LRIT3 is essential to localize TRPM1 to the dendritic tips of depolarizing bipolar cells and may play a role in cone synapse formation

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Title: LRIT3 is essential to localize TRPM1 to the dendritic tips of depolarizing bipolar cells and may play a role in cone synapse formation

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ABSTRACT

Mutations in LRIT3 lead to complete congenital stationary night blindness (cCSNB). The exact role of LRIT3 in ON-bipolar cell signaling cascade remains to be elucidated. Recently, we have characterized a novel mouse model lacking Lrit3 (no b-wave 6, (Lrit3<sup>nob6/nob6</sup>)), which displays similar abnormalities as patients with cCSNB with LRIT3 mutations. Here we compare the localization of components of the ON-bipolar cell signaling cascade in wild-type and Lrit3<sup>nob6/nob6</sup> retinal sections by immunofluorescence confocal microscopy. An anti-LRIT3 antibody was generated. Immunofluorescent staining of LRIT3 in wild-type mice revealed a specific punctate labeling in the outer plexiform layer (OPL), which was absent in Lrit3<sup>nob6/nob6</sup> mice. LRIT3 did not colocalize with ribeye or calbindin but colocalized with mGluR6. TRPM1 staining was severely decreased at the dendritic tips of all depolarizing bipolar cells in Lrit3<sup>nob6/nob6</sup> mice. mGluR6, GPR179, RGS7, RGS11 and Gβ5 immunofluorescence was absent at the dendritic tips of cone ON-bipolar cells in Lrit3<sup>nob6/nob6</sup> mice, while it was present at the dendritic tips of rod bipolar cells. Furthermore, PNA labeling was severely reduced in the OPL in Lrit3<sup>nob6/nob6</sup> mice. This study confirmed the localization of LRIT3 at the dendritic tips of depolarizing bipolar cells in mouse retina and demonstrated the dependence of TRPM1 localization on the presence of LRIT3. Since tested components of the ON-bipolar cell signaling cascade and PNA revealed disrupted localization, an additional function of LRIT3 in cone synapse formation is suggested. These results point to a possibly different regulation of the mGluR6 signaling cascade between rod and cone ON-bipolar cells.
INTRODUCTION

The visual ON-pathway is initiated at the first retinal synapse between photoreceptors and ON-bipolar cells. In darkness, glutamate released by the photoreceptors binds to the metabotropic glutamate receptor 6 (GRM6/mGluR6) on the ON-bipolar cell dendrites (Nakajima et al., 1993; Nomura et al., 1994). This binding leads to the activation of the heterotrimeric G-protein, Go (Nawy, 1999; Dhingra et al., 2000), which results in the closure of the transient receptor potential melastatin 1 (TRPM1) cation channel (Morgans et al., 2009; Shen et al., 2009; Koike et al., 2010). When the photoreceptors are stimulated by light, glutamate release is reduced, the cascade is deactivated, and TRPM1 channels open, resulting in cell membrane depolarization.

Mutations in several genes disrupt synaptic transmission between photoreceptors and ON-bipolar cells, leading to complete congenital stationary night blindness (cCSNB) (Zeitz, 2007; Zeitz et al., 2015). cCSNB has been associated with mutations in genes encoding proteins localized at the dendritic tips of ON-bipolar cells (Masu et al., 1995; Morgans et al., 2009; Orlandi et al., 2012; Peachey et al., 2012b; Orhan et al., 2013), including nyctalopin, a leucine-rich repeat protein (Morgans et al., 2006). Recently, we have identified mutations in LRIT3, a gene encoding the leucine-rich repeat, immunoglobulin-like and transmembrane domains 3 protein, leading to cCSNB (Zeitz et al., 2013). The corresponding protein localizes at the dendrites of ON-bipolar cells in human (Zeitz et al., 2013).

The exact role of LRIT3 in the mGluR6 signaling cascade remains to be elucidated. It has been shown that nyctalopin binds to TRPM1 (Cao et al., 2011; Pearring et al., 2011) and is essential for its correct localization (Pearring et al., 2011). Membrane transport of proteins such as TRPM1 involves cytoskeletal scaffolding proteins, many of which contain PDZ domains (Feng & Zhang, 2009). Nyctalopin, being mainly an extracellular protein, does not
contain an intracellular PDZ binding domain and is therefore probably not able to bring TRPM1 to the cell surface (Pearring et al., 2011). Interestingly, LRIT3 has a PDZ-binding motif and might fulfill this function (Zeitz et al., 2013).

