRISPR-Cas systems are used together with the appropriate template DNA. Since HEs have been long considered to be un-programmable, they were not used to develop therapeutic strategies, even though their relatively small size and low toxicity render them an interesting tool. Recent progress in programming HEs for eukaryotic target loci may overcome this issue.

Because the cellular DSB repair mechanisms are difficult to control in vivo and off target toxicity remains a major issue, current strategies to target exonic mutations in inherited retinal dystrophy genes in vivo are still comparably risky. The absence of appropriate in vivo model systems to study genome editing in the retina in detail further complicates that issue. Standard knock-out models with large deletions or insertions in the genome do not represent ideal systems to study precise genome editing. Disease models based on point mutations are rare and only available for few inherited retinal dystrophies.

Since most of the disease causing mutations in X-linked retinitis pigmentosa (XLRP) are located in the
RPGR-ORF15 exon and genome editing application are dependent on realistic animal model systems, the aim of this study was to develop a mouse model for XLRP containing a human-like mutation in the ORF15 exon that also contains specific features important for the development of therapeutic genome editing strategies. These features include a point mutation (deletion) similar to that in patients, other silent mutations for screening purposes and the introduction of an I-SceI recognition site in order to study this enzymes cutting efficiency in a therapeutically relevant system. Here we report the generation and characterization of a new mouse model of XLRP carrying a point mutation in exon ORF15 of the Rpgr gene and an I-SceI site downstream of exon ORF15, and provide evidence for NHEJ at the RPGR locus following I-SceI expression in cell culture and in photoreceptors in vivo following AAV mediated gene transfer.

MATERIALS AND METHODS

Ethics statement. The animals were maintained and experimental procedures complied in accordance with the Animal Welfare guidelines of the local German authorities and the ARVO Statement for the Use of Animals in Ophthalmic and Vision Research. The Institutional Animal Care and Use Committee of the local government approved all procedures (permission 22/2011 and 79/2016). Surgery and in vivo examinations were performed under anesthesia and all efforts were taken to avoid suffering of the animals. Mice were euthanized by cervical dislocation. Mice were kept and bred in a 12-hour light and/or dark cycle with unlimited access to food and water in the animal facility of the Max-Planck-Institute for Heart and Lung Research in Bad Nauheim or in the animal facility of the Institute of Biochemistry at the Justus-Liebig-University Giessen.

Generation of the B6J.Sv129-Rpgr<sup>tm1Sti</sup> mouse model. The generation of the targeting vector was done using the method of homologous recombination by gap repair. A <i>bacterial artificial chromosome</i> BAC plasmid (pBACe3.6) containing the genomic locus of <i>Rpgr</i> was purchased from ImaGene GmbH (Berlin, Germany). A part of the genomic RPGR locus containing exons 13, Orf14/15, as well as a loxP/FRT flanked Neomycin cassette for positive selection were combined via gap repair into the pKOIV2 plasmid (kindly provided by Thomas Braun, MPI Bad Nauheim, Germany), already containing a DTA (diphtheria toxin A) cassette (Fig 1, A). To introduce the 4 mutations, the <i>Orf15</i> sequence was cloned into a pL452 plasmid and the defined point mutations were integrated via mutagenesis polymerase chain reaction (PCR). The mutated target-sequence was cloned into the pKOIV2 by directed cloning via Sal-I and Sac-II restriction digestion, thus finalizing the targeting vector.

For homologous recombination in embryonic stem (ES) cells, the targeting vector was linearized by NotI and electroporated into male ES cells (C57BL/6/129sv hybrid cells). Transgenic clones were selected by G418 resistance and screened for correct homologous recombination by long range PCR using primers binding within the Neomycin sequence [TTCTGAGGGGTCAATTCTCTA-GAGCTCGC] and outside the recombination site [AGATCTGACGCCCTTCTCTGTGTTTCTGAAAG] (Fig 1, A, external sonde eS), and by standard PCR spanning 500 bp around the newly introduced Xbal site and the pathologic frame shift mutation (Fig 1, A, internal sonde iS) using primers [GGAAATGTAGTAGTG-GAGCA] and [GTGTATGGAAATATGTCGGAATTTG] (Fig 1, B). Positive clones were injected into C57BL/6J blastocysts and F1 hybrid offspring was tested for the presence of DNA changes by PCR as described above. Genomic DNA was isolated from tail to tip tissue with the DNeasy Tissue Kit (Qiagen, Hilden, Germany). The neomycin cassette was deleted from the genome via Cre-recombination of the F1 hybrids. Offspring was subsequently back-crossed into a C57BL/6J background.

