Early disruption of photoreceptor cell architecture and loss of vision in a humanized pig model of usher syndromes

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Abstract

Usher syndrome (USH) is the most common form of monogenic deaf-blindness. Loss of vision is untreatable and there are no suitable animal models for testing therapeutic strategies of the ocular constituent of USH, so far. By introducing a human mutation into the harmonizing USH1C gene in pigs, we generated the first translational animal model for USH type 1 with characteristic hearing defect, vestibular dysfunction, and visual impairment. Changes in photoreceptor architecture, quantitative motion analysis, and electroretinography were characteristics of the reduced retinal virtue in USH1C pigs. Fibroblasts from USH1C pigs or USH1C patients showed significantly elongated primary cilia, confirming USH as a true and general ciliopathy. Primary cells also proved their capacity for assessing the therapeutic potential of CRISPR/Cas-mediated gene repair or gene therapy in vitro. AAV-based delivery of harmonin into the eye of USH1C pigs indicated therapeutic efficacy in vivo.

Keywords gene therapy; impaired vision; photoreceptor morphology; pig model; Usher syndrome

Subject Categories Development; Genetics, Gene Therapy & Genetic Disease

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Introduction

Usher syndrome (USH) is the most common form of inherited deaf-blindness in humans (Mathur & Yang, 2015). USH is clinically and genetically heterogeneous, with at least 10 genes assigned to three clinical USH types (Moller et al., 1989; Toms et al., 2020). The most severe of them is USH1, characterized by profound hearing loss from birth on, vestibular areflexia, and pre-pubertal onset of retinitis pigmentosa (RP). In patients, the congenital sensorineural hearing impairment can be compensated with cochlear implants, but no therapeutic option is presently available for the ocular disease constituent. Thus, most USH cases lead to severe visual impairment and blindness over time, adding a substantial psychological component to the clinical symptoms (Lonborg-Moller et al., 2020).

Development of therapies and understanding of USH pathogenesis in the retina is arduous as existing rodent models for USH reflect the human deficits in the inner ear, but show only a very mild, if any, retinal phenotype (Lentz et al., 2007; Williams, 2008; El-Amraoui & Petit, 2014). This has been attributed to substantial differences between mouse and human in anatomy and cellular composition of the retina and stimulated discussion about alternative model species. Pigs reflect structure and function of the human eye much better than mice as they have a similar size, a comparable rod/cone ratio and a cone-rich region, the so-called visual streak (Chandler et al., 1999; Hendrickson & Hicks, 2002). Specifically, calyceal processes (CP), cellular extravaginations at the transition of the inner segment (IS) to the outer segment (OS) of photoreceptor cells (PRC) appear in many species, including human and pig, but not rodents (Wolfrum et al., 2010; Sahly et al., 2012; El-Amraoui & Petit, 2014). Various pig models have been described for retinal degeneration, with the majority of them induced by compounds or light (Li et al., 2009; Wang et al., 2011; Monés et al., 2016) or additive transgenesis (Petters et al., 1997; Sommer et al., 2011; Ross et al., 2012; Kostic et al., 2013). As the latter do not truly reflect the genetic constellation in patients, the phenotype was, however, difficult to interpret and limited the utility of these models.

We therefore hypothesized that introducing a patient-specific mutation into a partially humanized USH1C gene would facilitate a better genotype–phenotype correlation of an USH1C pig model. USH1C comprises 28 exons with alternative splicing giving rise to numerous transcript variants (Zwaenepoel et al., 2001; preprint: Nagel-Wolfrum et al., 2021). According to their domain composition, the transcripts are classified into the three protein subgroups harmonin_a, b, and c (Verpy et al., 2000; Sahly et al., 2012). Most characteristic for harmonin are several PDZ domains (Kim and Sheng, 2004). N-harm and CC (coiled-coil) domains are further integral constituents of all harmonin variants, while only harmonin_b comprises a second CC as well as a PST (proline-serine-threonine rich) domain (Verpy et al., 2000; Boeda et al., 2002; preprint: Nagel-Wolfrum et al., 2021). A gene therapy approach in the inner ear of a murine Ush1c model demonstrated the physiological relevance of distinct isoforms: harmonin_b1 appeared sufficient for restoring the vestibular system while only harmonin_b1 corrected deficits in the auditory function (Pan et al., 2017). The specific relevance of harmonin_b variants for hearing is also illustrated by mutations in the CC2 or the PST domain, causing autosomal recessive deafness (DFNB18A) rather than USH in human patients (Ouyang et al., 2002) or in a naturally occurring deaf-circler mouse line (Johnson et al., 2003). In contrast, USH-causing mutations are located in the common N-harm, PDZ, or CC1 domains (Fuster-Garcia et al., 2021).

The components of the USH1 and USH2 interactome are the main, but not exclusive interaction partners of harmonin (Reiners & Wolfrum, 2006; Mathur & Yang, 2015). The mutual interplay of USH proteins and the molecular properties of harmonin, however, remain elusive. In the inner ear, harmonin serves in the formation and functionality of the interstereociliary links in the bundles protruding from cochlear hair cells (El-Amraoui & Petit, 2005; Richardson et al., 2011). Harmonin is also consistently localized to the ribbon synapses of hair cells and PRCs (Reiners et al., 2005; Williams et al., 2009a; Gregory et al., 2011), whereas data on harmonin abundance in other compartments of the retina are contradictory in rodents (Reiners et al., 2003, 2005; Williams et al., 2009a; Sahly et al., 2012). In species with pronounced CP structures in PRC such as amphibians or primates, however, USH proteins, including harmonin, seem to co-localize with CP or, at least, appear in the transition zone of IS to OS (Sahly et al., 2012; Schieteroma et al., 2017). Very recently, we detected harmonin also in the OS of rods, in Müller glia cells (MGC), and in the outer limiting membrane of the human retina at the adhesive junctions between MGC and PRC (preprint: Nagel-Wolfrum et al., 2021).

Based on these considerations, we designed and generated an USH1C pig model, carrying the deleterious c.C91T/p.R31X non-sense mutation after humanization of exon 2 and its surrounding intronic sequences in the porcine USH1C gene. For the integration of exon 2 in the major splice variants USH1C_a, b, and c, the chosen mutation will affect all relevant USH1C transcripts and its disruption was, therefore, proposed to cause the full spectrum of USH. USH1C pigs revealed profound hearing impairment, pronounced vestibular dysfunction, and significant visual deficits. At the cellular level, the retinal phenotype was characterized by an early disruption of OS photosensitive disc architecture, elongated connecting cilia, and narrower cone synapses. Primary cells from USH1C pigs were sufficient for in vitro testing of human-specific gene repair or gene therapy. The latter approach did also restore retinal function in an in vivo pilot study.

Results

Genetic modification of the porcine genome by introducing a human segment carrying a patient-specific mutation into the USH1C gene

First, we assessed the translational potential of an USH pig model by examining the harmonin proteins and the underlying genetic USH1C structure across mammalian species. With the exception of a few small segments in the coding regions, harmonin shows an outstandingly high degree of conservation between the species (Fig 1A). Extended CCCTC-binding protein (CTCF)/cohesin-binding regions suggest that exons undergoing complex splicing are controlled by distantly located chromosomal segments in cis (Holwerda & de Laat, 2013; Ghirlando & Felsenfeld, 2016) rather than by local regulatory elements (Fig 1B). In contrast, a 1.5 kb segment comprising exon 2 and its surrounding intronic regions were predicted to be relevant in gene regulation (Fig 1C). Conservation is noticeable throughout the entire region, culminating in a 100% identity among all amino acid positions encoded by exon 2.
in all examined species (Fig 1D). We therefore modified the porcine USH1C gene by a corresponding 1.5 kb element including a disruptive c.C91T/p.R31X nonsense mutation (Zwaenepoel et al, 2001) as well as 3 intronic single nucleotide polymorphisms (SNPs) identified on the disease allele of a human USH1C patient. A modification vector was produced by bacterial recombineering out of a bacterial artificial chromosome (BAC) covering the porcine USH1C locus and combined with CRISPR/Cas components to introduce the desired modification in pig primary cells (Fig EV1A–C and Appendix Fig S1A–D).

Following previous attempts to generate pig models (Kurome et al, 2015; Vochozkova et al, 2019), single-cell clones (SSCs) were generated from primary cells, nucleofected with the modification components, and screened for the desired modification (Fig EV1E and Appendix Fig S1E and F). Somatic cell nuclear transfer (SCNT) was used for establishing founder animals from verified genetically modified SSCs (Fig EV1D). Female cell clones with both USH1C alleles modified were further lipofected with Cre-mRNA in order to excise the selection cassette prior to their usage in SCNT experiments (Fig EV1F). Out of 12 embryo transfers, 4 litters were produced, delivering a total of 18 piglets. Genetic analysis confirmed excision of the neo cassette in 5 out of 15 founder animals, constituting a remnant single lox site in intron 1 (Fig EV1G and H), the humanized fragment (Fig EV1I) as well as the correct conjunction

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Figure 1. Generation of USH131X pigs.
A Three main splice variants _a_, _b_3, and _c_ have been confirmed for human USH1C. According to an Entropy plot (upper panel), the coding regions appear highly conserved among mammalian species with some variability in inter-domain segments, but also within the _b_3-specific PST domain.
B Homology: A high degree of conservation occurs at the nucleotide level, including exons but also intronic elements. Regulation: High degrees of conservation often correlate with strong indication for transcriptional regulation properties. Thickness and color intensity are according to matches to 6 prediction tools (DNase sensitivity, ChIP-Chip, FANTOM, GeneHancer, FAIRE, PreMod). Coverage: Most of the USH1C gene and its surrounding regions are consistently abundant in the examined species (grey). Gaps (white) appear in all species, but mostly remain small (< 500 bp). Few intronic areas are consistently affected by the introduction of repetitive elements (triangles, with approximate length in kb). Rodents show a high evolutionary turnover, with large gaps or inserts > 1 kb in extended introns and adjacent intergenic regions of USH1C. Significant proportions of USH1C are indicated as CTCF/cohesion-binding regions that appear relevant for chromosomal interactions (orange blocks).
C Exon 2 and its surrounding intronic sequences constitute a consistent block in the alignment with high degree of sequence conservation, strong evidence for regulatory purposes and a lack of repetitive element insertion. For this reason, the entire segment was humanized (magenta), including the causative c.C91T/p.R31X mutation as well as three patient-specific SNPs (pins).
D All examined species show identical amino acid sequence on the protein level and high degrees of conservation at the intronic borders on the nucleotide level. The yellow box indicates R31, the amino acid that is mutated in R31X patients.
Source data are available online for this figure.
of the modification to the porcine sequence at the 3'-end of the modification in the animals (Appendix Fig S1G).

Strikingly, all cloned founder animals showed a pronounced circling phenotype at birth (Movie EV1), correlating with a dysfunctional vestibular system (Ernest et al, 2000; Johnson et al, 2003). For this reason, animals were raised in a rescue deck from birth on (Egerer et al, 2018). After 48–72 h of interval feeding by hand, circling essentially ceased and USH1C piglets were able to feed themselves from the nurturing unit. We interpret this as central vestibular compensation and re-weighting of multisensory input to the balance system.