Mouse models for cCSNB have been helpful in dissecting the visual ON-pathway in bipolar cells (Cao et al., 2011; Pearring et al., 2011; Orlandi et al., 2012; Orlandi et al., 2013). Seven mouse models with defects in four different genes have already been published (Masu et al., 1995; Pardue et al., 1998; Gregg et al., 2003; Pinto et al., 2007; Maddox et al., 2008; Morgans et al., 2009; Shen et al., 2009; Koike et al., 2010; Peachey et al., 2012a; Peachey et al., 2012b). Recently, we have characterized a novel mouse model lacking Lrit3 (no b-wave 6 (Lrit3<sup>nob6/nob6</sup>)), which displays similar abnormalities as patients with cCSNB due to LRIT3 mutations, with lacking or severely reduced b-wave amplitudes in the scotopic and photopic electroretinogram, respectively (Neuille et al., 2014).

Here we describe the localization of LRIT3 in mouse retina and compare the localization of known components of the mGluR6 signaling cascade in wild-type and Lrit3<sup>nob6/nob6</sup> retinas to better understand the function of LRIT3.
MATERIALS AND METHODS

Ethics statements

All animal procedures were performed according with the Council Directive 2010/63EU of the European Parliament and the Council of 22 September 2010 on the protection of animals used for scientific purposes and were approved by the French Minister of Agriculture (authorization A-75-1863 delivered on 09th November 2011). All efforts were made to minimize suffering.

Animal Care

The generation and characterization of the Lrit3 knock-out mouse has been described elsewhere (Neuille et al., 2014) (http://www.taconic.com/knockout-mouse/lrit3/tf2034_--model_lrit3). Ten wild-type (Lrit3+/+) and seven mutant (Lrit3nob6/nob6) six to seven-week-old male and female mice were used in this study. Mice were housed in a temperature-controlled room with a 12-h light/12-h dark cycle. Fresh water and rodent diet were available ad libitum.

LRIT3 antibody

To obtain an antibody to mouse LRIT3, the following procedure was performed by a company (Eurogentec, Seraing, Belgium). Two peptides corresponding to amino acids 180–195 and 613-628 of mouse LRIT3 (NP_001274153.1; peptide sequences: AVTPSRSPDFPPRII and CTSKFWEEDLSKETY, respectively) were synthesized and coupled to keyhole limpet haemocyanin (KLH). Two adult New Zealand White rabbits were immunized with both peptide-KLH conjugates. The rabbits received three more immunizations seven, ten and eighteen days later. Immune sera were collected ten days after the final immunization and stored at -20°C. We performed immunohistochemistry on Lrit3+/+ and Lrit3nob6/nob6 retinal sections with these two sera. One of them produced a specific
punctate signal in the outer plexiform layer of the Lrit3+/ retinal sections that was absent on Lrit3nob6/nob6 sections. However, strong non-specific signal and noise were also present on the whole Lrit3+/ and Lrit3nob6/nob6 sections (data not shown). In order to determine which of the two peptides resulted in the specific staining, we performed immunohistochemistry on Lrit3+/ retinal sections after preincubation of the serum with either the N- or the C-terminal peptide. The specific signal was absent when the serum was preincubated with the peptide localized at the N-terminus but noise remained (data not shown). Subsequently, affinity purification was performed with this peptide to decrease noise and to obtain a more specific antibody.