mRNA isolation and RT-PCR. RNA was isolated from collected neuroretinai homogenized by freeze and thaw cycles using liquid nitrogen. RNA was isolated using the AllPrep RNA/Protein Kit (Qiagen, Hilden, Germany).

For cDNA synthesis, 80 ng of total retinal RNA was reverse-transcribed using the PrimeScript cDNA Synthesis Kit (Takara, Saint-Germain-en-Laye, France). PCR amplification was performed using the PrimeScript Polymerase (Takara, Saint-Germain-en-Laye, France). The same primer pairs as for the initial screening (internal sonde iS) were used. Since the primers were placed outside the repetitive region and the PCR products were comparably small, we were able to generate sequencing data from gelpurified PCR products (Seqlab, Göttingen, Germany).

Western blot. Neuroretinai (n = 3) from 3 months old wild type, mutant, and age matched <i>Rpgr</i><sup>ko</sup> animals were harvested and lysed using the RIPA (radioimmunooprecipitation assay) buffer. Fifty micrograms of total protein extract was analyzed by SDS-PAGE and immunoblotting using the RPGR<sup>ORF15</sup> antibody, which identifies an epitope in the C-terminal region of the ORF15 frame, as described. The antibody was kindly provided by Anand Swaroop (NEI).
Fig 1. Generation of the mouse model. (A) Scheme of the recombination strategy of the targeting vector and the genomic mouse DNA applied for the generation of this mouse model. The locations of the mutational sites marked with asterisks and amplification sites for screening via PCR are indicated by brackets. The “floxed” neomycin-cassette is deleted via Cre recombination from the genome. Inserted mutations are specified by 1–4.
Two different techniques were used to prepare the retinal tissue.

(1) Enucleated eyes were embedded unfixed in OCT medium (optimal cutting temperature −20°C, Reichert Jung, Wetzlar, Germany) before preparing sections of 16 µm, which were subsequently fixed for 13 minutes at room temperature in 4% paraformaldehyde in 0.1 M phosphate buffered saline (PBS).

(2) Eyecups were harvested and immediately fixed for 45–70 minutes in 4% paraformaldehyde. Eyecups were then cryo-protected in graded sucrose solutions. Vertical sections (16 µm) were cut with a cryomicrotome (SLEE Medical GmbH, Mainz, Germany) and collected on Superfrost slides.

Slides were incubated over night at 4°C with S1 and S3 rabbit polyclonal RPGR antibodies, diluted 1:200, or with a cocktail of rabbit polyclonal RPGR antibody and acetylated tubulin antibody, diluted 1:300 and 1:5.000 respectively. Primary antibodies were diluted in PBS containing 3% normal donkey serum (NDS) and 0.5% Triton X-100; S1 and S3 rabbit polyclonal RPGR antibodies were kindly provided by Dr. Tiansen Li (NEI). Rabbit polyclonal RPGR-ORF15 antibody targeting the C-terminal region of the ORF15 exon (see reference 9 for further information) was used as well. Monoclonal acetylated tubulin antibody was commercial (# T 6793, Sigma-Aldrich, Darmstadt, Germany). After washing with 0.1 M PBS, sections were incubated with Alexa-Fluor488 donkey antirabbit (1:500) secondary antibody (Molecular Probes, UK, Europe, #A21206) for 1.5 hours at room temperature. Nuclei were stained by DAPI (1 µg/ml; Molecular Probes, UK, Europe, #D1306) during incubation with secondary antibodies. For double immunolabeling, the respective primary and secondary antibodies were applied as a cocktail. Sections were mounted with antifade medium (Aqua Poly/mounting medium, Polysciences, #18606) and examined by laser scanning confocal microscopy (Olympus FV10i, Hamburg, Germany; equipped with argon and HeNe lasers).