In general, USH1C pigs developed and acted normally, but stress-induced circling proved a consistent hallmark in the case animals found themselves challenged by new situations. At an age of 3–6 months, we occasionally observed a spontaneous nystagmus of the eyes (Movie EV2) in some animals. This is very likely caused by an asymmetrical tonic firing of vestibular afferents from the right and left labyrinth (Baloh et al, 2012; Dougherty et al, 2020) and has been anecdotally described in USH patients (Moller et al, 1989; Bonneau et al, 1993). 8-month-old USH1C pigs were fertile and 3 sows became pregnant after insemination with wild-type (WT) sperm and gave birth to F1 litters, comprising heterozygous offspring (Appendix Fig S2A). F2 USH1C piglets were produced by mating F1 animals with each other or by inseminating of the cloned founder sows with sperm from one of their heterozygous offspring. In all USH1C offspring, a circling phenotype appeared as it had been observed in the F0 generation produced by SCNT. During breeding, it became obvious that some of the founder animals carried two distinct mutations of the USH1C alleles. One constituted the c.C91T/p.R31X mutation, while the other was a large genomic disruption, deleting exon 2 of the USH1C gene (Appendix Fig S2B). Genomic evaluation by endpoint PCR and qPCR and transcriptional analysis of heterozygous animals confirmed this hypothesis (Appendix Fig S2C and D and Appendix Fig S3). Details on animals used for experimental and breeding purposes are shown in Appendix Table S1.

Motion analysis demonstrates vestibular dysfunction and visual deficits in USH1C pigs

To quantify changes in visually guided behavior, animals were challenged in distinct settings. In a barrier course in which animals had to bypass vertical shields to reach a food bowl (Barone et al, 2018), USH1C pigs did not show obvious differences to WT controls regarding duration or gait instability (Movies EV3, EV4 and Fig 2A and Appendix Fig S4). Interestingly, however, USH1C animals hesitated to enter the course in the dark (average 2.9 lux), turned around more often within the course, and had more frontal contacts with obstacles than WT control animals while these differences were less pronounced under light conditions. A more complex obstacle course was carried out on a cohort of 5 USH1C and 5 age-matched WT controls. Over an age of 7–12 months, animals had to repeatedly walk through a parcour in which they had to cross or bypass distinctly shaped hurdles in varying constellations (Movies EV5, EV6 and Appendix Fig S5). In the dark (0.1–12 lux), USH1C pigs had significantly more contacts with hurdles, while their speed was similar to WT (Fig 2B and Appendix Fig S6). In the light (50–150 lux), USH1C pigs did not only touch hurdles more often, but were also significantly slower than WT controls. Plotting the data points for individual animals supported the interpretation of a general difference between USH1C pigs and WT control animals (Appendix Fig S7). Remarkably, USH1C pigs had difficulties to pass hurdles taking a pronounced vertical movement and therefore requiring balance (such as the red cavaletti) as well as obstacles challenging stereopsis and peripheral vision (such as the hanging or standing blue barrel and the step board) (Fig 2C). While the interpretation of an impaired vestibular system was further supported by the clearly different gait pattern in USH1C pigs (Movies EV5 and EV6), the interpretation of a reduced visual function in these animals was corroborated by a more detailed analysis of the movement in the obstacle course (Fig 2D). It became obvious that USH1C pig trajectories were longer (TrajLength and TrajDistance) compared to WT, with more frequently abrupt changes in direction (lower Emax). In addition, they took more time to complete the course (TrajDuration) while their step length was shorter.

Clinical examinations confirm hearing loss and vision impairment in USH1C pigs

Auditory brainstem response (ABR) documented severe hearing deficits in USH1C pigs as early as 3 weeks and up to 2 years of age, indicated by a lack of response even upon click stimulus at sound pressure levels of up to 120 dB sound pressure level (SPL).
The findings of sensorineural hearing loss (Cvejic et al. 2009) correlated with changes in the arrangement of hair cell ciliary bundles and the lack of stereocilia in inner ear hair cells (Fig EV2B), as well as with the findings in Ush1c mouse models (Johnson et al., 2003; Lefevre et al., 2008) and the congenital deafness in human patients (Koenekoop et al., 1993). The hearing threshold of heterozygous USH1C+/− pigs was slightly elevated compared to WT littermates at distinct ages (Figs 3A and EV2).
EV2E and F). Hearing thresholds for both WT and heterozygous animals slightly declined with increasing test frequency, while thresholds in USH1C animals deteriorated further for higher frequencies. In line with their profound sensorineural hearing loss, USH1C piglets did not react on their mother’s call for nourishing (Movie EV7). Most of them needed nudging for waking up, while heterozygous littermates arrived at their mother within 15 s (Fig 3A). Consistently, some USH1C piglets were within the faster group, presumably awoken from being sideswiped by their littermates.

Clinical examination of the eyes in USH1C pigs at 1 year of age revealed changes comparable to early stages of USH in human patients. A reduction of cone photoreceptor function was evident in the central visual streak area analyzed by multifocal electroretinography (ERG, Fig 3B and C). Both rod and cone photoreceptor cell (PRC) responses were markedly attenuated in USH1C pigs compared to age-matched wild-type animals at 12 months of age (Fig 3D and Appendix Fig S8). USH1C animals showed >70% reduction in the scotopic standard flash a-wave amplitude ([mean value ± SD] 21 ± 18 µV in USH1C vs. 79 ± 2 µV in WT animals) and a >50% reduction in scotopic b-wave amplitude (75 ± 32 µV in USH1C vs. 154 ± 39 µV in WT animals). Responses after photopic standard flash stimuli showed a 50% reduction of a-wave amplitude ([mean ± SD] 10 ± 0.1 µV in USH1C vs. 20 ± 5 µV in WT animals) and a 60% reduction in the respective b-wave amplitude (86 ± 49 µV in USH1C vs. 215 ± 50 µV in WT animals). Confocal scanning laser ophthalmoscopy (csLO) of the retina and choroid layer showed no differences between WT and USH1C pigs (Appendix Fig S9). Interestingly, spectral domain optical coherence tomography (OCT) (Fig 3E) revealed only minor differences in retinal architecture between WT and USH1C pigs with good preservation of outer retinal layers containing the PRCs, suggesting a large therapeutic window.

Lack of harmonin in sub-cellular structures disrupts photosensitive disc architecture

Previous studies of USH1C/harmonin expression, particularly in the mouse retina, were inconsistent but indicated localization of harmonin in PRC, MGC, and secondary retinal neurons. (Reiners et al, 2003; Reiners & Wolfrum, 2006; Williams et al, 2009b; Sahly et al, 2012). In WT pigs, we detected harmonin in the PRC layer, the outer limiting membrane (OLM) and outer plexiform layer (OPL) (Appendix Fig EV3). More specifically, the protein was found in the PRC outer segment (OS), the OS base, in calyceal processes (CP) and synaptic pedicles. Harmonin abundance in MGC microvilli tips and the cell adhesion region of the OLM is in line with expression profiles and subcellular localization of harmonin in the human retina (preprint: Nagel-Wolfrum et al, 2021). In both, USH1C and WT pigs, USH1C transcripts were detected in multiple but not all organs (Fig 4A and B). Sanger sequencing confirmed correct splicing of the humanized exon 2 in USH1C pigs (Appendix Fig S10A and B). While in some tissues (colon, duodenum), RT-PCR indicated considerable proportions of transcripts lacking exon 13 (Fig 4C), splicing out of exons 13 or 14 appeared only at very low levels in the retina (Appendix Fig S10C). USH1C_b splice variants were found at low level in general, but clearly indicated a lack of exon 27 in all transcripts and alternative splicing of exon 20 (Appendix Fig S10D and E). We did not detect transcripts without exon 11, a core characteristic of USH1C_c, indicating that USH1C_a1 is the dominant transcript of USH1C in the pig retina. At the protein level, harmonin was consistently lacking in Western blot analysis (Fig 4D) and in tissue sections (Fig 4E), confirming the sufficient suppression of harmonin translation by the premature R31X nonsense mutation. No changes were observed in protein levels of SANS (encoded by USH1G), whrin (encoded by WHRN (USH2D)), and myosin VIIa (encoded by MYO7A (USH1B)) in the retina (Fig 4F). In contrast, glial fibrillary acidic protein (GFAP) was considerably upregulated in the USH1C retina already at an age of 3 weeks (Fig 5A and B), suggesting gliosis, a phenomenon often seen in inherited retinal diseases (Hippert et al, 2015) or injury-mediated degradation processes (Bringmann et al, 2009).

While the general structure of the retina appeared grossly intact in USH1C pigs (Figs 4E and EV3G, and Appendix Fig S9), subcellular analysis revealed specific disruptions in the architecture of compartments containing harmonin. In line with its proposed scaffold function in ribbon synapses (Gregory et al, 2013), the loss of harmonin results in significantly reduced extension of the cone
synaptic pedicles. This was confirmed by both the quantitative determination of cone-specific peanut agglutinin (PNA) stretches at synapses (Fig 5C) and the direct measurement of synapse width in transmission-electron microscopy (TEM) (Fig 5D). We also found significant increases in length of the connecting cilium (CC) in rod PRCs, as determined by quantitative measurements of the ciliary marker GT335 stretches (Fig 6A–C) as well as by direct examination of TEM ultrathin sections (Fig 6D and E). Aiming at identifying a
consistent relevance of harmonin for cilia formation, we examined primary cilia in dermal fibroblast of USH1C and WT in pigs and humans (Fig 6F). While the number of ciliated cells in USH1C pigs was reduced compared to WT (Fig 6G), the length of the primary cilia was consistently elongated in fibroblasts isolated from USH1C pigs and a USH1C^{ΔSIX/RROBS} patient after induction of ciliogenesis (Fig 6G and H). Most striking, however, was the deprivation of rod OS architecture in PRC. While this is normally characterized by a strictly horizontal stacking of parallel photosensitive membrane discs (Fig 6I) (Stoerbrand, 1953), we detected vertically oriented membrane discs (Fig 6J) in animals as early as 3 weeks of age. This aberrant structuring became even more prominent at 12 months when the stacking of discs was further infringed by interstitial gaps and vesicle-like structures at the OS base (Fig 6K–M). In contrast to rods, the OS architecture in cones remained normal throughout the first year of life (Fig EV3G).

**USH1C pigs facilitate testing of therapeutic approaches in vivo and ex vivo**

Besides its usage in studying disease progression and its underlying mechanisms, the USH1C pig will also support the development of novel treatment approaches. The defined genetic cause of the model, the disruption of exon 2, affects all relevant USH1C splice forms and, therefore, facilitates the testing of a broad spectrum of treatment options. We addressed fundamental questions by examining the tropism of distinct serotypes of adeno-associated viruses (AAV) for transducing cells of the retina in vivo, by testing the gene therapy and gene repair approaches in vitro and by exploring the potential to restore eye function in vivo after sub-retinal injection of AAV expressing USH1C mRNA.

For testing the distribution of AAV vectors, defined amounts (200 µl, 2 × 10^{10} vg/eye) of established AAV8, AAV9, or a modified Anc80 vectors, each of them expressing egFP under control of a CMV promoter (Vandenberghhe et al., 2013; Carvalho et al., 2018), were injected into the sub-retinal space of the eye in WT pigs followed by examination of egFP expression after 5 weeks. While AAV8 transduced only PRCs efficiently, AAV9 and Anc80 transduced both, PRC and MGC (Fig 7A, lower panel). Anc80 was much less effective in transducing cells of the retinal pigment epithelium (RPE) than AAV9 and, in particular, than AAV 8 (Appendix Fig S11), indicating that the target specificity in the eye can be controlled by the appropriate choice of the AAV capsid. No relevant other off-target expression of egFP was noted for the tested AAVs.

The capability for correcting the causative USH1C^{ΔSIX} mutation in the humanized exon 2 by CRISPR/Cas-mediated gene repair was evaluated by applying distinct constellations of human-specific gRNA and repair oligo-nucleotides into pig primary kidney cells (PKCs), isolated from USH1C founder animals (Figs 7B and EV4). The efficacy of transforming the nonsense codon TGA into an Arginin-encoding CGA by homology-directed repair (HDR) reached 40%, determined in DNA preparations from mixed cell clones. Upon clonal selection, the analysis of 68 single cell clones (SSCs) confirmed that 40% of the clones underwent gene correction, while non-homologous end-joining (NHEJ) was seen in 31% of the SSCs. Occasionally, HDR was accompanied by additional mutations, either small insertions/deletions or single nucleotide exchanges. For the specific positioning of the gRNA, a considerable proportion of mutations that appear in intron 2 would not affect the correct transcription and translation of USH1C.