**Preparation of retinal sections for immunohistochemistry**

Mice were killed by CO₂ administration and cervical dislocation. Eyes were removed and prepared following three methods. For method 1, we made two slits in a cross within the cornea and placed the eyeball in ice cold 4% (w/v) paraformaldehyde in 0.12 M phosphate buffer, pH 7.2 for 1 hour. After three 10-min-washes with ice cold phosphate buffer saline (PBS), we transferred the eyeball to cold 30% sucrose. Finally, the lens was removed and the eyecup was embedded in OCT (Sakura Finetek, AJ Alphen aan den Rijn, The Netherlands) and frozen in a dry ice-cooled bath of isopentane. For method 2, the anterior segment and lens were removed and the eyecup was fixed in ice cold 4% (w/v) paraformaldehyde in 0.12 M phosphate buffer, pH 7.2 for 20 min. The eyecup was washed three times in ice-cold PBS and cryoprotected with increasing concentrations of ice cold sucrose in 0.12 M phosphate buffer, pH 7.2 (10%, 20% for 1 h each and 30% overnight). Finally, the eyecup was embedded in 7.5% gelatin-10% sucrose and frozen in a dry ice-cooled bath of isopentane. For method 3, we made a hole just behind the ora serrata and placed the eyeball in 4% (w/v) paraformaldehyde in 0.12 M phosphate buffer, pH 7.2 for 5 min. We then removed the lens and the eyecup was again fixed for 20 min in paraformaldehyde at room temperature. The
eyecup was washed three times in PBS and cryoprotected with increasing concentrations of ice cold sucrose in 0.12 M phosphate buffer, pH 7.2 (10% for 1 h and 30% overnight). Finally, the eyecup was embedded in 7.5% gelatin-10% sucrose and frozen in a dry ice-cooled isopentane bath. Sections were cut at a thickness of 18 µm on a cryostat and mounted onto glass slides (Super-Frost, Thermo Fisher Scientific, Waltham, MA, USA). The slides were air dried and stored at -80°C.

**Immunostaining of retinal cryosections**

Primary antibodies used for immunostaining are listed in Table 1. The TRPM1 immunoreactive serum from a patient suffering from melanoma-associated retinopathy (MAR), the rat mGluR6 antiserum raised in sheep, the purified goat polyclonal antibody to mouse Gβ5 and the mouse RGS11 antibody raised in rabbit were used as previously described (Chen et al., 2003; Morgans et al., 2006; Morgans et al., 2007; Xiong et al., 2013). Immunohistochemistry on retinal sections was performed following two protocols depending on the section preparation. For eyecups embedded in OCT, we used a previously published protocol (Morgans et al., 2006). Briefly, eyecup sections were blocked by incubation with antibody incubation solution (AIS) (3% (v/v) normal horse serum, 0.5% (v/v) Triton X-100, 0.025% (w/v) NaN₃ in PBS) at room temperature for 60 min. Subsequently, the sections were incubated with primary antibodies in AIS for 1 h at room temperature. After washing in PBS, the sections were incubated with secondary antibodies coupled to Alexa Fluor 488 or Alexa Fluor 594 (Life Technologies, Grand Island, NY, USA) at a dilution of 1:1000 in PBS for 1 h at room temperature. The slides were stained with 4′,6-diamidino-2-phenylindole (DAPI) and subsequently cover-slipped with mounting medium (Aqua-Mount mounting medium, Thermo Fisher Scientific, Waltham, MA, USA). For eyecups embedded in gelatin-sucrose, sections were blocked by incubation at room temperature for 60 min in 10% (v/v) donkey serum,
0.3% (v/v) Triton X-100 in PBS. Subsequently, the sections were incubated with primary antibodies in blocking solution overnight at room temperature. After washing in PBS, the sections were incubated with secondary antibodies coupled to Alexa Fluor 488 or Cy3 (Jackson ImmunoResearch, West Grove, PA, USA) at a dilution of 1:1000 in PBS for 1.5 h at room temperature. The slides were stained with DAPI and subsequently cover-slipped with mounting medium (Mowiol, Merck Millipore, Billerica, MA, USA). None of the secondary antibodies used gave recordable signal when used without primary antibodies (data not shown).

**Image acquisition and quantification**

Fluorescent staining signals were captured with a confocal microscope (FV1000, Olympus, Hamburg, Germany) equipped with 405, 488, and 559 nm lasers.