Retinal histology. Enucleated eyes were fixed in yellow fix solution (45% formaldehyde, 45% PBS, 8% glutaraldehyde, and 2% picric acid) for 4 hours and rinsed in PBS overnight. The retinæ were then post-fixed in 1% osmium tetroxide (2 hours) at pH 7.3, dehydrated in graded ethanols, and embedded in EMBED-812 epoxy resin (all reagents from Science Services, Munich, Germany; automated LYNX-Tissue processor Leica, Bensheim, Germany). After 48 hours of heat polymerization at 60°C, semithin (0.5 µm) tissue sections were obtained and stained with toluidine blue. Pictures were taken with the Keyence BC8000 digital microscope (Keyence Inc, Japan).

Optical coherence tomography. The MICRON III fundus camera was used with modular spectral domain OCT technology (Phoenix Research Inc, CA), specifically developed for rodent applications (axial optical resolution 3 µm), and B scans were averaged over 10 frames. At least 3 animals per time point were analyzed over time. Total retina and outer nuclear layer (ONL) thickness were measured over the entire length of the scan (excluding the papilla in central scans) using the Insight OCT segmentation software (Phoenix Research Inc, Calif). Mean thickness and standard deviation was calculated from each group of animals. Significance was tested using a student t test with P < 0.05.

Electroretinography. Animals were anaesthetised by injection of xylazine (9 mg/kg body weight; 2% Xylazine, Ceva Santé Animale, Libourne, France) and ketamine (90 mg/kg body weight; 10% Ketavet # 9089.01.00, Bela-Pharm GmbH & Co. KG, Vechta, Germany). Full-field stimulation was performed using an Espion E3 Electrophysiology system with a Color-Dome full-field Desktop Ganzfeld stimulator (Diagnosys LLC, Lowell).

Mice were dark-adapted overnight and anesthetized under dim red light. Pupils were dilated with Tropicamide (1%, # 23.0311.03, Mydriaticum Stulln, Pharma Stulln, Stulln, Germany). For local anaesthesia Propar-akain-Pos (0.5%, Ursapharm, Saarbrücken, Germany) was used in addition to Methocel (2%, # 421503, OmnipVision GmbH, Puchheim, Germany) application to keep the cornea moist. A gold-wire ring electrode was used as recording electrode, and subcutaneous
needle electrodes as reference (between the ears) and ground (tail) electrode. Body temperature was maintained at 37°C using a heating pad.

Electroretinographies (ERGs) were performed as previously described. Briefly, single flash recordings were obtained both under dark-adapted (scotopic) and light-adapted (photopic) conditions. Light adaptation was performed with a background illumination of 30 cd/m² presented for 10 min. White stimuli were presented with increasing intensities, reaching from 10⁻⁴ cd s/m² to 25 cd s/m², divided into 10 steps of 0.5 and 1 log cd s/m². Twelve to fifteen responses were averaged with an interstimulus interval of 5 seconds or 17 seconds (for 1, 3, 10, and 25 cd s/m²).

Significance was tested using a student t test with P < 0.05.

**Targeting of the I-SceI site in vitro.** The ORF15 exon from the targeting vector that was used to generate the mouse model was cloned into the pcDNA 3.1 (-) vector, under control of a cytomegalovirus early enhancer promoter (CMV) promoter (Invitrogen/ThermoFisher SCIENTIFIC). Four micrograms of the vector was linearized with EcoRI and then transfected into HEK293 cells in a 6-well plate as mentioned earlier. Forty-eight hours following the transfection, 0.5 μg/mL Genetin (Life Technologies) was added to the medium for selection. Surviving cell foci were picked and transferred to 24 well plates. The selection process was repeated twice. The integration was analyzed using automated DNA sequencing by Sanger’s methods.

I-SceI cDNA sequence was cloned into a bicistronic expression cassette together with the GFP cDNA separated by a T2A linker sequence under the control of a CMV promoter. After transfection of the expression plasmid into the HEK293−mORF15 cell line, GFP positive cells were sorted 72 hours later by Fluorescent Activated Cell Sorting (FACS) and DSB repair events were analyzed using the T7 SURVEYOR assay (NHEJ events).