For exploring the potential of harmonic restoration by gene therapy, we transfected a vector encoding the human harmonin_a1 splice variant into primary skin fibroblasts derived from USH1C pigs. By using the number of ciliated cells as well as the length of the primary cilium as therapeutic readout (Fig 6G and H), we confirmed that treated fibroblasts differed significantly from untreated USH1C cells and reached the levels determined in WT fibroblasts for both parameters (Fig 7C).

The potential to compensate for the lack of harmonin in USH1C pigs was explored by applying a volume of 200 µl, 1 × 10^{11} vg Anc80.CMV.harmonin_a1 vector into the sub-retinal space of one eye while the other eye was sham-treated with PBS. In one 3.5-year-old USH1C pig, we detected harmonin in histological sections and in Western blot after 12 months in the treated eye (Fig 7D). Furthermore, amplitudes of the a- and b-waves were significantly increased in the scotopic and photopic ERG, compared to the sham-treated eye, similarly to another animal that was treated accordingly at an age of 3.5 years and examined after 3 months (Figs 7D and EV5).

**Discussion**

Combining hearing deficits, vestibular dysfunction, and vision loss, the USH1C pig is the first animal model to reflect the full syndromic phenotype in human USH1 patients. The early onset of symptoms...
Figure 4.
Figure 5. USH1C<sup>331X</sup> induce upregulation of GFAP and a synaptic phenotype in USH1C pig photoreceptor cells.

A, B Upregulation of GFAP in the retina of USH1C pigs at an age of 3 weeks (3 w). (A) Representative immunofluorescence staining of GFAP in Müller glia cells which extend throughout almost the entire retina from the OLM (arrow) to the ganglion cell layer (GCL) of the retina. The consistent increase of GFAP expression in the Müller glia of USH1C pigs indicates Müller cell activation and gliosis. IS, inner segment; ONL, outer nuclear layer; OPL, outer plexiform layer; INL, inner nuclear layer; IPL, inner plexiform layer; scale bar: 20 µm. (B) Left panel: Western blot analysis of GFAP protein expression in 3 w old USH1C piglets and age-matched controls. Anti-actin Western blot was used as loading control. Right panel: Quantification of Western blot bands in 4 gels by the LI-COR Odyssey system revealed a strong increase GFAP expression in 3w USH1C piglets when compared to age-matched WT controls (2 piglets, 3w, 1 retina each, 2 TRs, error bars represent SD of the mean, two-tailed Student’s t-tests, *P < 0.05).

C, D Reduced synaptic width in USH1C pigs at an age of 1 year (1 y). (C) Fluorescent microscopic analysis of the cone synapse phenotype. Left panel: Representative images of longitudinal sections through WT and USH1C pig retinae stained for the pre-synaptic marker PSD-95 (green) and by fluorescent peanut agglutinin (PNA, red) for cone synaptic pedicles (white arrows) and counter-stained by DAPI for nuclear DNA (blue). Scale bar 10 µm. Middle panel: Higher magnification of a PNA-stained cone synaptic pedicle. Synapse width was determined as the maximum extension of consistent PNA signals. Scale bar 1 µm. Right panel: Measuring cone synaptic pedicle width in WT and USH1C pigs by applying a Fiji script to PNA-stained sections indicated reduced synaptic width (2 pigs, 1 y, 1 retina each, number of examined synapses indicated, error bars represent SD, two-tailed Student’s t-tests, ***P < 0.001). (D) Determining cone synaptic pedicle width by TEM. Left panel: Representative images of retinal cross sections. Scale bar 500nm. Right panel: Quantification of synaptic width confirmed the significantly reduced width of cone synaptic pedicles in USH1C pigs. (1 pig, 1 y, 1 retina each, number of examined synapses indicated, error bars represent SD, two-tailed Student’s t-tests, *P < 0.05).
makes it suitable for studying the basic mechanisms of the disease as well as for evaluating strategies of therapeutic intervention. The persistence of a largely intact retinal architecture throughout the first year of life provides a wide therapeutic window and the potential to revert the pathogenesis by targeting dysfunctional, but still viable photoreceptor cells. The specific design of the genetic modification allows the examination of a broad diversity of therapies, including classical additive gene therapy, human-specific gene repair, and translational read-through or cell replacements. The establishment of breeding populations (Appendix Fig S2), the provision of distinct therapeutic options in vitro and the success of AAV-based delivery of harmonin in exploratory experiments in vivo (Figs 7, EV4, and EV5) collectively demonstrate the potential of the model for evaluation of therapeutic interventions. Consequently, USH1C pigs can now be used for systematic pre-clinical examination in vivo, similar to recent ventures in genetically engineered pig models (Langin et al, 2018; Regensburger et al, 2019; Moretti et al, 2020). In such studies, housing USH1C pigs under standardized conditions will minimize environmental confounding, whereas the pronounced genetic diversity of conventionally produced pigs needs consideration. While presenting a statistical challenge for preclinical proof-of-principle studies, this heterogeneity is, of course, much more likely to reflect the scenario in clinical trials and might thus lead to a more robust predictive power of the model in the translation of new therapeutic strategies (Ebermann et al, 2010).

The manifestation of an ocular phenotype in USH1C pigs is striking, taking into account the multiple efforts that have been made in creating such models in other species with so limited success (Williams, 2008). Our results reinforce some of the multiple sentiments that have been postulated as reasons for the lack of a vision phenotype in rodents. The onset of retinal dysfunction in pigs within the first year of life argues against the notion that the limited life span of mice prevents the developing of a similar phenotype in the mouse eye. While a potentially distinct function of the USH proteins in the retinae of different species cannot be readily excluded, the degree of sequence conservation of harmonin and its immediate interaction partners is remarkable (Fig 1 and Appendix Fig S12), insinuating a conserved and consistent nature of the USH1 interactome across species.

Crucially, porcine harmonin appears to be localized to CP (Fig EV3D) which is thought to stabilize the OS structure in the PRC of different species, although not in rodents (El-Amraoui & Petit, 2014). A recent publication demonstrated the absence of CP in cone PRCs and a pronounced outgrowth of disc membranes at the base of rod OS upon partial knock-down of the Ush1f-encoded protein pcdh15 in the clawed toad (Schiетromaa et al, 2017). While the complete lack of harmonin in pig did neither show outgrowth of disc membranes nor affect the formation of CP to a similar extent, it explicitly disturbed the photosensitive disc architecture in rods (Fig 6I–N). Considering the actin-binding properties of harmonin, the changes in disc architecture and CC dimensions (Fig 6A–E) link the retinal pathology in USH1C pigs to the proposed role of the actin cytoskeleton as driving force in disc neogenesis and in the structural maintenance of the PRC OS (Corral-Serrano et al, 2020; Spencer et al, 2020). Specifically, ciliogenesis has been shown altered upon dysregulation of the actin cytoskeleton at a molecular level (Kim et al, 2015), an effect that has been also associated with neurodegenerative diseases (Karam et al, 2015). The consistent elongation of CC in PRC and primary cilia in dermal fibroblasts (Figs 6 and 7C) further confirms the tight interaction of the ciliary actin system with axonemal microtubules in PRCs (Wolfbum & Schmitt, 2000; Corral-Serrano et al, 2020) and the general role of the actin system in the organization of ciliary axonemes (Kiesel et al, 2020). Although precise mechanistic insights into the role of harmonin in cilia function require further studies, our data provide clear evidence that USH is a true retinal ciliopathy in its fundamental aspects (Buja-kowska et al, 2017).

Remarkably, the significantly disturbed photosensitve disc architecture in rod OSs and the ciliary aberrations (Fig 6) are associated with a significant (c. 70%) reduction in rod-derived ERG responses (Fig 3D). The apparent decline in cone-derived ERG (c. 50%) and the reduced width of cone synaptic pedicles.

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**Figure 6. Changes in ciliary structures and outer segment architecture of USH1C photoreceptor cells.**

A–C Ciliary length measured by fluorescence microscopic analysis at 1 year of age (3y). (A) Representative immunostaining of the ciliary marker Gt335 (green) in longitudinal crosssections of PRC ciliary regions in WT and USH1C retinas. Counterstaining by DAPI, scale bar 10 μm. (B) Gt335 staining at higher magnification and principle of quantifying ciliary length by Gt335 staining. Scale bar 500nm. (C) Length of connecting cilia (CC) appear elongated in USH1C when applying a Fiji script to immunostained histological sections (2 pigs, 1y, 1 retina each, number of examined connecting cilia is indicated, error bars represent SD, two-tailed Student’s t-tests, ***P < 0.0001). **

D–E Measuring CC length in TEM sections. (D) Representative images of longitudinal cross sections and principle of determining CC length. Scale bar 500 nm. (E) Elongation of CC length is confirmed in TEM sections of USH1C PRC. (1 pig, 1y, 1 retina each, number of examined connecting cilia indicated, error bars represent SD, two-tailed Student’s t-tests, ***P < 0.0001). **

F–H Length of primary cilia in dermal fibroblasts. (F) Staining of ciliary structures in fibroblasts stained with ARL13B (ciliary shaft, green) and PCNT2 (ciliary base, red), DAPI (nucleus, blue). Scale bar upper panel 5 μm, lower panel 2.5 μm. (G) Left panel: The number of ciliated cells is decreased in porcine fibroblasts of USH1C pigs, compared to WT controls (6 TR, number of examined cells indicated, error bars represent SD, two-tailed Student’s t-tests, *P < 0.05). Right panel: The length of primary cilia in porcine fibroblasts is increased in USH1C (6 TR, number of cells examined indicated, Kolmogorov-Smirnov (KS) test, ***P < 0.0001). (H) Similarly, ciliary length of fibroblasts from a human USH1C patient is elongated, compared to a healthy control (7 TR, number of cells examined indicated, KS-test, ***P < 0.0001). **

I–M Outer segment (OS) PRC architecture by TEM of longitudinal retinal sections. (I) At 3w, WT show normal sub-cellular organization, including parallel stacking of photosensitive discs and calyceal processes (black arrows, representative image from 2 pigs, 2 retinas, scale bar 600 nm). (J) In 3w USH1C rods, membrane discs appear also in vertical orientation (white arrows, representative image from 2 pigs, 2 retinas, scale bar 750 nm). At 1 y, interstitial gaps appeared in disc stacks of the OS in USH1C pigs (K, arrow, scale bar 550 nm) and vesicles are found at the OS base (L, asterisk, scale bar 500 nm). (M) The disc architecture in 1 y animals was substantially distorted (arrows, representative image from 2 pigs, 2 retinas, scale bar 850 nm).

N Localization of cellular anomalies in an USH1C rod highlighted in orange.
Figure 6.
(Fig 5C and D) indicate the early onset of cone-specific alterations as well. Both, structural and clinical observations thus support the notion of a rod-cone dystrophy phenotype in USH1C pigs, reflecting the situation in human patients quite well (Koenekoop et al., 1993; Khateb et al., 2020). The early onset of PRC malfunction was in line with the up-regulation of GFAP in MGC (Fig 5A and B) and with reports of visual deficits in early childhood of USH patients (Stingl et al., 2019). Interestingly, the changes in subcellular morphology, the reduction in electrophysiological PRC function and the observed changes in visually guided behavior (Fig 2) come along with only very mild effects on overall retinal morphology (Fig 3E and Appendix Fig S9). It needs to be considered, however, that the retinal phenotype in USH1 patients varies widely, ranging from intact central retinal architecture and excellent visual acuity at 45 years to advanced central retinal disease and legal blindness by their early 30s (Lenassi et al., 2014). Such a phenotypic heterogeneity is observed in most if not all forms of inherited retinal diseases and is thought to arise from genetic or environmental confounding factors or a combination thereof (Veleri et al., 2015). The possibility to examine the USH1C pig model under standardized conditions will help to discern the effect of the mutation from other influencing factors.

