NHE8 Is Essential for RPE Cell Polarity and Photoreceptor Survival

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A new N-ethyl-N-nitrosourea (ENU)-induced mouse recessive mutation, identified by fundus examination of the eye, develops depigmented patches, indicating retinal disorder. Histology data show aberrant retinal pigment epithelium (RPE) and late-onset photoreceptor cell loss in the mutant retina. Chromosomal mapping and DNA sequencing reveal a point mutation (T to A) of the Slc9a8 gene, resulting in mutant sodium/proton exchanger 8 (NHE8)-M120K protein. The lysine substitution decreases the probability of forming the 3rd transmembrane helix, which impairs the pore structure of the Na⁺/H⁺ exchanger. Various RPE defects, including mislocalization of the apical marker ezrin, and disrupted apical microvilli and basal infoldings are observed in mutant mice. We have further generated NHE8 knockout mice and confirmed similar phenotypes, including abnormal RPE cells and late-onset photoreceptor cell loss. Both in vivo and in vitro data indicate that NHE8 co-localizes with ER, Golgi and intracellular vesicles in RPE cells. Thus, NHE8 function is necessary for the survival of photoreceptor cells and NHE8 is important for RPE cell polarity and function. Dysfunctional RPE may ultimately lead to photoreceptor cell death in the NHE8 mutants. Further studies will be needed to elucidate whether or not NHE8 regulates pH homeostasis in the protein secretory pathways of RPE.

Vision depends on the photoreceptor cells of the retina to catch photons to start the visual process¹. The retinal pigment epithelium (RPE), adjoining the outer segments (OS) of photoreceptor cells in the back of the retina, is essential for maintaining the homeostasis and survival of photoreceptor cells²,³. Photoreceptor cell death in age-related macular degeneration (AMD) is the leading cause of untreatable blindness in industrialized countries⁴,⁵. The fundamental molecular and cellular mechanisms that regulate and maintain life-long function of the RPE and photoreceptor cells are still not well understood⁶. Understanding these mechanisms is critical for elucidating the events that trigger pathological cascades towards photoreceptor cell death, which occurs in human eye diseases such as retinitis pigmentosa (RP) and AMD⁵,⁷,⁸.

RPE is a monolayer of polarized cells that plays many essential roles in the maintenance and homeostasis of photoreceptor cells⁹. The apical ends of RPE abut and engulf photoreceptor OS through phagocytosis whereas the basolateral sides lie on Bruch’s membrane and transport nutrients, digested metabolic wastes, ions and water between the retina and the choroidal vasculature in the back of the eye. RPE provides the recycle of retinoids for the phototransduction pathway as well as the blood-retinal barriers. Due to phototoxicity, the daily renewal of photoreceptor OS is profoundly important to the survival of photoreceptors and maintenance of retinal health⁸. In the last decade, significant progress has been made in understanding some of the mechanisms that control RPE phagocytosis, especially OS recognition and engulfment signaling molecules including the Gas6/MerTK pathway⁵,¹¹. Late stage of phagocytosis is the recycling and degradation mediated by the endosome/lysosome pathway¹²,¹³; the underlying mechanisms that determine the recycling and degradation of proteins in RPE are still not well understood¹²,¹⁴. Specific markers for polarized RPE cells have been identified for their unique functions in the RPE¹⁵. However, mechanisms that control and maintain RPE polarity and function also require further investigation⁶,¹⁵. In mice, photoreceptor cells start to form OS at postnatal day 10¹⁶; RPE establishes cell polarity at birth and phagocytosis occurs at around the age of two weeks when mice open eyes¹⁷,¹⁸.

From a forward genetic study of new retinal degeneration mouse mutations, we have found that mutations of the Slc9a8 gene, which encodes the sodium/proton exchanger 8 (NHE8), lead to slow photoreceptor cell death. NHE8 is a member of the solute carrier family 9 (SLC9). Sodium/proton exchangers (NHEs) are a large group of monovalent cation/proton antiporters that predominately move Na⁺ in exchange for H⁺, and are involved in diverse physiological processes including the regulation of intracellular pH, absorption of sodium into epithelia, salt tolerance, cell volume, cell adhesion, cell proliferation, organelle biogenesis, and protein trafficking⁹. NHEs can be divided into two subfamilies based on protein cellular localization and sequence alignments; one is a cell
surface subfamily, including mammalian NHEs 1–5, and the other is an intracellular subfamily localized to organelle membranes, including mammalian NHEs 6–9\(^{21,22}\). NHE8 belongs to the intracellular subfamily, and previous studies have reported that NHE8 is expressed on the apical membrane of intestinal epithelial cells to prevent infectious bacterial adherence\(^{22}\). Our current work suggests that NHE8 is essential for the survival of photoreceptor cells and plays an important role in the retina by regulating RPE cell polarity and function.