**Targeting of the I-SceI site in vivo.** The expression cassette CMV.I-SceI-T2A-GFP (the same as for the in vitro experiment) was cloned into an SSV9 vector. AAV2/5.CMV.I-SceI-T2A-GFP was produced in the Vector Core at the University Hospital of Nantes (http://www.vectors.nantes.inserm.fr) and the titer was determined by dot blot and expressed as vector genomes (vg)/mL. Viral preparation was manufactured using the HEK293 cell transfection method and purified by ultracentrifugation on iodixanol gradient. Iodixanol gradient was exchanged by PBS and the titer was determined by quantitative PCR.

For transfer of AAV vectors into the subretinal space, 2-month-old mice (B6J.Sv129-Rpgr<sup>tm1Sti</sup>) were anesthetized and pupils dilated as for the ERG measurements. A syringe (Hamilton, Reno, NV) with a blunt, GA34 12 mm needle (# 207434, Hamilton Company) was inserted tangentially through the conjunctiva and sclera and placed under visual control in the nasal half of the retina into the subretinal space (ie, between the retina and retinal pigment epithelium). One micro-liter vector suspension was injected to produce a bulous retinal detachment. Correct bleb formation was verified by OCT imaging immediately following the procedure.

Two months after injection, GFP expression was verified in vivo using the MICRON III camera (Phoenix Research lab). Animals were subsequently euthanized and the retina was dissociated. Dissociation was performed using the Papain-kit (Worthington Biochemical Corp) according to the protocol of the manufacturer with slight modifications. Briefly, retinae were kept for 90 minutes in Hanks balanced salt solution buffer at 37°C before being digested with Papain for 45 minutes at 37°C under constant shaking. DNase digestion was performed in Earlé’s balanced salt solution together with Ocomucoid and bovine serum albumin, and cells resuspended after centrifugation in PBS.

Dissociated cells were sorted by FACS according to their GFP expression profile. Cells from the noninjected retinae were also FACS sorted according to their size and granulation (photoreceptor enrichment). DSB repair events were analyzed using the T7 SURVEYOR assay (NHEJ events).

**RESULTS**

**Generation of the B6J.Sv129-Rpgr<sup>tm1Sti</sup> mouse model.** For the generation of the mouse model, a targeting vector containing a pathological point mutation (c.2793delA) was used for homologous recombination into the Rpgr locus of murine ES cells (C57BL/6/129sv hybrid cells) (Fig 1, A, mutation 2). To obtain an extended altered amino acid chain caused by the frame-shift mutation, the conditioned formation of a premature stop codon was prevented by introduction of an additional single base substitution (c.3071subT-A) (Fig 1, A, mutations 3). Furthermore, an XbaI restriction site (silent mutation, no change of encoded amino acid) was introduced for later screening purposes (c.2650subT-C) (Fig 1, A, mutations 1). Since the newly generated mouse model was created to be used for the development of therapeutic strategies based on genome editing, a recognition site for the I-SceI homing-endonuclease was integrated into the genome downstream of the Orf15 stop codon in the mutant sequence (Fig 1, A, mutation 4).
Successfully recombined stem cells were implanted into surrogate BALB/c mother mice and offspring were screened for chimeric events and sequence changes on the X chromosome. F1 offspring carrying the targeted mutations on the X chromosome was crossed with Cre-deleter mice to remove the neomycine selection cassette and further breeding was performed into the C57BL/6J background. Sequence changes were verified at the genomic level through Sanger sequencing during the breeding process (Fig 1, B). The entire sequence of the putative mutant full length cDNA is provided in the Supplementary file 1.

The resulting mutant protein is considered to contain RK repeats instead of EG repeats and is slightly shorter compared to the wild type protein (Fig 1, C). The amino acid sequence of the putative full length RPGR ORF15 isoform is shown in Supplementary file 2.