In our work, we give also first indications for a potentially impaired hearing ability in heterozygous USH1C carriers (Figs 3A and EV2E and F). As the results were consistent at distinct ages, in different groups of animals and across the examined frequencies, we interpret this as a slight but significant biological effect. It must be considered that we are not aware of reports on impaired hearing in human USH1 carriers and so we cannot seriously estimate if these findings are of clinical relevance for human USH1 carriers. Taking into account the very low numbers of patients carrying distinct USH1C mutations, however, we would definitely exclude an evolutionary benefit of heterozygous USH1C defects, as it has been reasoned for the spreading of genes such as HBB (Piel et al., 2010) or CFTR (Gabriel et al., 1994; Pier et al., 1998) in the human population.

For the different aging processes in pig and human, a direct correlation of ages between the species is misleading. Instead, we propose to correlate the manifestation of fertility at 5–8 months in pigs, compared to 12–15 years in human beings, suggesting that a 12-month-old pig may fairly correspond to a biological age of 15–25 years in humans (http://www.age-converter.com/, https://unleashpetphotography.com/animal-age-calculator). Accordingly, the phenotype in USH1C pigs appears to reflect patients well within the published spectrum of disease severity and progression. As a consequence, we would also suppose that translating clinically established endpoints (as measured e.g., by ERG or OCT) from human to pig will support the assessment of future pre-clinical studies in our model. Vice versa, the multi-disciplinary examination of the model described here defined novel disease hallmarks, which may give rise to alternative endpoints for future clinical trials. The value of additional read-out parameters such as motion analysis (Fig 2) is corroborated by a pivotal study documenting the success of the voretigene neparvovec AAV-based drug in the multi luminance mobility test (MLMT) as primary endpoint (Russell et al., 2017).

The specific wavelength sensitivity of cones (Kostic & Arsenijevic, 2016) and the complex interaction of vision with other sensory organs required substantial effort in the investigation of visually guided behavior. Parcour tests were conducted under distinct light conditions, but calculated luminance levels in the pig eye suggested that the experiments took place at the lower and upper limit of mesoscopic sensing (Zele & Cao, 2014). Albeit this prevents a clear discrimination of rod- and cone-based vision in the motion analysis, essential conclusions can be drawn. First, the similarity of core parameters such as duration and snout contacts in the hurdle parcour between USH1C animals and WT controls in the dark confirms that pigs substantially rely on non-vision sensing and denies an exclusive or dominant effect of vestibular dysfunction on movement. Second, the clear differences in illuminated environment and

**Figure 7. Therapeutic approaches.**

A. AAV transduction of retinal layers. AAV8, AAV9, and AncB0 encoding eCFP under the ubiquitous CMV promotor were injected subretinally into the eyes of mature WT pigs in vivo. Upper panel: Epifluorescence of dissected eyes demonstrates eCFP fluorescence in the bleb region of whole mount preparations of all three AAV samples. Dashed circles indicate bleb size. Lower panel: Epifluorescence images of longitudinal retina sections revealed eCFP expression in the PRC layer (OS, IS and ONL) for all AAVs. Increased exposure (4×) documented transduction of MGC (arrows) by AAV0 and AncB0 but not by AAV8 (representative images from 1 pig, 1 retina each, 2 TR, scale bars 25 µm).

B. CRISPR/Cas-mediated gene repair was tested in primary cells of USH1C pigs (see also Fig EV4). Left panel: The cutting site (red arrow) of the most favorable gRNA was directly at the transition from exon 2 (pink) to the downstream intron. Positions of the correcting mutation (blue) and blocking mutation (magenta) are indicated. Lower left: Distinct ssODN binding to the sense (green) or anti-sense strand (orange) were compared. Upper right: Sanger electropherograms confirmed the correct introduction of the correcting (blue) and blocking (magenta) mutations and indicated by standing NHEJ mutations (grey, overlapping nucleotide peaks) at the gRNA cutting site. Lower right: Frequencies of HDR (blue) and NHEJ (grey) in mixed cell populations were similar for the tested ssODN templates (left). 68 single cell clones treated with urt1.3, indicated that HDR appeared either exclusively (dark blue) or was accompanied by intronic (blue/white shaded) or exonic (blue/gold) mutations (right). In some clones, HDR transformed only the blocking mutation site which is more closely located to the cutting site (white/gold). NHEJ events appeared as insertions (dark grey) or deletions (light grey). The frequency of unmodified clones is depicted in brown.

C. Effects of harmonin_a2 expression were investigated in dermal fibroblasts. Left panel: Transfection of USH1C primary cells with an expression plasmid (USH1C+) increased the number of ciliated cells toward WT level (3 TR, number of examined cells is indicated, error bars indicate SD, two-tailed Student’s t-test, *P < 0.05). Middle and right panel: Length of cilia in USH1C primary cells was shifted toward WT level (3 TR, number of examined cells is indicated, error bars indicate SD, Kolmogorov-Smirnov test, *P < 0.05, ****P < 0.0001).

D. Efficacy of AAV gene therapy was examined in vivo by sub-retinal injection of AncBO virus capsid expressing harmonin_a2 under a ubiquitous promoter into a USH1C pig (see also Fig EV5). Left panel: Sub-retinal injection restored harmonin expression in the AAV-treated eye but not in the sham-treated eye (3 TR). Protein extract from a WT retina served as positive control, actin staining as loading control. The 72 kD band correlates with harmonin_a isoforms while higher MW indicates putative dimer formation. Middle panel: Immunofluorescence reveals absence of harmonin in the retina of the PBS-injected control eye and harmonin abundance (green) in the retina of the AAV-treated eye. (Representative image from a single animal. Blue: DAPI, scale bar 25 µm). Right panel: Photopic gERG reveal increased a- and b-wave response to single flash and flicker stimulation in the AAV-treated (red), compared to the sham-treated (black) eyes of two USH1C pigs.
Figure 7.
the characteristic difficulties that USH1C pigs have in the recognition of defined objects (Fig 2A-C) document their substantial problems in orienting within a structured 3D space. Although an influence of the bilateral vestibular dysfunction on orientation cannot be excluded (Kremmyda et al., 2016), the faster and smoother locomotion (Fig 2D) of WT control pigs, specifically under light conditions, demonstrates how a normal visual function promotes effective clearance of obstacles and “elegant” motion.

Taken together, the generation of an USH1C pig model has substantially improved our molecular understanding of the disease. We now have several lines of evidence how harmonin supports PRC function and the particular integrity of the OS. The early onset of basic defects in the retina and the impairment during the first year provides an ideal foundation for following the natural history of USH1C. Combining quantitative motion analysis with clinical examination tools will promote the definition of read-out parameters in pre-clinical studies and in clinical trials for treating the Usher syndrome, but also in trials involving patients with other progressive blinding or audiovestibular diseases. The long-term maintenance of retinal structures and, moreover, the success of exploratory gene therapy approaches in older animals is highly relevant as it indicates a large window of therapeutic opportunity for USH1 patients.

Material and Methods

Regulatory statement

Work on USH pigs has been conducted under the supervision of the responsible regulatory authorities: the Regierung von Oberbayern has approved animal experiments involving somatic cell nuclear transfer, maintenance of pigs, and longitudinal monitoring at LMU Munich under the file number AZ 02-17-136. The State Veterinary Administration of the Czech Republic approved animal experiments on maintenance, sub-retinal intervention and longitudinal monitoring at IAPG Libechov under the experimental protocol number 75/2019. Human primary cells were provided by University of Tuebingen, Germany, work on the cells was carried under approval of the ethics committee of the Landesaerztekammer Rheinland-Pfalz under the file number 837.413.17. Informed consent was obtained from all subjects providing skin biopsies. The experiments on human fibroblasts conformed to the principles set out in the WMA Declaration of Helsinki and the Department of Health and Human Services Belmont Report.

Antibodies, oligos, and gene synthesis

see Supplemental Materials and Methods.

Generating USH1C pigs

Bioinformatics evaluation

USH1C loci were identified in macaque, marmoset, cattle, sheep, pig, horse, cat, dog, rabbit, mouse, and rat by BLAT using human exonic sequences and extracted from the respective reference genomes (www.ensembl.org). Multi-species alignments were done by a combination of DiAlign and CHAOS (Brudno et al., 2004). Proposed regulatory elements were extracted from the UCSC Genome browser (genome.ucsc.edu) as DNAse sensitive elements (Thurman et al., 2012). ENCODE ChIP-chip (Wang et al., 2013), FANTOM5 enhancer elements ( Andersson et al., 2014), GeneHancer elements (Fishlinevich et al., 2017), FAIRE ENCODE regions ( Giresi et al., 2007) and PreMod ( Ferretti et al., 2007). Any identified regulatory element was localized within the alignment in BioEdit (https://bioedit.software.informer.com/7.2/) and a density plot was created. Both, a homology plot at nucleic acid level and a protein conservation plot were calculated in BioEdit on the basis of the nucleotide summary assessment and entropy calculation functions, respectively. Coverage maps of the augmented alignment were plotted by JalView ( Waterhouse et al., 2009).

Personalized R31X allele

For introducing a patient-specific segment, we analyzed a 1.6 kb fragment from an individual carrying an USH1C<sub>R31X</sub> mutation. The PCR product was amplified with the primers hush1c<sub>2f</sub> and hush1c<sub>2r</sub>, cloned into a plasmid vector and sequenced. For avoiding the misinterpretation of potential PCR mistakes, the sequence of the R31X-allele was based on 6 distinct clones and the other allele on 3 distinct clones. Consistently appearing polymorphisms were defined as SNPs.

Targeting construct

The porcine USH1C gene has been annotated to chr2:42Mb, with the exons largely corresponding to the human USH1C transcript variant b3, GenBank no. NM_153676.4 (Appendix Fig S1). Distinct porcine primary cell lines (Richter et al., 2012) were examined for SNPs in the target region and 3 distinct gRNAs, rk1, rk3, and rk4 that were not affected by cell line-specific SNPs were examined for their power to introduce NHEJ-based mutations in pig primary cells lines; for this, cells were nucleofected and after 48 h the target locus was amplified with the primers ushwt1f and ushwt1r, cloned into plasmids and sequenced. Targeting experiments were based on rk4 as it proved superior (12% mutation rate) to rk1 (6%) and rk3 (10%). BACs CH242-515C3 and CH242-199J14, were identified to completely carry the porcine USH1C locus via the Sscrofa 10.2 reference genome and the PigPre BAC map (www.ensembl.org) and purchased from CHORI BACPAC Resources Center (BACPAC Genomics, Emeryville, CA, USA). A sequence comprising a 278 bp 5’-arm, the 1,524 bp humanized fragment including the TGA stop codon in exon 2 and 3 intronic SNPs as well as a 278 bp 3’-arm were commercially synthesized and provided in a pUC plasmid derivative by GeneArt (Invitrogen by Thermo Fisher Scientific, Carlsbad, CA, USA). A cassette with a PGK/EM7 driven neo gene, providing neomycin resistance in mammalian cells and kanamycin resistance in bacteria (Klymiuk et al., 2012), was cloned into the plasmid via the NotI/XhoI sites located between the 5’-arm and the humanized fragment. The entire fragment was excised from the backbone via Ascl and introduced into the CH242-515C3 BAC by bacterial recombination, according Vochozova et al. (2019) using the SW106 E. coli strain (Warming et al., 2005). Correct modification of the BAC was proven by PCR for recombination at the 5’- and 3’-ends with the primer pairs 5arm1f – 5arm1r and 3armwfl – 3armwfr, and for the removal of the corresponding porcine segment with the primer pair ushwt1f – ushwt1r. Further, the overall integrity of the BAC and changes in the restriction enzyme (RE) recognition sites that arise...
from to the modification were confirmed by the RE pattern after digestion with XbaI (Fig EV1C) and SpeI (Appendix Fig S1D).