**Results**

The \(r15\) mutation, identified from a fundus screen of ENU-induced mutant mice, displayed a recessive retinal degeneration\(^{23}\). At the age of 4 weeks, compared to the normal fundus image of heterozygous mice (\(r15/\+)
(1a), homozygous mutant mice (\(r15/r15\)) showed depigmented patches (Fig. 1b). More severe depigmentation was observed in aged mutant mice (Fig. 1c). Histological sections revealed that the number of outer nuclear layers (ONL) in heterozygous mice was around 10–12 without noticeable changes from the ages of 4 weeks to 18 months (Fig. 1d). In contrast, 4-week old homozygous mutant mice displayed aberrant RPE cells with vacuole-like structures while the number of ONL remained comparable to the control at about 10–12 layers (Fig. 1e). The loss of photoreceptor cells was observed in homozygous mutant mice only after the age of 6 weeks. The ONL decreased to 6–8 layers in 6-month old homozygous mutant mice (data not shown) and about 3–4 layers in 18-month old homozygous mutant mice, which also had pigmented aberrant cells submersed in the subretinal space (Fig. 1g). We measured the ONL thickness of the heterozygous and homozygous mutant retinas from mice at the ages of 4 weeks, 6 months and 18 months (Fig. 1h,i,j). At 4 weeks of age, there is no significant difference in ONL thickness between the \(r15\) heterozygous and homozygous mice (\(p > 0.5\)); however, significantly reduced ONL thickness was observed in 6-month old (\(p < 0.05\), except the \(-2\) mm data point) and 18-month old homozygous (\(p < 0.001\)) mutant mice compared to the heterozygous controls. To confirm the pigmented aberrant structures are aberrant RPE cells, immunostaining was performed in 20-month old retinal sections. Compared to normal RPE65 positive RPE cells in the control heterozygous retina (Fig. 1k''), RPE65 positive aberrant pigmented structures were observed in the photoreceptor cell layer in the homozygous mutant (Fig. 1l''). Thus, depigmentation in fundus images likely results from pathological changes of RPE in \(r15\) homozygous mutant retinas; dysfunctional RPE might ultimately contribute to the late-onset photoreceptor loss.

We further investigated the cellular defects of RPE in the \(r15\) mutant mice. Double labeled images of rhodamine-phalloidin and DAPI revealed a typical hexagonal array of RPE cells containing single or double nuclei, with a sharp F-actin network at cell boundaries in the whole mount RPE of a 4-week old wild-type sample (Fig. 2a). In contrast, an age-matched \(r15\) homozygous mutant displayed disorganized RPE cells with irregular shape and size and disrupted F-actin network at some cell boundaries (Fig. 2b); moreover, the \(r15\) mutant RPE flat-mount image was fuzzy, reflecting substantial defects and changes of RPE cells observed by histology analysis (Fig. 1e). Electron microscope images showed abundant microvilli at the apical surface next to organized outer segments of photoreceptors as well as typical basal infoldings next to the Bruch’s membrane in 5-month old wild-type RPE cells (Fig. 2c); however, RPE from 5-month old homozygous \(r15\) mice had neither apical microvilli nor distinct basal infoldings (Fig. 2d). These morphological data indicate that \(r15\) mutant RPE cells lose polarity and lack the distinct structures and organization that are needed for performing functions such as phagocytosis. Therefore, dysfunctional RPE likely causes the late-onset death of photoreceptor cells in \(r15\) mutant mice.