Expression of the mutant allele and protein in the mouse model. Expression of the targeted Rpgr ORF15 allele was verified on RNA and protein level. Messenger RNA (mRNA) from neuroretine of mutant and wild type animals was used as a template for cDNA synthesis via reverse transcription. A PCR product of 500bp in size was generated from the region containing the pathological mutation (2793delA) (Fig 2, A), and was sequenced to verify the alteration of the sequence (Fig 2, B). Translation of mRNA into protein was verified on whole retina extracts from animals at the age of 3 months using an antibody that recognizes an epitope in the C-terminal part of the ORF15 exon, thus allowing to recognize the RPGR ORF15 isoform specifically (Fig 2, C). As a control, protein from age matched Rpgr wild type retina was used. Bands with sizes below and above 150 kD were detected in the wild type retina, indicating the presence of several ORF15 variants with alternative amounts of the ORF15 acidic domain. In contrast, in the mutant retina, the total amount of RPGR protein was lower, and some of the isoforms appeared to be absent from the blot. The largest band, widely considered to be the RPGR-ORF15 full length band was visible in both samples, with the size in the mutant lower, and some of the isoforms appeared to be absent from the blot.

Expression and localization of the different RPGR isoforms was subsequently analyzed on retinal frozen sections of 3-month-old animals (Fig 3). First, we employed the same ORF15 specific antibody as was used in the Western blot together with an antibody staining acetylated tubulin, which marks the CC (Fig 3, A and B). The difference in immunolabeled RPGR-ORF15 protein between wild type and mutant retina was obvious. In the wild type retina, RPGR ORF15 immunoreactivity (green color) was of rod-like shape with brighter small spots (Fig 3, A, zoomed inserts). It was visible at the OS-axoneme side of almost every single Actub immunoreactive CC (red color). At the distal end of the immunolabeled CC, a small area of co-localization of RPGR ORF15 and Actub was visible (orange color). Similarly, albeit more spot-like, staining of RPGR ORF15 was detectable at the OS-axoneme side of the Actub immunoreactive CC in the mutant retina, the majority of it being correctly localized to the CC but with reduced immunoreactivity (zoomed inserts). In addition, exceptionally long rod-like accumulations of immuno-labeled RPGR ORF15 have been observed on the distal side of the CC, more frequently in the mutant than wild type retina. Since background signal was observed in both wild type and mutant outer retina, mislocalization of RPGR ORF15 was not assumed.

To further characterize the expression and localization of both isoforms, we also used S1 and S3 antibodies that have been widely used to stain RPGR proteins in the retina. The S1 antibody, which recognizes both the RPGR ORF15 as well as the RPGR ORF15 variant, detected RPGR protein in both wild type and mutant animals at the IS-side of the CC co-localizing with the Actub antibody (Fig 3, C and D). The reduced RPGR ORF15 immunoreactivity in mutant animals was due to the more spot-like appearance (Fig 3, D).

Staining with the antibody S3, which recognizes only the RPGR ORF15 variant, revealed clear immunolabeling in the wild type and mutant retina co-localizing with the Actub antibody at the IS-side of the CC (Fig 3, E and F). In the mutant retina RPGR ORF15 staining had a bright and rod-like appearance and was visible in every Actub immunoreactive CC. In comparison to the wild type retina, it appeared to be slightly more intense. The spot-like appearance of RPGR ORF15 immunoreactivity in the wild type retina resulted in a less prominent fluorescence than RPGR ORF15 immunolabeling (Fig 3, C).

In summary, these data demonstrate the expression and translation of mutant RPGR ORF15 mRNA. The RPGR ORF15 isoform is unaltered in its location but seems to be potentially upregulated when detected by
the RPGR\textsuperscript{const} antibody. The 2 RPGR\textsuperscript{ORF15} antibodies applied in this study to detect the RPGRORF15 protein revealed different subciliary localizations in both the wild type and mutant retina. While this might be due to technical reasons with the postfixation technique, it can also be associated with different epitopes recognized in different RPGR\textsuperscript{ORF15} isoforms as seen in our immune blot results (Fig 2, C). Further studies are ongoing to shed more light on this observation.