**Generating pigs**
Plasmids carrying the modified BAC, the gRNA rk4 and Cas9 were prepared endotoxin-free and transfected into pig primary cells PKC1 and PKCm by nucleofection (Amaxa by Lonza, Basel, Switzerland). Single cell clones were generated as described (Richter et al, 2012) and propagated toward a 2 × 96-well scale. One aliquot was used for analysis by the loss-of-wild-type-allele approach (LOWA) (Vochozkova et al, 2019) and the other served as a backup for potential SCNT. qPCR was carried out on a LightCycler96 (Roche Life Science, Basel, Switzerland) using FastStart Essential DNA Green Master (Roche Life Science) and primer pair ush1c_qf1 – ush1c_qr1 for the target site as well as the primer pair o4_qf2 – o4_qr1 for a reference site in the POU5F1 gene and ng_qf6 – ng_qr4 for a second reference in the NANOG gene. Verified single cell clones were used as donors in SCNT and embryos were transferred to synchronized gilts, according to established procedures (Albl et al, 2015). Birth was introduced in pregnant foster mothers by Estramate at day 115. (MSD Animal Health, Merck, Kenilworth, NJ, USA).

**Tissue sampling**
For isolation of pig primary kidney cells and sampling of tissue, animals were sedated with ketamine 100 mg/ml (Ursotalin®, Serumwerk Bernburg, Germany) and azapenone 40 mg/ml (Stresnil®, Elanco Animal Health, Bad Homburg, Germany), according to the manufacturer’s specifications. Fully anesthetized animals were euthanized by intravenous injection of T61® (MSD Animal Health). Pig primary kidney cells were isolated according to standard procedures (Richter et al, 2012) (see below). Tissue samples for histology, electron microscopy, and molecular analyses were collected according to established sampling guidelines adapted to porcine biomedical models (Albl et al, 2016). Samples for molecular analysis were frozen on dry ice and stored at −80°C. Tissue was powdered using hammer and anvil first and pestle and mortar afterward. All instruments and samples were cooled in liquid nitrogen to avoid thawing of the samples during processing. Powdered tissue was transferred to a pre-cooled tube and stored at −80°C.

**Breeding**
After reaching fertility and stabilization of cycle, USH1C F0 sows were inseminated with WT sperm. When pregnant, sows were trained and reassured intensively for farrowing by regular physical contact with caretaker. F1 offspring were genotyped (see below). F1 boars were mated to their mothers and sisters for producing USH1C-null pigs and with wild-type sows to broaden the genetic background of the future breeding herd. Some litters from hom x het matings were with their mother for suckling, but piglets were kept in a separate box otherwise. Upon calling by the mother, the barrier between the boxes was opened and the time until piglets passed the gate was determined. Non-responding piglets were nudged by the caretaker after one minute to avoid impaired nurturing.

**Molecular analysis**

**Genomic level**
Genomic DNA was isolated from tail biopsies by using the Easy DNA kit (Invitrogen), DNAeasy kit (Qiagen, Hildesheim, Germany) or Nexttec kit (Nexttec GmbH, Leverkusen, Germany). For verifying the abundance of the modification element, amplicons generated with the primer pair hush_for3 – hush_rev3 were sequenced. The presence of the neo selection cassette was detected by endpoint PCR with ushar2f – usharm2r and sequencing of the amplicon while Cre-mediated excision of neo was detected with neoF03 – neoREV3 and sequencing of the amplicon.

**Transcriptional analysis**
RNA was isolated from powdered tissue samples using TRIzol (Invitrogen). Approximately 100 mg of powdered tissue was grinded up in 1 ml of TRIzol by Polytron PT2500E (Kinematica, Luzern, Switzerland) in a pulsatil manner to avoid overheating. Further steps were carried out as suggested by the TRIzol protocol and RNA was stored at −80°C. For cDNA synthesis, samples were treated first with DNase I (Invitrogen) to remove a possible contamination with genomic DNA. Then, RNA was reversely transcribed into cDNA with SuperScriptTM III Reverse Transcriptase (Invitrogen). For determining USH1C transcripts, cDNA was used in RT–PCRs with the primer pair ushrt2f – ushrt2r or ushrt2f – urt4r, located in the consistently transcribed exons 1 and 8 or 6, for detecting all splice variants of the gene. Primer pairs ushrt11f – ushrt11r and ushrt13f – ushrt13r were used to amplify USH1C_a and USH1C_c variants (pig specific primers binding on exons according the splice pattern of human GenBank entries NM_005709.4 and NM_001297764.2, respectively). Primer pairs ushrt16f – ushrt16r and ushrt18f – ushrt18r were used to amplify USH_b splice variants (pig specific primers binding on exons according the splice pattern of human GenBank entry NM_153676.4). Afterward, Sanger sequencing was performed on PCR amplicons.

**Protein analysis**
Tissue was powdered and then processed as previously described (Sedmak & Wolfrum, 2011). In brief, retinal tissues were lysed in modified RIPA buffer (50 mM Tris–HCl, 150 mM NaCl, 0.1% SDS, 2 mM EDTA, 1% NP-40, 0.5% sodium-deoxycholate, 1 mM sodium-vanadate, 30 mM sodium-pyrophosphate, pH 7.4). Protein lysates were separated by SDS–PAGE gel electrophoresis, followed by semi-dry Western blotting as previously described in (Overlack et al, 2011). Western blots were analyzed and quantified in an Odyssey infrared imaging system (LI-COR Biosciences, Lincoln, NE, USA).

**Morphology & histology**

**Immunohistochemistry**
Porcine eyes were fixated in melting isopentane for cryosectioning as described (Overlack et al, 2011). Samples were sectioned with a MICROM HM 560 Cryo-Star cryostat (Fisher Scientific by Thermo Fisher Scientific, Waltham, MA, USA) and placed on poly-L-lysine-precocated coverslips. After drying, samples were incubated with 0.01% Tween 20 PBS, then washed with PBS and covered with blocking solution (0.5% cold-water fish gelatin, 0.1% ovalbumin in PBS) for a minimum of 30 min followed by overnight incubation at 4°C with primary antibodies (for details on antibodies and fluorescent dyes see below). For GFAP staining samples were post-fixated with 4% paraformaldehyde (PFA) in PBS. Cryosections were washed repeatedly and incubated with secondary antibodies in blocking solution containing DAPI (Sigma-Aldrich, St Louis, MO,
USA) for 1 h at room temperature. After washing, sections were mounted in Mowiol (Roth, Karlsruhe, Germany). Slides were analyzed on a Leica DM6000B microscope (Leica, Bensheim, Germany); images were processed with LAS-AF Leica imaging software and ImageJ/Fiji software (Schindelin et al, 2015) or Adobe Photoshop CS (Adobe Systems, San Jose, CA, USA).

Quantitative analyses of connecting ciliary length and cone synaptic width: For measurements of connecting cilia length and cone pedicle synapses we stained cryosections through pig retinæ with either GT335, a marker for tubulin glutamylation in the connecting cilia or FITC-PNA, a marker for the extracellular matrix of cones, respectively. For the image analysis, we developed Fiji scripts by ImageJ/Fiji software for automatic analysis of the fluorescent images.

The script for measuring connecting cilia length is:

```
run("8-bit");
run("Smooth");
run("Auto Threshold", "method=Yen ignore_black white");
run("Analyze Particles...", "size=10–75 pixel display add");
```

The script for measuring synapse width is:

```
run("8-bit");
run("Mean...", "radius=2 stack");
run("Subtract Background...", "rolling=50 stack");
//setThreshold(0, 11);
run("Convert to Mask", "method=Otsu background=Light black");
run("Measure").
```

Scanning electron microscopy (SEM)
The temporal bones of the pigs were dissected to expose the middle and inner ears. Their cochleæ were dissected immediately and the organ of Corti was exposed. The tectorial membrane was removed prior to fixation. Dissected cochleæ were fixated in 2.5% glutaraldehyde, 4% PFA in 0.1 M Sörensen’s phosphate buffer. After several washing steps with 0.1 M Sörensen’s phosphate buffer, cochleæ were dehydrated in ethanol, critical-point dried, and gold sputtered in an argon atmosphere. Specimen were imaged with a Philips ESEM XL30 scanning electron microscope (Philips, Eindhoven, Netherlands).

Transmission electron microscopy (TEM)
For conventional TEM dissected eyeballs were pre-fixated for 2 h in buffered 2.5% glutaraldehyde containing sucrose and post-fixated in buffered 2% OsO4 as previously described (Karlstetter et al, 2014). After dehydration in ethanol and passing through propylenglycol as intermediate, samples were embedded in Renlam®-M-1 resin. Flatmount samples were heat-polymerized and glued on the top of empty Araldit blocks. Ultrathin sections of the embedded specimens were prepared with an Ultracut S ultramicrotome (Leica) and collected on Formvar-coated copper or nickel grids. Sections were counter-stained with heavy metals before imaging and analyzing in a Tecnai 12 BioTwin TEM (FEI by Thermo Fisher Scientific, Hillsboro, OR, USA), equipped with a SIS Megaview3 CCD camera (EMSIG, Münster, Germany) was used. The images were processed using Adobe Photoshop CS (Adobe Systems). Connecting cilia length or synapse width were analyzed directly in the TEM using the measuring tool of the Olympus SIS image analysis system (Soft Imaging Systems, Muenster, Germany). Quantification steps have been performed with Fiji/ImageJ.

Primary cell culture
Pig primary kidney cells and primary dermal fibroblasts were isolated according to standard procedures (Richter et al, 2012). In brief, kidneys were taken from euthanized animals, washed in PBS containing Penicillin/Streptomycin and minced. Tissue was treated with collagenase and filtered cells were cultivated in DMEM. Pieces of porcine skin were disinfected in 98% Ethanol and after short washing in PBS placed in 10% Betaisodona solution. After repeated washing steps in PBS, skin was minced. Kidney or skin samples were transferred to cell culture dishes and completely covered with MEM containing 10% heat-inactivated fetal calf serum (FCS) and 1% penicillin-streptomycin (PS) and incubated at 37°C and 5% CO2. 5 days later 10% FCS were added. Outgrown cells were transferred to new complete medium for further culturing. For ciliogenesis experiments, 5 × 105 cells were seeded per well (in medium containing 10% FCS and 5% PS) and 24 h later cultured in OPTI-MEM reduced-serum medium (Invitrogen by Thermo Fisher Scientific). After 48 h of starvation, cells were washed in PBS and fixed with 2% PFA in PBS. After washing and permeabilization with 0.1% Triton-X in PBS, cells were blocked with 0.5% cold-water fish gelatin, 0.1% ovalbumin in PBS followed by incubation with primary antibodies overnight at 4°C. After washing, coverslips were incubated with secondary antibodies for 1 h at room temperature. Samples were washed and then mounted in with Mowiol (Roth).

The primary ciliary length and number of ciliated fibroblasts were quantified by using Fiji. Lengths were measured directly in immunostained images taken with a 63-x objective. For this purpose, the Arl13b-positive axoneme was measured from the tip to the cilia base, which was labeled for pericentrin, a common marker for the ciliary base.