To identify the causative gene, we performed a genome-wide linkage analysis. The \(r15\) mutation was mapped to mouse chromosome 2. Based on linkage data from genomic DNA samples of 177 meioses, we further mapped this mutation into a 6 Mb interval region between the markers D2Mit51 and D2Mit229 on the Ensembl Mouse Genome Server (Fig. 3a). Sequencing data revealed that the \(r15\) retinal phenotype was correlated with a missense mutation (T to A) of the \(Slc9a8\) gene, which results in substitution of methionine (M) by lysine (K) at residue 120 of the \(Slc9a8\) protein (Fig. 3b), mutant \(Slc9a8\)-M120K. The probability of forming transmembrane helices predicted by the program TMHMM (v. 1.0) (http://www.cbs.dtu.dk/) suggested that wild-type \(Slc9a8\) contained 11 transmembrane helices (Fig. 3c) but mutant \(Slc9a8\)-M120K abolished the 3rd transmembrane helix (Fig. 3d). Thus, mutant \(Slc9a8\)-M120K with a disrupted 3rd transmembrane helix likely impairs its function as a sodium-proton exchanger in the RPE. According to standard genetic nomenclature guidelines, the \(r15\) mouse mutant line should be named \(Slc9a8^{r15/M120K}/r15^{r15/M120K}\). To be consistent with previous studies of sodium-hydrogen exchangers in the literature, we use \(Slc9a8\)-M120K in the paper to describe this mutant mouse line.

To confirm that a dysfunctional \(Slc9a8\)-M120K is the sole cause for defective RPE and photoreceptor cell death rather than any other unknown mutation occurring in the ENU-induced \(r15\) mutant line, we generated \(Slc9a8\) null mutant mice (\(Slc9a8^{-/-}\)) from \(Slc9a8\) knockout ES cell clones made by the International Knockout Knockout Mouse Consortium (www.mousephenotype.org/martsearch_lkmc_project/about/eucomm). The \(Slc9a8\) gene targeting vector was constructed using the knockout-first-reporter tagged insertion (promotorless cassette) strategy. Insertion of a promotorless cassette into the intron between exons 3 and 4 of the \(Slc9a8\) gene would result in a truncated \(Slc9a8\) fusion protein, which expresses only the N-terminal 92 amino acids (encoded by the first three exons) followed by the inserted LacZ+ neo in the \(Slc9a8\) knockout mice.

Similar to the recessive retinal phenotypes in \(r15\) (\(Slc9a8^{-/-}\)) homozygous mutant mice, \(Slc9a8\) knockout mice displayed depigmented patches in the fundus photo (Fig. 4b), disrupted RPE layer with vacuoles (Fig. 4d,f), irregularly sized and shaped RPE cells (Fig. 4j), and an obvious loss of photoreceptor cells at the age of 6 months but not 4 weeks (Fig. 4d,f). Heterozygous \(Slc9a8^{+/-}\) mice had no obvious retinal phenotypes (data not shown). The ONL thickness was measured in wild-type (WT) and \(Slc9a8^{-/-}\) mutant retinas at the ages of 4 weeks and 6 months (Fig. 4g,h). At 4 weeks of age, there is no significant difference in ONL thickness between the control and \(Slc9a8^{-/-}\) mice (\(p > 0.5\)); 6-month old \(Slc9a8^{-/-}\) mice display significantly reduced ONL thickness (\(p < 0.05\)). Since the \(r15\) mutant and \(Slc9a8\) knockout mice display similar recessive retinal phenotypes, \(Slc9a8^{-/-}\) is likely a dysfunctional point mutation.

In order to elucidate the mechanistic role of \(Slc9a8\), we characterized the localization of \(Slc9a8\) proteins in the RPE in vivo. Immunohistological data showed that \(Slc9a8\) proteins displayed an intracellular vesicular expression pattern (Fig. 5a) in RPE cells; \(Slc9a8\) colocalized with giantin (a marker for the Golgi complex) as well as expressed in other intracellular vesicles, but not at RPE cell boundaries (Fig. 5a,b). Tight junctions labeled by ZO-1 antibody (green) were colocalized with F-actin in RPE cells (Fig. 5c).