Significant reduction of retinal layer thickness starting at 9 months. Initially, the retina was analyzed at young age (3—6 months) and old age (15—18 months) for changes to the gross retinal morphology (Fig 4, A—D). In young mutant retinae, the gross morphology was similar to healthy retinae, with the exception that the outer segments looked slightly disorganized (Fig 4, A and B). Older mutant retinae displayed significant structural changes compared to healthy retinae, including a loss of column like organization of the nuclei, and outer segment destruction (Fig 4, C and D). No morphologic changes in the inner nuclear layers (ganglion cell layer) were observed in mutant animals (data not shown). Quantification of photoreceptor cell nuclei showed a slight loss of nuclei at early ages (3—6 months) in mutant retinae, which increased over time and aggravates at late ages (18—21 months), where only about 50% of nuclei were left (Fig 4, E).

In order to monitor photoreceptor degeneration over the entire lifetime of the animals, we performed in vivo imaging of the retina using OCT (Fig 5). Retinae were examined starting at 3 months up to 24 months of age every 3 months in the center and midperiphery (5 disk diameter distance; Fig 5, A and B). From the earliest time point onwards (even at 1 month, data not shown), the outer retinal bands, which can be nicely seen in wild type retinae, were not
Fig 4. Analysis of the retinal morphology. Semithin sections reveal morphologic alterations in the retinae of mutant mice. (A, B) Retina of a wild type animal at 3 and 18 months of age. (C, D) Retina of mutant animals at 3 and 18 months of age. (E) For the quantification of photoreceptor nuclei the number of nuclei in a section of 100 μm was counted at defined distances from the optic nerve head (n = 2 animals per group). Standard deviations are shown. INL: inner nuclear layer; IS: inner segments; ONL: outer nuclear layer; OPL: outer plexiform layer; OS: outer segments; RPE: retinal pigmented epithelium.
**Fig 5.** *In vivo* imaging of the retina using optical coherence tomography (OCT). (A) Retina of wild type and mutant animals were examined in the central or peripheral part (5 disk diameter distance) of the retina. The white brackets in the B scans indicate the outer retinal bands region. The green or black line in the fundus photographs highlights the area where the OCT scans was obtained. Scale bar: 50 μm. (B) Timeline of OCTs from the peripheral scans from wild type animals (3 and 15 months) and from mut at 3, 6, 9, 12, 15, 18, 21, and 24 months. (C) Entire retina and ONL thickness in wild type and mutant animals were quantified between 3 and 24 months of age (n = 4 animals per group). CH: choroid; ELM: external limiting membrane; GCL: ganglion cell layer; INL: inner nuclear layer; IPL: inner plexiform layer; IS: inner segments; ISe: inner segments ellipsoid;
discernable in the mutant retina (Fig 5, A, white brackets in OCT scans). These findings confirm the observation of outer segment disorganization in mutant animals as seen in Fig 4, B and represent an early biomarker for morphologic changes in the retina.

The thickness measurements of the entire retina as well as ONL alone show a progressive thinning of the layers, which becomes significant at the age of 9 months (Fig 5, B and C). Both, entire retina and ONL thickness in the center as well as the periphery are initially slightly thinner in mutant retina with a first decline at 9 month followed by a plateau phase until 15 months. Beyond that age, entire retina and ONL thickness rapidly decline until only minimal ONL thickness is left at 24 months of age and the entire retina thickness is only 50% of normal.

Significant reduction of functional changes starting at 12 months. Retinal function was analyzed using standard ERG recordings at different ages between 1 and 21 months (Fig 6). In the dark-adapted ERG, responses show progressing decrease of the a-wave amplitude, starting to be significant at the age of 12 months (Fig 6, A, B, and H). At 21 months, only about 35% of normal a-wave amplitudes were reached in mutant animals. B-wave amplitudes also decrease progressively starting at 12 months of age (Fig 6, C). The b/a-wave ratio remained unchanged at about 2, indicating a progressive degeneration of photoreceptors and second order neurons over time (data not shown). Albeit the amplitudes were decreased, characteristic waveforms remained in a-waves as well as in b-waves including oscillatory potentials (Fig 6, A and B). Interestingly, a-wave latency was increased in older mutant animals compared to age matched control mice (data not shown).

Under photopic conditions, b-wave amplitude was also slightly, but not significantly decreased in mutant animals (Fig 6, D and E). Photopic flicker responses were severely reduced in older animals (18–21 months) but not in young animals (3–6 months) (Fig 6, F and G).