For localization studies as well as for quantification of LRIT3, confocal images were acquired with a 40x objective compatible with oil (lens NA: 1.3) imaging pixels of 310 nm and 77 nm in width and height for zoom 1 and 4, respectively, and using a 0.52 µm step size. For localization, each image corresponds to the projection of three optical sections. For figures, brightness and contrast were optimized (ImageJ, version 1.49; National Institutes of Health, Bethesda, MD, USA). The method for quantification of LRIT3 on 4 independent $Lrit3^{+/-}$ retinal sections was adapted from a previously published protocol (Ramakrishnan *et al.*, 2014), using Imaris (Bitplane, Zurich, Switzerland) and is illustrated in Figure S1. Optical sections centered on the outer plexiform layer (OPL) were stacked to obtain a 3D reconstruction of the entire retinal section thickness at this level. An initial region of interest (ROI) was drawn around the OPL (Figure S1A and S1B). We used peanut agglutinin (PNA) staining as a marker to define the ROI of cone ON-bipolar cell dendritic tips (Figure S1C). The mean intensity per voxel of LRIT3 staining at the dendritic tips of cone ON-bipolar cells...
was obtained by quantifying LRIT3 staining enclosed within this region. The ROI corresponding to LRIT3 staining at the dendritic tips of rod bipolar cells was formed by drawing spheres of 2 µm radius around LRIT3-labeled puncta that are not associated with PNA (Figure S1D). The mean intensity per voxel of LRIT3 staining at the dendritic tips of rod bipolar cells was then measured. Finally, these two intensities were compared in the same retinal section.

For quantification of PNA, confocal images were acquired with a 20x objective compatible with oil (lens NA: 0.85) imaging pixels of 621 nm in width and height, and using a 1.55 µm step size. We reconstructed three Lrit3+/+ and three Lrit3noba6/noba6 retinas by assembling multiple confocal images of the same retina. For each Lrit3+/+ mouse, a Lrit3noba6/noba6 mouse of the same breeding pair was chosen and their retinas were processed in parallel and in the same way. Quantification of PNA in the OPL and in inner and outer segments (IS/OS) was adapted from a previously published protocol (Xu et al., 2012) and was performed with ImageJ (version 1.49). The background value was defined as the mean intensity per pixel in a ROI comprising the inner nuclear layer (INL) in which no PNA staining has been described. This value was then subtracted from the measurements of OPL and IS/OS intensities to give corrected measurements. For quantification of PNA in the OPL, the corrected mean intensity per pixel was measured for a segmented line of 30 pixels thickness drawn along the OPL. To obtain the number of PNA-labeled synaptic clefts between cones and corresponding cone ON-bipolar cells per mm of OPL (density of particles), the line was straightened and a threshold applied to isolate each labeled structure. The same threshold was applied for the wild-type and the mutant retinas of the same pair. We then counted the number of labeled structures with a size between 4 and 45 µm² and divided this value by the length of the selected line. The corrected mean intensity per pixel and the density of particles were compared between wild-type and mutant mice in each pair. To quantify the PNA staining in the IS/OS compartment,
the corrected mean intensity per pixel was measured in a ROI surrounding these compartments. The values obtained were compared between $Lrit3^{+/+}$ and $Lrit3^{nob6/nob6}$ mice of each pair.

**Preparation of retinal lysates and Western blotting**

Retinas were prepared according to a previously published protocol (Cao et al., 2012). Mice were killed by CO$_2$ administration and cervical dislocation. Whole retinas were removed from 3 $Lrit3^{+/+}$ and 3 $Lrit3^{nob6/nob6}$ mice and lysed by sonication in ice-cold PBS supplemented with 150 mM NaCl, 1% Triton X-100, and protease inhibitor (complete protease inhibitor tablets, Roche, Meylan, France). Supernatants were collected after a 15 min centrifugation step at 21,000 g at 4°C and total protein concentration was measured by using a bicinchoninic acid protein assay kit (BCA Protein Assay Kit, Thermo Fisher Scientific, Rockford, IL, USA). SDS sample buffer (pH 6.8) containing 8 M urea was added to the supernatants and 8.5 µg of protein were subjected to 4-15% SDS-PAGE (Bio-Rad Laboratories, Hercules, CA, USA). Protein bands were transferred onto nitrocellulose membranes (GE Healthcare, Little Chalfont, UK). Membranes were blocked for 1 hour at room temperature in 5% dry milk diluted in PBS, 0.1% Tween 20 and subjected to primary antibodies in 1% dry milk diluted in PBS, 0.1% Tween 20 overnight at 4°C. After washing in PBS, 0.1% Tween 20, the membranes were incubated with HRP-conjugated secondary antibodies (Jackson ImmunoResearch) in 1% dry milk diluted in PBS, 0.1% Tween 20, and subsequently bands were detected by using a chemiluminescent substrate (ECL2, Thermo Fisher Scientific). Primary antibodies were mouse anti TRPM1 antibody raised in sheep diluted at 1:1000 (Cao et al., 2011) and anti β-actin antibody raised in mouse diluted at 1:10000 (MAB1501, Merck Millipore, Darmstadt, Germany). Bands intensities were quantified by using ImageJ (version 1.49). For each lane, TRPM1 staining intensity was normalized to β-actin staining intensity
and compared between 3 *Lrit3*+/+ and 3 *Lrit3*<sup>nob6/nob6</sup> mice. The experiment was repeated 7 times.