Since a previous work reported that \(Slc9a8\) played a key role in the control of protein trafficking in cultured HeLa M-cells in vitro\(^{24}\), we investigated whether \(Slc9a8\) played an important role in the protein trafficking needed for RPE cell polarity. We examined the distribution of \(Slc9a8\) delivered by Adeno-associated virus (AAV) in human ARPE-19 cells in vitro. \(Slc9a8\)-SE pFluorin proteins, expressed from AAV5, were predominantly present in vesicular intracellular structures of ARPE-19 cells (Fig. 5d), and were colocalized with the Golgi marker TGN46 (Fig. 5e) and the endoplasmic reticulum (ER) marker calreticulin (Fig. 5f).
Figure 1 | The r15 mutant mice display late-onset recessive retinal degeneration with aberrant RPE cells. Fundus photos from a 4-week old r15 heterozygous mouse (a), a 4-week old r15 homozygous mutant mouse (b) and a 6-month old homozygous mutant mouse (c); retinal veins are indicated by black arrowheads and retinal arteries indicated by white arrows. Toluidine blue stained retinal sections from 4-week old heterozygous (d) and homozygous (e) mutant mice, and 18-month old heterozygous (f) and homozygous (g) mutant mice; white arrows indicate RPE; ONL, outer nuclear layer; INL, inner nuclear layer. Spider graphs show ONL thickness in r15 heterozygous mice (black lines, n = 3) and r15 homozygous mice (grey lines, n = 3) at the ages of 4 weeks (h), 6 months (i), and 18 months (j). Immunostaining of 20-month old r15 heterozygous (k) and homozygous (l) retinal frozen sections indicate that mislocalized pigmented structures are labeled by an anti-RPE65 antibody (red, white arrows); nuclei are labeled with Dapi (blue); differential interference contrast (DIC) merged images (k’ and l’) are underneath the corresponding images; RPE, retinal pigment epithelium; PS, photoreceptor segments; Scale bars for panels d-g and k-l’, 20 μm.
We further examined the expression of ezrin, a suitable apical marker for monitoring RPE cell polarity, even though it could be detected at a very low level in the basal-lateral sides. As expected, ezrin proteins were predominantly located at the apical sides of the RPE in the wild-type mouse retina (Fig. 6a,c,e). However, aberrant lateral-basal distribution of ezrin proteins in RPE was detected as early as two weeks of age in NIH8/- mice (Fig. 6d). Ezrin also appeared in apical, lateral and basal sides of RPE in 4-week old NIH8-M120K (Fig. 6f) and NIH8/- mice. Consistent with previous morphological data (Fig. 1,2,4), RPE cell polarity is disrupted in NIH8-M120K and NIH8/- mutant retinas.

Discussion

This work demonstrates that NHE8 is essential for the survival of photoreceptor cells, and it plays an important role for the polarity and function of RPE cells. Both NHE8 knockout and NIH8-M120K mutant mice display late-onset loss of photoreceptor cells and impaired RPE cells including altered cell morphology, mislocalized ezrin proteins and a loss of apical microvilli. Since RPE cell defects occur before the loss of photoreceptor cells in both NIH8 mutant mouse lines, the slow loss of photoreceptors is likely a consequence of impaired RPE function. Based on the fact that NHE8 proteins are localized in protein trafficking compartments such as the Golgi, ER and other unknown intracellular vesicles in RPE, we hypothesize that NHE8 may regulate pH homeostasis in the protein trafficking pathway and/or in intracellular vesicles. However, further studies will be needed to test this hypothesis. The underlying mechanism for altered distribution of the cell polarity marker ezrin is unknown; it is possible that NHE8 loss-of-function mutations perturb ezrin’s trafficking, sorting and/or targeting to its final destination. A loss of NHE8 function alone is sufficient to impair RPE and leads to the slow loss of photoreceptor cells in mice. The presence of other NHE isoforms and other pH regulators are unable to compensate the functional loss of NHE8 in the retina. It will be important to investigate whether or not NHE8 plays an essential function in humans. These NIH8 mutant mice provide an excellent model system to study intracellular pH homeostasis that is largely unknown in the RPE.

All key functions of the RPE rely on its cell polarity, such as precisely localized secretory receptors for phagocytosis and polarized structures to regulate the transport of nutrients and waste products. Ezrin is an apical marker of the RPE due to its high abundance in apical microvilli, even though very low levels of ezrin are present in basal infoldings of RPE. Ezrin is needed for the proper formation of microvilli and basal infoldings of RPE and for the function of NHE3. Thus, NIH8 mutations will cause many indirect consequences to impair the function of RPE. Transporters such as proton-coupled monocarboxylate transporters (MCTs) are specifically positioned apically or basolaterally to facilitate the transport of lactate and H+ out of the retina to regulate pH, ion homeostasis and the high-energy demands of photoreceptor cells. MCT3 and MCT4 harbor dominant sorting information for the basolateral membrane. It will be interesting to study whether or how MCTs are affected in NIH8 mutations.