DNA repair at the I-SceI locus in vitro and in vivo. In order to perform a proof of principle experiment demonstrating the possibility to edit the genome at the I-SceI locus, we employed a HEK293 cell line containing the C-terminal part of the murine ORF15 exon (Fig 7, A). A vector cassette containing the I-SceI and GFP cDNA linked by a T2A peptide under the control of a CMV promoter was used (Fig 7, B). Upon transfection, we subsequently enriched the transfected cell population to be analyzed by cell sorting of GFP positive cells (FACS). Upon enrichment, the surveyor assay (T7E1) was performed to visualize NHEJ events, which were at relatively high levels in the enriched cell population (Fig 7, C). Since nonenriched samples lower levels of NHEJ events, this method to isolate endonuclease expressing cells is important to obtain larger number of DNA editing events.

Upon successful proof of concept in in vitro experiments, we further evaluated the possibility to use I-SceI mediated genome editing at the ORF15 locus in vivo. One microliter each of AAV vectors (titer = 3 × 10^{11} vg/mL) containing the I-SceI-T2A-GFP expression cassette either under the control of the ubiquitous CMV promoter (AAV2/5) or the photoreceptor specific RK promoter (AAV2/8) was subretinally administered into the Rpgr<sup> EDIT</sup> mouse retinae at 8 weeks of age (Fig 8, A). Two months later, GFP expression was verified by in vivo fluorescent imaging (Fig 8, B), and retinal cells were dissociated and FACS enriched for GFP expression (Fig 8, C). Nonjected eyes were sorted for size and granulation. The enriched cells were subjected to the surveyor assay (T7E1) showing evidence of NHEJ activity at the target locus only in the injected eyes. Control vectors containing exclusively the GFP cDNA under the control of a CMV promoter were also injected into control animals, and did not show any sign of NHEJ events in the surveyor assay (data not shown).

**DISCUSSION**

In this study, we describe a new mouse model for human XLRP, which contains a single pathologic point mutation in the Rpgr-Orf15, a genetic alteration found frequently in human patients. We observed expression and limited translation of the mutant Rpgr-Orf15 transcript, and a slowly progressive degeneration starting at the photoreceptor level at about 9 months of age, both morphologically and functionally. Furthermore, we show for the first time in vivo genome editing at the Rpgr-Orf15 locus by using the I-SceI endonuclease.

The uniqueness of the presented mouse model lies in the presence of an I-SceI site close to the Rpgr gene locus, which can be used in subsequent experiments to study the efficacy of genome editing approaches based on homing endonucleases (HE) or other systems. Since the I-SceI endonuclease has a longstanding history in showing high efficacy and low toxicity in eukaryotic cells, this nuclease can now be used as a reference standard for studies using the current state of the art RNA based endonuclease systems (ie, CRISPR-Cas). Recent developments in the research for HE point even toward programmable variants, making our model even more relevant for future studies. Furthermore, due to its small size, HE including I-SceI is the single nuclease class that can be packaged into one single AAV vector together with the template DNA, which allows us to study the importance of using a 1-vector system compared to a 2-vector system, a question that is currently unanswered due to the lack of useful models and nucleases. In vivo studies addressing this question are currently ongoing.

Even though the Rpgr-Orf15 sequence is optimal for genetic modifications due to its highly repetitive nature, replacement of the entire exon by homology-directed repair or microhomology-mediated end-joining could be envisaged as one treatment approach for a large number of patients with different mutations within that exon. Therefore, targeting this exon by HE or other systems in a relevant animal model is of high interest to the scientific community.

In our mouse model, we observed protein translation from the mutant Rpgr<sup> EDIT</sup> transcript. This is in slight contrast to the situation observed in the rd9 mouse model, which contains a 32 bp deletion in the ORF15

ONL: outer nuclear layer; OPL: outer plexiform layer; OS: outer segments; RPE: retinal pigmented epithelium; RNFL: retinal nerve fiber layer; * P < 0.05. (Color version of figure is available online.)
region leading to a frame shift similar to our model and which shows only residual levels of the mutant protein in western blot and immunofluorescence assays. Furthermore, normal amounts of the RPGR<sup>const</sup> isoform were produced and are correctly localized in the CC in our mouse model, while no S3 antibody staining was detected in the rd9 mouse. A possible reason for this difference could be a differential regulation of the
RPGR^const isoform in both disease models, a mechanism already seen in selective KO models of either Rpgr^const or Rpgr^Orf15. Further studies to decipher this potential mechanism are currently ongoing.