**Statistical analyses**

Statistical analyses were performed using SPSS Statistics (version 19.0, IBM, Armonk, NY, USA). A Student’s t-test was used to compare TRPM1 protein amounts between *Lrit3*+/+ and *Lrit3*<sup>nob6/nob6</sup> mice, the intensity of LRIT3 staining at the dendritic tips of rod bipolar cells versus cone ON-bipolar cells in *Lrit3*+/+ mouse as well as PNA staining in the OPL and in the IS/OS complex between *Lrit3*+/+ and *Lrit3*<sup>nob6/nob6</sup> mice. Tests were considered as significant when p<0.05.
RESULTS

Validation of the LRIT3 antibody

Immunofluorescent staining of LRIT3 on Lrit3\(^{+/+}\) mouse retinal sections revealed punctate labeling in the OPL (Figure 1A, two pictures on the left and Figure 1B left). On Lrit3\(^{nob6/nob6}\) mouse retinal sections, this punctate labeling was absent, demonstrating the specificity of the LRIT3 labeling (Figure 1A, two pictures on the right and Figure 1B, right). Some non-specific signal remained in the inner plexiform and the ganglion cell layers (IPL and GCL, respectively) in both Lrit3\(^{+/+}\) and Lrit3\(^{nob6/nob6}\) mice (Figure 1A, LRIT3 staining alone). The specific staining in the OPL most likely represents the dendritic tips of rod bipolar cells (arrows, Figure 1B, left) and cone ON-bipolar cells (arrowheads, Figure 1B, left).

LRIT3 localizes at the dendritic tips of depolarizing bipolar cells in the mouse retina

In order to precisely localize LRIT3 in mouse retina, and particularly in the OPL, we performed co-immunolocalization studies with the anti-LRIT3 antibody and antibodies against ribeye, a component of the photoreceptor synaptic ribbon (Schmitz et al., 2000; tom Dieck et al., 2005), calbindin, which labels the post-synaptic processes of horizontal cells (Haverkamp & Wassle, 2000), PKCa to label rod bipolar cells (Negishi et al., 1988; Greferath et al., 1990) and mGluR6 to label dendritic tips of ON-bipolar cells (Vardi et al., 2002; Morgans et al., 2007; Cao et al., 2008).

The LRIT3 puncta did not colocalize with ribeye but were nested within the arcs of the synaptic ribbons (Figure 2A). LRIT3 staining was also adjacent to calbindin staining but without colocalization (Figure 2B). Thus, LRIT3 was present in the OPL but did not localize presynaptically and was absent in the horizontal cells. PKCa was present in both the dendritic tips as well as in the cell bodies of rod bipolar cells, whereas LRIT3 staining was solely
concentrated in the OPL (Figure 2C). Strong colocalization of LRIT3 and mGluR6 confirmed that LRIT3 is present at the dendritic tips of rod and cone ON-bipolar cells (Figure 2D).

**TRPM1 localization is disrupted at the dendritic tips of ON-bipolar cells in Lrit3nob6/nob6 retina**

In order to test our hypothesis that LRIT3 is required for the correct localization of TRPM1, we performed immunohistochemistry on Lrit3+/+ and Lrit3nob6/nob6 mouse retinal sections with serum from a patient suffering from melanoma-associated retinopathy (MAR) that was previously demonstrated to show specific TRPM1 immunoreactivity on mouse retinal sections (Xiong et al., 2013). In Lrit3+/+ retinal sections, the serum revealed punctate labeling in the OPL (Figure 3A and 3C, left) that colocalized with mGluR6 staining (Figure 3B and 3C), confirming the localization of TRPM1 to the dendritic tips of rod bipolar cells (arrows, Figure 3B and 3C, left) and cone ON-bipolar cells (arrowheads, Figure 3B and 3C, left). We also observed immunofluorescent labeling of bipolar cell bodies (asterisks) in the INL (Figure 3A and 3C, left). In Lrit3nob6/nob6 mice, the intensity of the TRPM1 puncta co-localizing with mGluR6 was dramatically decreased, but somatic staining (asterisks) seemed unchanged (Figure 3A and 3C, right), suggesting that TRPM1 localization at the dendritic tips of ON-bipolar cells is dependent on LRIT3.