The polarized trafficking machinery, composed of secretory organelles (ER and Golgi complex) and endosomal compartments, is critical for the generation and maintenance of the asymmetric distribution of plasma membrane proteins. The trans-Golgi network (TGN) is where cargoes undergo sorting, packaging and delivery to different destinations such as the plasma membrane, the endosomes, or secretory granules. Each of these transport routes probably uses a specialized and dedicated machinery. However, the underlying mechanism for how polarized protein trafficking contributes to RPE plasma membrane polarity is still poorly understood. NIH8 is known to contribute to the maintenance of the unique acidic pH values of the Golgi and post-Golgi compartments in cultured cells. The pH values gradually decrease from ~7 to ~5, starting in ER, Golgi, TGN and secretory granules, and from early and late endosomes (pH ~ 6.5) to lysosomes (pH ~ 4.5) in general. This progressive acidification is essential for post-translational modifications, including sorting of newly synthesized proteins into the secretory pathway. Studies from cultured HeLa M-cells in vitro have reported that NIH8 plays a key role in the control of protein trafficking and endosome morphology. The majority of epitope-tagged NIH8 was found in the trans-Golgi network of HeLa M-cells, but a proportion was also localized to multivesicular bodies (MVBs). Depletion of endogenous NIH8 or overexpression of a nonfunctional point mutant protein NIH8-E225Q affected endosome morphology but did not affect the overall pH inside dense MVBs. Our unpublished data show that endosome marker EEA1 seems not colocalize with NIH8 in RPE cells. So the function of NIH8 in RPE cells likely differs from its role in HeLa M-cells. NIH8 is also known to be an apically expressed membrane protein in intestinal epithelial cells for controlling sodium absorption and bicarbonate secretion in the intestine. However, NIH8 protein is hardly detected in either the apical or basolateral membrane of RPE (Fig. 5a,b). Intracellular pH changes in polarized protein trafficking of RPE have not been investigated in vivo nor in cultured RPE cells in vitro. NIH8 mutants may be useful for exploring pH homeostasis in the polarized trafficking machinery needed for RPE cell polarity as well as the luminal pH regulation in the secretory pathway of RPE during development and aging.

Methods

Animals. All studies and examinations were conducted in accordance with a protocol approved by the Animal Care and Use Committee (ACUC) at University of California, Berkeley. ENU-mutagenesis and breeding of the r15 mice were performed as previously described. NIH8 knockout mice were acquired from the Mutant Mouse Regional Resource Center (MMRRC) at UC Davis. The targeting vector was designed to insert a promoterless cassette into the intron between exons 3 and 4 of the NHE8 gene.
Figure 3 | Identification of the causative gene, Slc9a8, for the r15 mutation. (a) Linkage analysis of the r15 mutation was based on the phenotypes and genotypes of 177 meioses. The mutation was mapped to a 6 Mb region between markers D2Mit51 and D2Mit229 on mouse chromosome 2. Positions of the markers are based on the Ensembl Mouse Genome Server. (b) DNA sequencing data revealed a missense (T to A) mutation of the Slc9a8 gene, resulting in a substitution of methionine (M) by lysine (K) at the 120th amino acid of the NHE8 protein in the r15 mutation. (c) Wild-type NHE8 protein was predicted to form 11 transmembrane helices by the program TMHMM (v. 1.0) (http://www.cbs.dtu.dk/). (d) NHE8-M120K mutant protein decreased the probability of forming the 3rd transmembrane helix containing residue 120.
Figure 4 | NHE8 knockout mice display retinal phenotypes similar to the r15 mutation. Fundus photos of 2-month old wild-type (a) and NHE8-/- (b) littersmates. The NHE8-/- fundus reveals depigmented patches, similar to that observed in the r15 mutant mice. Histological sections of 4-week old wild-type (c) and NHE8-/- (d) littersmates as well as 6-month old wild-type (e) and NHE8-/- (f) littersmates. The NHE8 knockout RPEs displayed vacuoles and abnormal pigmented cellular structures (d,f, indicated by white arrows). At the age of 6 months, the wild-type control had 10–12 outer nuclei layers (ONL) of photoreceptor cells, while NHE8 knockout had only 5–7 layers. Spider graphs show ONL thickness in wild-type (black lines, n = 3) and NHE8-/- mice (grey lines, n = 3) at the ages of 4 weeks (g) and 6 months (h). Phalloidin stained images show typical pentagonal/hexagonal cell shape with intact F-actin network of flat-mount RPE in 1-month-old wild-type mice (i), while variable cell size and abnormal cell shape with disrupted F-actin network of flat-mount RPE in NHE8-/- (j) littersmates. Scale bars for panels c–f, 50 μm; for panels i and j, 20 μm.