Clinical examination
Anesthesia
For examination with ERG, mfERG, OCT, AF, FA, and ABR on 3-week old animals were anesthetized by intramuscular injection of 2 mg/kg azaperone (Stresnil®, Elanco, Germany), 0.02 mg/kg atropine sulfate (B. Braun, Germany) and 20 mg/kg ketamine (Ursotamin®, Serumwerke Bernburg, Germany). Anesthesia was continued by intravenous injection of propofol (Fresenius, Germany) according to effect. After endotracheal intubation, pigs were mechanically ventilated and cardiovascular function was monitored throughout the procedures. To exclude eyeball...
movement during examinations, a peripheral muscle relaxant (Rocuronium, Inresa, Germany) was applied. For ABR on older animals, animals were starved overnight before intramuscular application of TKX (tiletamine 4 mg/kg, zolazepam 4 mg/kg (Zoetil 100; Virbac), ketamine 5 mg/kg (Narketan 10; Chassot), and xylazine 1 mg/kg (Rometar 2%; Spofa).

Auditory brain stem response (ABR)
In 3-week-old USH1C and WT piglets, ABR was recorded with standard electrodiagnostic equipment (Viking Quest®; Natus, Planegg, Germany) and Natus® TIP-300 insert earphones (AF). Recording stainless steel needle electrodes were positioned subcutaneously ipsilateral to the stimulated ear over the mastoid at the base of the ear (−) and at the vertex in the midline of the skull corresponding to Cz (+). A ground electrode was positioned in the neck. Electrode impedance was < 2 kOhm. Click stimuli (100 μs, alternating polarity, 11.1 Hz) were delivered at supramaximal stimulation intensity (100 dB SPL) to one ear and masking noise was applied to the contralateral ear (−40 dB SPL). Filter settings were 100 Hz for high pass and 3 kHz for low pass filters. The ABR represented the averaged signal of 1,000–1,500 10 ms recordings. ABR was recorded twice for each ear to ensure reproducibility of the ABR peaks in WT animals (n = 3) or absence of recognizable peaks in USH1C-null animals (n = 3), respectively. If there was no recordable ABR at 100 dB SPL in USH1C-null animals, ABR was also conducted at 120 dB SPL (twice for each ear). Additional testing comprised ABR at decreasing steps of 10 dB in WT piglets. Hearing threshold was determined as the minimal click intensity that still evoked a noticeable potential negative peak after peak V. ABR for other time points were conducted with four subcutaneous needles placed in vertex, forehead, and mastoids with specialized audiometric equipment. 8 weeks and 2 years old animals were examined with eABRUSB (eABR—BioMed Jena GmbH, calibrated by standard norm IEC 318-4) and headphones (Sennheiser Momentum M2 In-Ear G Black-Red, 18Ω) were used on the 8 weeks old piglets. Click stimuli (duration 100 μs, steps 10 to 5 dB) and tone pips (duration 100 μs, frequencies 2, 4, 8, 16 kHz, steps 10 to 5 dB) were applied. The evoked signal was filtered by band-pass filter of range from 300 Hz to 3 kHz. All the responses were averaged 200–400 times for each intensity and analyzed with eAudio software. On 3 months, 6 months and 2 years old pigs, acoustic stimuli were generated with an evostar 2 system (ERA system: Evoselect/Evostar 2, Pilot Blankenfelde medizinisch elektronische Geräte GmbH) and presented to the animal via headphones (beyerdynamic, DT48, 5Ω, Germany). Click stimuli (duration of electric pulse 150 μs) and the tone pips (duration 2 ms, frequencies of 0.5, 1, 2, 3, 4 kHz, intensity steps from 10 to 5 dB) were applied. The signal from an electrode was filtered by band-pass filter over the range of 100 Hz to 2.5 kHz. The response was averaged 200–500 times on each intensity. This signal was processed with and analyzed using Evostar software. The threshold response to each frequency and click was determined as the minimal tone/click intensity that still evoked a noticeable potential peak in the expected time window of the recorded signal. This evaluation process was the same for both systems.

Electroretinography (ERG)
Full-field ERG was recorded with a corneal Kooijman electrode (Roland Consult, Brandenburg an der Havel, Germany). Platinum needles were placed subcutaneously at the temple and top of the head between the ears as reference and ground electrodes respectively. Stimuli were brief white flashes (4 ms) delivered from a light source within the Kooijman electrode (Kooijman & Damhof, 1986). The RETImap system (Roland Consult) was used for stimulus generation and data acquisition. The recording protocol used was based on, and extended from, the International Society for Clinical Electro-physiology of Vision standard for human clinical full-field ERG. A band-pass filter was used to remove frequencies outside 1–300 Hz. An interstimulus interval (ISI) of 1 s was used with 20 responses recorded and averaged for each single flash protocol. For scotopic ERG, the protocol consisted of 30 min dark-adaptation, followed by recording of a dark-adapted single flash series with increasing luminance from −3.0 log cd*s*m⁻² to 2.0 log cd*s*m⁻². After completion of the dark-adapted intensity series, animals were light adapted for 10 min to a steady white background of 30 cd*s*m⁻² with the same light source. Light-adapted standard flash was recorded by using white flash stimuli at 0 log cd*s*m⁻². The same luminance was used to record a 28 Hz flicker response for 300 ms. For multifocal ERG, the mERG was recorded using the same RETImap system (Roland Consult) using a 61-hexagon stimulus and according to the standard of the International Society for Clinical Electrophysiology of Vision (Hood et al, 2012). A 61-hexagon stimulus was presented on a 51-cm cathode ray tube monitor with a frame frequency of 60 Hz. It encompassed the central visual field of 44° horizontally and vertically. The scaled-size hexagons (distortion factor 4) were light-modulated according to a binary pseudorandom m-sequence. The light state had a luminance of 120 cd/m² and the dark state of 1 cd/m². Refractive errors were not corrected. Pupil dilation was achieved with one drop of tropicamide–phenylephrine mixture into the conjunctival sac of each eye about 15 min before the exam. Contact lens electrodes (ERG-jet®, CareFusion, San Diego, USA) were applied. Amplifier gain was set to 50,000 and the band-pass from 5 to 100 Hz. There were eight recording cycles of 47 s each. The 61 traces are displayed separately in a false-color-coded amplitude map identifying P1 amplitude (measured from the N1 trough to the following maximum positive peak; N1 amplitude was measured from the electroneutral starting point to the first negative deflection). All traces were also grouped into one response value and the averaged first-order kernel function displayed as single trace. Luminance levels in the pig eye were calculated from the light intensity measured at the floor. Given the complex physical correlation between light source, reflection, and sensing in an indoor 3D space, it is challenging to determine corresponding luminance levels in the pig eye, but considering the general reflexion parameter p as 0.2 seems a fair approximation.

Optical coherence tomography (OCT)
Spectral-domain OCT B-scans of central retina in animals were obtained using a Spectralis HRA + OCT device (Heidelberg Engineering, Heidelberg, Germany) under general anesthesia and during full pupil dilation. During the acquisition, the corneal surface was protected with methylcellulose eye drops while lids were held open using lid specula. After 30deg and 55deg infrared and red-free images were recorded using scanning laser ophthalmoscopy (SLO), animal received a flush of approximately 5 mg/kg fluorescein and the angiography was recorded using the 55 degree lens at multiple time points (0.5, 1, and 5 min).
Behavior testing

Barrier course

The design was adapted from a previous publication (Barone et al., 2018). Ten barriers (boards of 94 × 76 cm) were placed in a distance of one meter in a straight aisle (12 × 2.2 m) alternating in the middle, the left or right positions. There were four different arrangements of barriers, which were used alternatingly (Appendix Fig S4). Pigs were trained in the course until they walked straight through the course without pausing or hesitating for at least two times in a row. After a training period of four weeks, the animals were tested in the course weekly. The pigs entered the course individually in random order, they had to pass the barriers, and at the end, they were rewarded with fruit juice in a food bowl. The test was performed before feeding to ensure a higher motivation in the pigs. After some runs in the light, the test was conducted alternately in the light (average of 135 lux) and in the dark (average of 2.9 lux). Illumination level was measured at three different sites in the course (start, middle and end) in animals’ head height. Each run was documented by video. The pigs were marked with numbers on their backs to facilitate recognition. The times pigs took for passing the course, from snout crossing the start line until the pig was touching the food bowl, were measured. All barrier contacts by the pigs were counted. Contacts were divided into two subgroups. Frontal contacts mean the pig was touching the barrier with its snout or head, while contacts were called lateral when the pig was touching the obstacle barrier with its shoulders or hips. Each turning around or walking back before reaching the end was counted. It was distinguished between turns in front of the course and turns within the course. Such runs were excluded. After turning around, animals were sent back to the course, and the new run was documented with time and contacts when they reached the end. Walking in circles was counted as a separate event as well. As long as they were just circling and not walking back, the run was not excluded, time measurement was continuing. When the pig was circling in front of the start line, those circles were also counted, but time measurement only started when the animal was crossing the start line. Pigs were excluded from parcour testing when they were lame or when they had already been outside of their pen on the experiment day for another reason. When a pig was turning around or circling so often that it needed longer than two minutes to walk to the endpoint, the run was also excluded. The numbers of turns/circles within those two minutes were nonetheless counted. Pigs were not included in the study when motivation was lacking, for example, in the case a pig did not head to the endpoint reliably after six weeks of training or when it seemed more interested in playing with the barriers than walking straight to the food bowl.

Obstacle course

An obstacle course to test the visual capacity of USH1C pigs and WT pigs was conducted. The course was 6.15 m long, the width was 1.58 m in the beginning and 2.48 m in the end. Most obstacles were built from obstacles used for equestrian sport, other obstacles were a stepboard and barrels. Obstacles were rearranged for almost each test day (Appendix Fig S5).

Pigs were trained until they walked straight through the obstacle course to the food trough without stopping or hesitating. After a training period of three weeks, the pigs were performing the course once a week. The pigs entered the course individually in random order, they had to pass the obstacles and at the end they were rewarded with a cookie in a food trough. The test was conducted under light and dim conditions. Each run was documented by video. The pigs were marked with numbers to facilitate recognition. The times the pigs needed to pass the course were measured. All contacts with the obstacles or the wall were counted. When a pig nudged one obstacle more than once in a playful way in the same area, this was counted as one touch. When a pig touched one obstacle at different sites, each contact was counted separately. When a pig swept along an obstacle with its snout, this was counted as two touches. When a pig turned around before reaching the end or when a pig was lame, the pig was excluded for the test day. For analyzing the trajectory, the dimensions of the course and the gaps of the slatted floor were measured and scaled at 1:100 (Macromedia FreeHand MX, STUDIOMX 2004). Using the slatted floor as a ruler, each footstep of each animal in each qualified run were plotted in the map. X and Y coordinates of footsteps as well as the corresponding times when the foot touched the ground were transferred to an excel file. Trajectories were analyzed from these data sets with the R package trajr. Briefly, we first imported and smoothed all trajectories to avoid any source of noise by applying a Savitzky-Golay smoothing filter (Savitzky & Golay, 1964; Luo et al., 2005). Then, we called all the functions of the trajr packages (McLean & Skowron Volponi, 2018) to characterize the trajectories into measures of speed, length, distance, duration and measures of straightness or tortuosity. The trajectory significance was tested using Kruskal–Wallis test. For statistical evaluation, data were checked for consistency and normality. Fisher’s exact test or Pearson’s test were used to analyze cross tabulations. Generalized linear models with Poisson distribution, Median tests, bootstrap-t tests based on 5000 Monte Carlo simulations, t-tests with and without the assumption of homogeneity, Mann–Whitney U-tests were used to test continuously distributed variables. All reported tests were two-sided, and P < 0.05 were considered statistically significant. Analyses were performed by use of NCSS_10 (NCSS LLC, Kaysville, UT, USA), STATISTICA_13 (Statsoft, Tulsa, OK, USA), MATHEMATICA_12 (Wolfram Research, Hanborough, UK), Champaign, IL (2018) and PASW 24 (SPSS by IBM Corp, Armonk, NY, USA), and Prism_5 (GraphPad Software, San Diego CA, USA).