We observed thinning of the ONL and reduction of photoreceptor nuclei at later stages of the disease. In the context of a therapeutic setting, such late reductions of photoreceptor cells and function may not be advantageous, since readout parameter following treatment would become significant only at late stages. On the other hand, this long-lasting presence of photoreceptors generates potentially a long therapeutic window. More importantly, the early alterations to the photoreceptor morphology, that is, the absence of the outer retinal bands on OCT scans early in the disease process, represent a useful readout parameter to study treatment outcome at an early age.

ONL thinning has already been shown for other Rpgr mouse models, for example, in the rd9 mutant mice becoming clear and significant around 12 months.
The Rpgr knock-out mouse model developed by Hong et al.\(^\text{23}\) also showed a thinning of the photoreceptor nuclear layer with a loss of 2 rows of nuclei on average by the age of 6 months, an observation that was also reported by other groups.\(^\text{28,29}\) These findings suggest that frame shift mutations in the Orf15 (like the ones occurring in the rd9 model and the model presented here) lead to milder forms of degeneration compared to knock-out mutations, which may be related to a residual function of the altered protein.

Concerning the function of the photoreceptors, ERG amplitudes in our model are reduced in scotopic a-wave as well as b-wave, starting at the age of 12 months, coinciding with the start of photoreceptor nuclear cell loss. Photopic flicker responses are similarly reduced in older animals. These observations indicate that both types of photoreceptors, rods and cones are involved in the pathology. The impact of aging on function is relatively small, since age-matched wild type animals only show a slight reduction of the measured amplitudes. Similarly, functional analysis showed reduced amplitudes for all known Rpgr mouse models at later stages of the disease.\(^\text{25,23,28,30,31}\) Similar to our model, the rd9 model shows a moderate but constant reduction of a-wave as well as b-wave amplitudes until the age of 24 months.\(^\text{25}\) The knock-out mouse model by Hong et al. shows reduced a-wave amplitudes for rods as well as cones at the age of 6 months,\(^\text{23}\) while the model established by Brunner et al. shows only a mild reduction of the scotopic a-wave from the age of 3 months onwards and no alteration of the scotopic b-wave.\(^\text{28}\)

The pathologies observed in our model or in the published rd9 mouse are less severe than those observed in human patients. One base pair frame shifts with long C-terminal protein chain alteration seem to be reason
for the most severe phenotypes in humans, but only a mild phenotype in mice (this study). Recent in vivo analysis of human retinas with SD OCT technology revealed early ONL thinning outside the fovea despite the presence of identifiable ISe (inner segment retinoid) and OS structures, which was also observed in other studies. In contrast, the murine ISe and OS structures can be distinguished in wild type mice, not visible in mutant retinae. This has also been observed in the knock-out mouse generated by Huang et al. Since current understanding of functional correlates identifies ISe as a valid morphologic indicator of functional photoreceptors in OCT scans of humans, this is not the case for murine OCT scans. In young affected animals of our model, ERG recordings are unchanged, yet the ISe band is not detectable from the beginning.

In summary, the newly generated mouse model displays a degenerative phenotype with thinning of retinal layers starting at 9 months and loss of photoreceptor function beginning at 12 months of age, indicating the activation of a similar or at least closely related pathologic pathway like in human patients. Therefore, this model is an eligible model for the development of gene-therapeutic applications in the future. In addition, it may help to gain further insight into the pathological mechanisms involved in retinal degeneration and the expression pattern of mutant Rpgr-Orf15. Moreover, the influence of point mutations in the ORF15 repetitive region on expression and splicing of the mRNA, as well as the biochemical reason for the toxicity of emerging proteins could be addressed by the model in further experiments. Finally, the use of this model to study genome editing in the degenerating retina in vivo using all nucleases classes currently available renders the model very important.