To investigate whether the decrease in TRPM1 localization at the dendritic tips of depolarizing bipolar cells is also reflected by reduction of total amount of TRPM1 protein, we compared TRPM1 protein levels in Lrit3+/+ and Lrit3nob6/nob6 retinas by Western blot analysis. TRPM1 protein was still present in the retina of Lrit3nob6/nob6 mice and the protein amount was not significantly altered (Figure 3D and 3E). These findings suggest that elimination of LRIT3 results in a severe reduction of TRPM1 at the dendritic tips of all ON-bipolar cells but does not significantly affect the total TRPM1 protein amount.
Other components of the ON-bipolar signaling cascade are mislocalized in \( \text{Lrit}^{\text{nob6/nob6}} \) retina

Interestingly, mGluR6 staining in \( \text{Lrit}^{\text{nob6/nob6}} \) retina showed an apparently normal punctate labeling of the dendritic tips of rod bipolar cells (arrows, Figure 3B and 3C), but immunofluorescence seemed nearly absent at the dendritic tips of cone ON-bipolar cells (arrowheads, Figure 3B and 3C, left \emph{versus} right) with some mGluR6 immunofluorescence appearing instead over bipolar cell bodies in the INL. This dramatic decrease of staining at the dendritic tips of \( \text{Lrit}^{\text{nob6/nob6}} \) cone ON-bipolar cells was also observed for GPR179, RGS7, RGS11 and Gβ5 (arrowheads, Figure 4 A-D), which are also components of the mGluR6 signaling cascade and normally localize at the dendritic tips of both rod and cone ON-bipolar cells (Morgans \emph{et al.}, 2007; Rao \emph{et al.}, 2007; Orlandi \emph{et al.}, 2012; Peachey \emph{et al.}, 2012b; Orhan \emph{et al.}, 2013). Co-immunostaining of mGluR6 with \( \text{G}_\alpha \) (Figure 5A) and with cone arrestin (Figure 5B) confirmed these findings. While \( \text{G}_\alpha \) is a marker of the somas and dendrites of both rod and cone ON-bipolar cells (Vardi, 1998), cone arrestin labels cone photoreceptors including the cone pedicles (Sakuma \emph{et al.}, 1996), which are contacted by the dendritic tips of cone ON-bipolar cells. On \( \text{Lrit}^{+/-} \) retinal sections co-stained with \( \text{G}_\alpha \), mGluR6 was present as small puncta at the dendritic tips of rod bipolar cells (arrows, Figure 5A, left) and as larger patches, where each patch represents the dendritic tips of multiple cone ON-bipolar cells contacting a single cone pedicle (arrowheads, Figure 5A, left). On \( \text{Lrit}^{+/-} \) retinal sections co-stained with cone arrestin, mGluR6 was again normally present as small puncta at rod bipolar cell dendrites (arrows, Figure 5B, left) and as larger patches at the dendritic tips of cone ON-bipolar cells (arrowheads, Figure 5B, left). Interestingly, on \( \text{Lrit}^{\text{nob6/nob6}} \) sections, in the presence of unaltered \( \text{G}_\alpha \) staining, cone-associated mGluR6 staining was absent whereas rod-associated mGluR6 puncta were still present (arrows, Figure 5A, right). The cone arrestin immunofluorescence showed that cone pedicles are present in
the \textit{Lrit3}^{nob6/nob6} retina, but that mGluR6 in close vicinity of cone arrestin was absent (Figure 5B, right).