Therapy approaches

AAV transduction of retinal layers

Distinct AAVs (AAV8.CMV.eGFP, AAV9.CMV.eGFP, and Anc80.CMV.eGFP) were generated by the viral core facility at Boston Children’s Hospital Viral Core at the Massachusetts Eye and Ear Infirmary. AAVs were injected subretinally into the eyes of WT pig at an age of 30 months using an ophthalmologic surgery microscope (Hi-R NEO 900A, Haag-Streit). Using inhalation anesthesia (Morpheus E, Siare), morphology of the retina was examined via OCT iVue (Optovue, Fremont, CA, USA) on both eyes before the injection. 200 µl of AAVs (2 × 10¹⁰ AAV particle (vg)) were applied in the subretinal space with Extendible 41G subretinal injection needle (23 gauge/0.6 mm) (DORC, Zuidland, Netherlands). After 5 weeks injection sites were evaluated by OCT (see above) and funduscopy, confirming the concise injection sites. Following enucleation, the eyecup was dissected and fixed with 4% PFA in
PBS. Flattened whole mount preparations of the eyecup were analyzed by epifluorescence using a handheld device (Bluestar Flashlight, Nightsea, Lexington, MA, USA). Samples were processed for cryosectioning and analysis as described above.

**Gene repair in vitro**

Different constellations of the gRNAs urg1 and urg2 with correcting ssODN urt1, urt2, urt1.1, urt1.2, urt1.3, urt1.4, and urt1.5 were tested in primary fibroblasts. For this, plasmids encoding Cas9 and the respective gRNA as well as correcting oligonucleotides were nucleofected under standard conditions (Richert et al., 2012). After 24–48 h of cultivation, DNA was isolated from mixed cell batches and the efficacy of NHEJ and HDR were determined by examination of Sanger electropherograms of amplicons generated by PCR with the primer pair huUSH1f – huUSH2r and sequenced with primer ush1s in the web-based ICE v2 CRISPR Analysis Tool (https://www.synthego.com/products/bioinformatics/crispr-analysis by Synthego, Menlo Park, CA, USA). For examination of specific allelic constellations, cells nucleofected with urg1 and urt1.3 were seeded onto 96-well plates and single cell clones were produced. DNA was isolated from independent clones and PCR amplicons were generated by primer pair ushHS1f – huUSH1r, sequenced with primer ush1s, and analyzed by Sanger sequencing.

**Gene therapy in vitro**

For restoring gene expression, 2.5 × 10⁵ primary porcine fibroblasts were transfected with 0.4 μg of an endotoxin-free pDest_Harm_a1/S/F plasmid (Invitrogen), expressing splice form a1 of the human USH1C gene under the control of the CMV promoter. Transfection was performed with the 4D-NucleaseFactorTM X Unit (Lonza), using the P2 Primary Cell 4D-NucleaseFactorTM X Kit S and program FS113, according to the manufacturer’s protocol. After transfection, cells were seeded in 10% FCS and analyzed for ciliogenesis as described above.

**Gene therapy in vivo**

1 × 10¹¹ vg Anc80.CMV.harmonin_a1 in 200 μl PBS (manufactured by Boston Children’s Hospital Viral Core) was injected subretinally into the right eye, while the left eye was sham-treated with 200 μl PBS vehicle, according to the procedures described for the AAV transduction experiment (see above). Animal #10056 was injected at an age of 2.5 years and experiment was terminated after 12 months after gFhERG examination. Eyes were collected from the vesicular and non-vesicular areas of the retina for immunofluorescence and Western blot analyses. Animal #10054 was injected at an age of 3.5 years. After intermediate gFhERG examination 3 months after injection, the experiment is ongoing.

**Data availability**

This study includes no data deposited in external repositories.

**Expanded View** for this article is available online.

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**Author contributions**

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**Zdenka Ellederova:** Methodology; Project administration. **Hannah Auch:** Validation; Investigation; Methodology. **Andrea Bahr:** Supervision; Investigation; Methodology. **Petra Runa-Vochozkova:** Supervision; Investigation; Methodology. **Janet Fadi:** Investigation. **Vanessa Arnold:** Investigation. **Taras Ardan:** Investigation; Methodology. **Miroslav Veith:** Investigation; Methodology. **Jiri Popelar:** Investigation; Methodology. **Valeri Zakharchenko:** Methodology. **Joshua Linnert:** Investigation; Methodology. **Stepanka Suchankova:** Validation; Investigation; Methodology. **Jiri Popelar:** Supervision. **Eduardo Rodriguez-Bocanegra:** Validation. **Julia Dlugaczyk:** Supervision. **Hans Straka:** Formal...
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In addition to the CRediT author contributions listed above, the contributions in detail are:

SG contributed to molecular analysis and breeding of animals, designed and performed behavior tests and contributed to writing of the manuscript. JS contributed to molecular and morphological analysis, and ciliation analysis in primary fibroblasts. KAW contributed to tissue sampling and processing, imaging (IF and SEM), data analysis and interpretation. ZE contributed to breeding attempts sub-retinal intervention and ABR measurements. HA evaluated Gene Repair. AB contributed intervention on pigs and anesthesia. PR-V contributed to Gene Repair study. JF contributed to genetic modification. VA contributed to quantitative analysis of the synaptic phenotype in cones. TA managed intervention on pigs. MV performed subretinal injection. GS performed data analysis. BK contributed to Somatic Cell Nuclear Transfer and performed Embryo Transfer. CE performed RT–PCR on USH1C transcripts. JK participated in the molecular and morphologic analysis of eyes after USH1C gene therapy. ABR performed ERG measurements. JI contributed ciliation analysis in primary fibroblasts. MK and VZ performed Somatic Cell Nuclear Transfer. AF, SS, and JP performed ABR measurement. ER-B participated in ophthalmologic analysis. AD contributed to the molecular and morphological analysis of animals, performed the vitreoretinal surgery and contributed writing of the manuscript. UW designed and coordinated the study, performed molecular and morphological analysis, interpreted data and wrote the manuscript. NK designed and coordinated the study, performed bio-informatic and molecular analysis, interpreted data and wrote the manuscript. ER-B contributed to analysis of ERG and OCT measurements.

Disclosure and competing interests statement

U.W., K.N.W., E.W. & N.K. are holders of patent US2020/0337279 A1.

References

Albl B, Haesner S, Braun-Reichhart C, Streckel E, Renner S, Seeliger F, Wolf E, Wanke R, Blutke A (2016) Tissue sampling guides for porcine biomedical models. Toxicol Pathol 44: 414 – 420

Andersson R, Cehbard C, Miguel-Escalada I, Hoof I, Bornholdt J, Boyd M, Chen Y, Zhao X, Schneideri C, Suzuki T et al (2014) An atlas of active enhancers across human cell types and tissues. Nature 507: 455 – 461

Baloh RW, Haimagyi GM, Zee DS (2012) The history and future of neuro- ophthalmology. Continuum (Minneap Minn) 18: 1001 – 1015

Barone F, Nannoni E, Elmi A, Lambertini C, Scorpio DG, Ventrella D, Vitali M, Maya-Vetencourt JF, Martelli C, Benfenati E et al. (2018) Behavioral assessment of vision in pigs. J Am Assoc Lab Anim Sci 57: 350 – 356

Boeda B, El-Amraoui A, Bahlouli A, Goodyear R, Daviet L, Blanchard S, Perfettini I, Fath KR, Shorte S, Reiners J et al. (2002) Myosin Vila, harmonin and cadherin 23, three Usher I gene products that cooperate to shape the sensory hair cell bundle. EMBO J 21: 6689 – 6699

Bonneau D, Raymond F, Kremer C, Klossek JM, Kaplan J, Patte F (1993) Usher syndrome type I associated with bronchiectasis and immotile nasal cilia in two brothers. J Med Genet 30: 253 – 254

Bringmann A, Landiev I, Pannicke T, Wurm A, Holllorn M, Wiedemann P, Osborne NN, Reichenbach A (2009) Cellular signaling and factors involved in Muller cell gliosis: neuroprotective and detrimental effects. Prog Retin Eye Res 28: 423 – 451

Brudno M, Steinkamp R, Morgenstern B (2004) The CHAOS/DIALIGN WWW server for multiple alignment of genomic sequences. Nucleic Acids Res 32: W41 – W44

Bujakowska KM, Liu Q, Pierce EA (2017) Photoreceptor cilia and retinal ciliopathies. Cold Spring Harb Perspect Biol 9

Canvalho LS, Xiao R, Wassmer SJ, Langsdorf A, Zinn E, Pacouret S, Shah S, Comander JJ, Kim LA, Lim L et al. (2018) Synthetic adeno-associated viral vector efficiently targets mouse and nonhuman primate retina in vivo. Hum Gene Ther 29: 771 – 784

Chandler MJ, Smith PJ, Samuelson DA, MacKay EO (1999) Photoreceptor density of the domestic pig retina. Vet Ophthalmol 2: 179 – 184

Corral-Serrano JC, Lamers IJC, van Reeuwijk J, Duijkers L, Hoogendoorn ADM, Yildirim A, Argrou N, Ruirogk RAA, Letteboer SJF, Butcher R et al (2020) PCARE and WASF3 regulate ciliary F-actin assembly that is required for the initiation of photoreceptor outer segment disk formation. Proc Natl Acad Sci USA 117: 9922 – 9991

Cvejic D, Steinberg TA, Kent MS, Fischer A (2009) Unilateral and bilateral congenital sensorineural deafness in owner pure-bred white cats. J Vet Intern Med 23: 392 – 395

Dougherty JM, Carney E, Emmady PD (2020) Vestibular dysfunction. Treasure Island, FL: StatPearls

Ebermann I, Phillips JB, Liebau MC, Koenekoop RK, Schermer B, Lopez I, Schafar E, Roux A-F, Dafiger C, Benrd A et al (2010) POZD7 is a modifier of retinal disease and a contributor to digenic Usher syndrome. J Clin Invest 120: 1812 – 1823

Egerer S, Fiebig U, Kessler B, Zakhartchenko V, Kuroe M, Reichbart B, Kupatt C, Klymiuk N, Wolf E, Denner J et al. (2018) Early weaning completely eliminates porcine cytomegalovirus from a newly established pig donor facility for xenotransplantation. Xenotransplantation 25: e12449

El-Amraoui A, Petit C (2005) Usher I syndrome: unravelling the mechanisms that underlie the cohesion of the growing hair bundle in inner ear sensory cells. J Cell Sci 118: 4593 – 4603

El-Amraoui A, Petit C (2014) The retinal phenotype of Usher syndrome: pathophysiologic insights from animal models. C R Biol 337: 167 – 177

Ernest S, Rauch GJ, Haffter P, Geisler R, Wieland T, Petit C, Nicolson T (2000) Mariner is defective in myosin VI: a zebrafish model for human hereditary deafness. Hum Mol Genet 9: 2189 – 2196

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Ferretti V, Poitras C, Bergeron D, Coulombe B, Robert F, Blanchette M (2007) PReMod: a database of genome-wide mammalian cis-regulatory module predictions. Nucleic Acids Res 35: D122 – D126