The NHE8 protein is associated with the Golgi complex and the intracellular vesicles. Fluorescent staining results reveal that NHE8 proteins (green) display intracellular vesicular-like signals (a), white arrowheads indicate enriched F-actin (red) at apical/lateral tight junctions. NHE8 proteins (green) are partly colocalized with giantin protein (red, a marker for the Golgi complex) in wild-type RPE, indicated by white arrows (b). Tight junctions labeled by anti-ZO-1 antibody (green) are colocalized with F-actin (c). Scale bars for panels a–c, 10 μm. Human ARPE-19 cells were infected with AAV expressing wild-type NHE8-SE pHluorin, and cells were fixed for staining with antibodies for Golgi and ER; NHE8-SE pHluorin expression (green) displays a more prominent perinuclear pattern as well as vesicular distribution throughout the cell (d); staining with anti-TGN46 (red) reveals colocalization of NHE8 and the Golgi complex (e) in the perinuclear region; staining with anti-calreticulin (red) also suggests a colocalization of NHE8 and ER in the cytosol (f). Scale bars for panels d–f, 20 μm.

Slc9a8 gene, which would result in a truncated NHE8 fusion protein that includes the N-terminal peptide (encoded by the first three exons) and the inserted LacZ and neo. For genotyping, a primer pair of forward 5’-GAA GTA GGT CTC TCA CTG G and reverse 5’-AAT GTC TGG CCT CCA CC generates a wild-type PCR fragment (~500 bp) and a larger knockout band (~575 bp) due to the insertion of a loxp site in the targeting vector.

Fundus examination and histology. Fundus examination and retinal histology were performed as previously described41.

Genomic linkage analysis. A female founder was identified from a screen of ENU-induced F3 mutagenized mice by fundus examination, and chromosome mapping was performed according to previously described methods41,42. Homozygous r15 mutant mice in the C57BL/6 strain background were mated with wild-type C3A.BLIA-Fde6b/’/+ mice to produce G1 hybrid mice; G1 hybrids were further mated with homozygous r15 mutant mice to produce second-generation (G2) mice. The G2 mice were examined for retinal phenotype, and genomic DNA samples were extracted from tail snips for genome-wide linkage analysis using a total of 59 microsatellite markers. After the chromosomal linkage was identified, we further performed fine mapping. Marker sequence information (D2Mit51, D2Mit145, D2Mit229, D2Mit344 and D2Mit200) was obtained from the Ensembl Mouse database.

DNA sequencing. Retinas were dissected from homozygous r15 mutant mice and total RNA was isolated using the TRIzol® Reagent (Invitrogen Life Technologies). The Superscript® First-Strand Synthesis System for RT-PCR kit (Invitrogen Life Technologies) was used to synthesize cDNAs. The coding region of the NHE8 gene was amplified by various primer pairs with Platinum® pfX DNA polymerase (Invitrogen Life Technologies). PCR fragments with overlapping regions were sequenced at the UC Berkeley DNA sequencing facility.

Immunofluorescence studies. Immunofluorescence staining of retinal frozen sections was performed as previously described41. For staining RPE whole mount, mouse eyes were fixed in 4% formaldehyde/PBS, the cornea and lens were removed, followed by antibody staining and the remaining eye cups were flat-mounted for imaging. A rabbit polyclonal antibody for NHE8 was generated using the C-terminal 14 amino acid peptide as immunogen; other antibodies used for immunostaining were: RPE65 mouse monoclonal antibody (generously provided by Dr. Debra Thompson, University of Michigan), ezrin rabbit polyclonal antibody (Cell Signaling Technology), ZO-1 rabbit polyclonal antibody (Invitrogen Life Technologies),
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**Figure 6** NHE8 is important for the polarity of ezrin expression in RPE cells. 
Immunostaining of ezrin proteins (green, a marker for the apical side of RPE) and F-actin (red) in wild-type (a) and NHE8-/- retinal sections (b) at the age of postnatal day 7 (P7) show very similar ezrin distribution in the apical RPE as well as some lateral sides. At P14, ezrin distribution is mostly restricted to the apical side of RPE in the wild-type (c), while some extended ezrin expression in the lateral sides towards the basal sides is observed in the NHE8-/- retina (d, white arrows). Panels e–g show the immunodistribution of ezrin proteins in the wild-type (e), r15 mutant (f) and NHE8-/- (g) retinas of mice at the age of 4 weeks. White arrows indicate mislocalized ezrin at basal and lateral sides of r15 and NHE8-/- RPEs. Scale bar, 10 µm. 

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**Author contributions**

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**Additional information**

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