**Peanut agglutinin staining is disrupted in the OPL in \textit{Lrit3}^{nob6/nob6} retina**

We performed immunostaining on wild-type and mutant retinas with peanut agglutinin (PNA), which binds specific glycosylated residues localized extracellularly to cone inner and outer segments (IS and OS, respectively), and at the synaptic cleft between cone pedicles and cone ON-bipolar cells located in the OPL (Blanks & Johnson, 1983; 1984; Wu, 1984) and was shown to overlap with GPR179 (Ray \textit{et al.}, 2014). In wild-type mice, PNA staining overlapped with mGluR6 staining at the dendritic tips of cone ON-bipolar cells in the OPL (arrowheads, Figure 6A-C, left). In contrast, PNA staining was dramatically decreased in the OPL on \textit{Lrit3}^{nob6/nob6} retinal sections (Figure 6A-C, right). On low-power views of retinal sections, this decrease is obvious in the OPL with no apparent difference of staining intensity in the IS and OS of the cones (Figure 6D). Quantitatively, PNA fluorescence intensity was significantly decreased (64\%, \(p=0.041\)) in the whole OPL of \textit{Lrit3}^{nob6/nob6} retinas compared to wild-type mice (Figure 6E). More strikingly, the number of synaptic clefts stained with PNA between cone pedicles and corresponding cone ON-bipolar cells, per mm of OPL, was dramatically decreased with only 1\% remaining (\(p=0.008\)) in \textit{Lrit3}^{nob6/nob6} retinas compared to wild-type (Figure 6F). In contrast, a measured 23\% decrease of the PNA staining at the IS/OS complex in \textit{Lrit3}^{nob6/nob6} retinas compared to wild-type was not statistically significant (\(p=0.515\)) (Figure 6E).

**LRIT3 staining is similar at the dendritic tips of rod and cone ON-bipolar cells**

As localization of several components of the cone to cone ON-bipolar cell synapse seemed to be disrupted in \textit{Lrit3}^{nob6/nob6} retina whereas their localization at the rod to rod bipolar cell synapse seemed to be unaffected, and because LRIT3 staining at the dendritic tips of cone
ON-bipolar cells seemed to be more intense than that at the dendritic tips of rod bipolar cells, we performed LRIT3 staining quantification at the dendritic tips of rod and cone ON-bipolar cells in 4 Lrit3+/+ mice. LRIT3 staining is 7% less intense at the dendritic tips of rod bipolar cells compared to the dendritic tips of cone ON-bipolar cells but this difference was not significant (p=0.504).
DISCUSSION

In this work, we have produced a specific antibody against mouse LRIT3, localized the corresponding protein in the mouse retina and studied how the localization of most of the known components of the ON-bipolar signaling cascade are affected in a mouse model of cCSNB lacking Lrit3 (Lrit3\textsuperscript{nob6/nob6}). Our results show that LRIT3 localizes at the dendritic tips of both rod and cone ON-bipolar cells, that the correct localization of TRPM1 is dependent on LRIT3 and that the localization of other partners is disrupted in Lrit3\textsuperscript{nob6/nob6} retina particularly at the cone to cone ON-bipolar cell synapse.

**LRIT3 localizes at the dendritic tips of ON-bipolar cells in mouse retina**

The punctate labeling observed in the OPL of wild-type retina with the mouse LRIT3 antibody was absent on Lrit3\textsuperscript{nob6/nob6} retina, demonstrating the specificity of this antibody and validating the knock-out mouse model. This labeling resembles those obtained on human retina with an LRIT3 antibody (Zeitz \textit{et al.}, 2013) as well as those obtained with antibodies directed against various components of the ON-bipolar signaling cascade on mouse retina such as mGluR6, GPR179, NYX, TRPM1, RGS7, RGS11, Gβ5, and R9AP (Masu \textit{et al.}, 1995; Bahadori \textit{et al.}, 2006; Morgans \textit{et al.}, 2006; Morgans \textit{et al.}, 2007; Rao \textit{et al.}, 2007; Cao \textit{et al.}, 2008; Cao \textit{et al.}, 2009; Morgans \textit{et al.}, 2009; Jeffrey \textit{et al.}, 2010; Orlandi \textit{et al.}, 2012; Peachey \textit{et al.}, 2012b; Orhan \textit{et al.}, 2013). Colocalization studies with either ribeye or calbindin (Haverkamp & Wassle, 2000; Schmitz \textit{et al.}, 2000; tom Dieck \textit{et al.}, 2005) confirmed the absence of LRIT3 presynaptically in photoreceptors and in horizontal cells. In contrast, LRIT3 localizes at the dendritic tips of both rod and cone ON-bipolar cells where it colocalizes with mGluR6.