Fishilevich S, Nudel R, Rappaport N, Hadar R, Plaschkes I, Iny Stein T, Rosen N, Kohn A, Twik M, Safran M et al (2017) GeneHancer: genome-wide integration of enhancers and target genes in GeneCards. Database (Oxford) 2017: bax028

Fuster-Garcia C, Garcia-Bohorquez B, Rodriguez-Munoz A, Aller E, Jaijo T, Karam A, Tebbe L, Weber C, Messaddeq N, Morl...
ameliorates skeletal and cardiac muscle failure in pig and human models of Duchenne muscular dystrophy. Nat Med 26: 207 – 214

Nagel-Wolfrum K, Fadl BR, Becker MM, Wunderlich KA, Schaefer J, Sturm D, Gue R, Kaplan L, Goldmann T, Brooks M et al (2021) Characterization of USH1C/harmonin in the human retina provides insights into pathophysiology and therapy options for Usher syndrome. bioRxiv https://doi.org/https://doi.org/10.1101/2021.08.27.457962 [PREPRINT]

Ouyang XM, Xia XJ, Verry E, Du LL, Pandya A, Petit C, Barkley T, Nance WE, Liu XZ (2002) Mutations in the alternatively spliced exons of USH1C cause non-syndromic recessive deafness. Hum Genet 111: 26 – 30

Overlack N, Klicik D, Bauss K, Markier T, Kremer H, van Wijk E, Wolfrum U (2011) Direct interaction of the Usher syndrome 1C protein SANSS and myomoglobin in the retina. Biochim Biophys Acta 1813: 1883 – 1892

Pans B, Askew C, Galvin A, Heman-Ackah S, Asai Y, Indzhikulian AA, Jodelka FM, Hastings ML, Lentz JJ, Vandenberghe LH et al (2017) Gene therapy restores auditory and vestibular function in a mouse model of Usher syndrome type 1c. Nat Biotechnol 35: 264 – 272

Petters RM, Alexander CA, Wells KD, Collins EB, Sommer JR, Blanton MR, Pier GB, Grout M, Zaidi T, Meluleni G, Mueschenborn SS, Banting G, Ratcliff R, Pan B, Askew C, Galvin A, Heman-Ackah S, Asai Y, Indzhikulian AA, Jodelka FM, Hastings ML, Lentz JJ, Vandenberghe LH et al (2017) Gene therapy restores auditory and vestibular function in a mouse model of Usher syndrome type 1c. Nat Biotechnol 35: 264 – 272

Petters RM, Alexander CA, Wells KD, Collins EB, Sommer JR, Blanton MR, Rojas C, Hoo Y, Flowers WL, Banin E et al (1997) Genetically engineered large animal model for studying cone photoreceptor survival and degeneration in retinitis pigmentosa. Nat Biotechnol 15: 956 – 970

Piel FB, Patil AP, Howes RE, Nyanngi OA, Gethin PW, Williams TN, Weatherall DJ, Hays S (2011) Global distribution of the sickle cell gene and geographical confirmation of the malaria hypothesis. Nat Commun 1: 104

Pier GB, Grout M, Zaidi T, Meluleni G, Mueschenborn SS, Banting G, Ratcliff R, Evans MJ, Collodje WH (1998) Salmonella typhi uses CFTR to enter intestinal epithelial cells. Nature 393: 79 – 82

Regensburger AP, Fonteyne LM, Jüngst J, Wagner AL, Gerhalter T, Nagel AM, Heiss R, Fienkenthaler F, Quraishi M, Neurath MF et al (2019) Detection of collagen by multispectral optoacoustic tomography as an imaging biomarker for Duchenne muscular dystrophy. Nat Med 25: 1905 – 1915

Reiners J, Reidel B, El-Amraoui A, Boed A, Huber I, Petit C, Wolfrum U (2003) Differential distribution of harmonin isoforms and their possible role in Usher-1 protein complexes in mammalian photoreceptor cells. Invest Ophthalmol Vis Sci 44: 5006 – 5015

Reiners J, van Wijk E, Márker T, Zimmermann U, Jürgens K, te Brinke H, Overlack N, Roepman R, Knipper M, Kremer H et al (2005) Scaffold protein harmonin (USH1C) provides molecular links between Usher syndrome type 1 and type 2. Hum Mol Genet 14: 3933 – 3943

Reiners J and Wolfrum U (2006) Molecular analysis of the supramolecular usher protein complex in the retina. Harmonin as the key protein of the Usher syndrome. Adv Exp Med Biol 572: 349 – 353

Richardson GP, de Monvel JB, Petit C (2011) How the genetics of deafness illuminates auditory physiology. Annu Rev Physiol 73: 311 – 334

Richter A, Kurome M, Kessler B, Zakhartchenko V, Klymiuk N, Nagashima H, Wolf E, Wuenisch A (2012) Potential of primary kidney cells for somatic cell nuclear transfer mediated transgenesis in pig. BMC Biotechnol 12: 84

Ross JW, Fernandez de Castro JP, Zhao J, Samuel M, Walters E, Rios C, Bray-Richardson GP, de Monvel JB, Petit C (2012) Generation of an inbred miniature pig model of retinitis pigmentosa. Invest Ophthalmol Vis Sci 53: 501 – 507

Russell S, Bennett J, Wellman JA, Chung DC, Yu Z-F, Tillman A, Wittes J, Pappas J, Elci O, McCague S et al (2017) Efficacy and safety of voretigene neparvovec (AAV-5010) in patients with RPE65-mediated inherited retinal dystrophy: a randomised, controlled, open-label, phase 3 trial. Lancet 390: 849 – 860

Sahl I, Dufour E, Sichtroma C, Michel V, Bahliou A, Perfettini I, Pepermans E, Esteve A, Carette D, Aghaie A et al (2012) Localization of Usher 1 proteins to the photoreceptor cilicary processes, which are absent from mice. J Cell Biol 199: 381 – 399

Savitzky A, Golay MJ (1964) Smoothing and differentiation of data by simplified least squares procedures. Anal Chem 36: 1627 – 1639

Sichtroma C, Parain K, Estivalet A, Aghaie A, Boutet de Monvel J, Picaud S, Sahel J-A, Perron M, El-Amraoui A, Petit C (2011) Usher syndrome type 1– associated cadherins shape the photoreceptor outer segment. J Cell Biol 216: 1849 – 1864

Schindelin J, Rueden CT, Hiner MC, Eliceiri KW (2015) The ImageJ ecosystem: an open platform for biomedical image analysis. Mol Reprod Dev 82: 518 – 529

Sedmak T, Wolfrum U (2010) Intraflagellar transport molecules in ciliary and nonciliary cells of the retina. J Cell Biol 189: 171 – 186

Sedmak T, Wolfrum U (2011) Intraflagellar transport proteins in ciliogenesis of photoreceptor cells. Biol Cell 103: 449 – 466

Sjostrand FS (1953) The ultrastructure of the outer segments of rods and cones of the eye as revealed by the electron microscope. J Cell Comp Physiol 42: 15 – 44

Sommier JR, Estrada JL, Collins EB, Bedell M, Alexander CA, Yang Z, Hughes G, Mir B, Gilger BC, Grob S et al (2013) Production of ELOVL4 transgenic pigs: a large animal model for Stargardt-like macular degeneration. Br J Ophthalmol 95: 1749 – 1754

Spencer WJ, Lewis TR, Pearring JN, Arshavsky VY (2020) Photoreceptor discs: built like ectosomes. Trends Cell Biol 30: 904 – 915

Stingl K, Korubenchak A, Hahn G, Kernstock C, Hipp S, Zobor D, Kohl S, Bonnet C, Mohand-Saïd S, Audo I et al (2019) Full-field electroretinography, visual acuity and visual fields in Usher syndrome: a multicentre European study. Doc Ophthalmol 139: 151 – 160

Thurman RE, Ryens E, Humbert R, Vierstra J, Mauroman T, Haugen E, Sheffield NC, Stegachis AB, Wang H, Vernet B et al (2012) The accessible chromatin landscape of the human genome. Nature 489: 75 – 82

Toms M, Pagarkar W, Moosajee M (2020) Usher syndrome: clinical features, molecular genetics and advancing therapeutics. Ther Adv Ophthalmol 12: 2515841420952194

Vandenberghe LH, Bell P, Maguire AM, Xiao R, Hopkins TB, Maguire AM, Xiao R, Hopkins TB, Grant R, Bennett J, Wilson JM (2013) AAV9 targets cone photoreceptors in the nonhuman primate retina. PLoS One 8: e53463

Velini S, Lazar CH, Chang B, Sieving PA, Banin E, Swaroop A (2015) Biology and therapy of inherited retinal degenerative disease: insights from mouse models. Dis Model Mech 8: 109 – 129

Verpy E, Leibovici M, Zwaenepoel I, Liu X-Z, Gal A, Salem N, Mansour A, Blanchard S, Kobayashi I, Keats BJ et al (2000) A defect in a PDZ domain-containing protein expressed in the inner ear sensory hair cells, underlies Usher syndrome type 1C. Nat Genet 26: 51 – 55

Vochkovka P, Simmet K, Jemiller EM, Wunsch A, Klymiuk N (2019) Gene editing in primary cells of cattle and pig. Methods Mol Biol 1619: 271 – 289

Wang J, Zhuang J, Iyer S, Lin XY, Greven MC, Kim BH, Moore J, Pierce BG, Dong X, Virgil D et al (2013) Factorbook.org: a Wiki-based database for transcription factor-binding data generated by the ENCODE consortium. Nucleic Acids Res 41: D171–D176

Wang K, Fernandez de Castro J, Vukmanic E, Zhou L, Emery D, Demarco PJ, Kaplan HJ, Dean DC (2011) Selective rod degeneration and partial cone inactivation characterize an iodoacetatic acid model of Swine retinal degeneration. Invest Ophthalmol Vis Sci 52: 7917 – 7923

Warming S, Costantino N, Court DL, Jenkins NA, Copeland NG (2005) Simple and highly efficient BAC recombineering using galK selection. Nucleic Acids Res 33: e36
Waterhouse AM, Procter JB, Martin DM, Clamp M, Barton GJ (2009) Jalview Version 2—a multiple sequence alignment editor and analysis workbench. *Bioinformatics* 25: 1189–1191

Williams DS (2008) Usher syndrome: animal models, retinal function of Usher proteins, and prospects for gene therapy. *Vision Res* 48: 433–441

Williams DS, Aleman TS, Lillo C, Lopes VS, Hughes LC, Stone EM, Jacobson SG (2009a) Harmonin in the murine retina and the retinal phenotypes of Ush1c-mutant mice and human USH1C. *Invest Ophthalmol Vis Sci* 50: 3881–3889

Williams DS, Aleman TS, Lillo C, Lopes VS, Hughes LC, Stone EM, Jacobson SG (2009b) Harmonin in the murine retina and the retinal phenotypes of Ush1c-mutant mice and human USH1C. *Invest Ophthalmol Vis Sci* 50: 3881–3889

Wolfrum U, Coldmann T, Overlack N, Mueller C, Vetter JM, Nagel-Wolfrum K (2010) Subcellular localization of usher syndrome proteins in the human retina. *Invest Ophthalmol Vis Sci* 51: 2494

Wolfrum U, Schmitt A (2000) Rhodopsin transport in the membrane of the connecting cilium of mammalian photoreceptor cells. *Cell Motil Cytoskeleton* 46: 95–107

Zele AJ, Cao D (2014) Vision under mesopic and scotopic illumination. *Front Psychol* 5: 1594

Zwaenepoel I, Verpy E, Blanchard S, Meins M, Apfelstedt-Sylla E, Gal A, Petit C (2001) Identification of three novel mutations in the USH1C gene and detection of thirty-one polymorphisms used for haplotype analysis. *Hum Mutat* 17: 34–41

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