**TRPM1 localization at the dendritic tips of ON-bipolar cell is dependent on LRIT3**
The cation channel TRPM1 is the endpoint of the ON-bipolar cell signaling cascade at the dendritic tips of ON-bipolar cells. TRPM1 opening during light stimulation results in the depolarization of ON-bipolar cells and the formation of the ERG b-wave (Morgans et al., 2009; Shen et al., 2009; Koike et al., 2010). In some mouse models of cCSNB, TRPM1 staining is severely reduced or absent at the dendritic tips of depolarizing bipolar cells. This is the case in the Trpm1 knock-out mouse itself (Morgans et al., 2009; Koike et al., 2010), but also in three different Grm6 mutants: the Grm6 knock-out model (Masu et al., 1995; Xu et al., 2012), the nob3 mouse (Grm6 splice mutation (Maddox et al., 2008; Cao et al., 2011)) and the nob4 mouse (Grm6 missense mutation (Pinto et al., 2007; Cao et al., 2011)). Similarly, the TRPM1 staining at the dendritic tips was also absent in the nob mouse (Nyx deletion (Pardue et al., 1998; Gregg et al., 2003; Pearring et al., 2011)). These results suggest that mGluR6 and nyctalopin, encoded by Nyx, are required for the correct localization of TRPM1.

In contrast, other mouse models deficient for proteins participating in the same signaling cascade (Rgs7−/−/Rgs11−/−, nob5 (deficient for GPR179)) do not show such a mislocalization of TRPM1 (Orlandi et al., 2012; Ray et al., 2014). Therefore these proteins likely play a role in regulating G-protein signaling, rather than a role in trafficking of TRPM1. Although it has been previously shown that nyctalopin is essential for the correct localization of TRPM1 at the dendritic tips of ON-bipolar cells, it was suggested that nyctalopin alone is probably not able to bring the cation channel to the cell surface as it is mainly an extracellular protein (Pearring et al., 2011). We hypothesized that LRIT3 with its intracellular PDZ-binding domain might interact with scaffolding proteins to traffic TRPM1 to its correct localization and, together with nyctalopin, LRIT3 maintains TRPM1 at this localization (Zeitz et al., 2013; Neuille et al., 2014). Indeed, here we show that TRPM1 localization at the dendritic tips of both rod and cone ON-bipolar cells is dramatically decreased in Lrit3−/−/nob6 mice, while the total quantity of TRPM1 was not significantly altered, indicating that localization but not the
protein amount of TRPM1 is dependent on LRIT3. Previous studies have also investigated the localization and protein amount of TRPM1 in mouse models for cCSNB. Quantitative analysis of total intensity of TRPM1 in all retina layers in the knock-out model for mGluR6 was slightly lower compared to wild-type mouse (15% reduction by immunostaining and 23% reduction by Western blot) (Xu et al., 2012). However the authors concluded that the main effect of eliminating mGluR6 is the severe reduction of TRPM1 localization at the dendritic tips of ON-bipolar cells (intensity 30% reduced and number of puncta 60% decreased in the OPL) (Xu et al., 2012). However, this decrease of the total protein amount was not found in another study using the same mouse model (Ray et al., 2014). In the two other mouse models with mGluR6 mutations, nob3 and nob4, in addition to the lack of TRPM1 staining at the dendritic tips of ON-bipolar cells, a moderate 30% reduction of TRPM1 levels in both mouse lines was described by Western blot analysis (Cao et al., 2011). In contrast, the absence of TRPM1 at the dendritic tips of ON-bipolar cells in nyctalopin mutant (nob) mice was accompanied with a severe reduction in total TRPM1 protein quantified by Western blot analysis (Pearring et al., 2011). The observation that the total amount of TRPM1 protein seems unchanged in Lrit3<sup>nob6/nob6</sup> mouse compared to the nob mouse suggests that LRIT3 is necessary for correct trafficking of TRPM1, but that nyctalopin in necessary for both trafficking and either synthesis or stability of TRPM1. It has been shown that nyctalopin, mGluR6, GPR179 and TRPM1 form a macromolecular complex (Cao et al., 2011; Pearring et al., 2011; Orlandi et al., 2013; Ray et al., 2014). It is not clear if LRIT3 directly interacts with TRPM1 or via another component of the complex. Motif analysis suggests that the predicted intracellular PDZ-binding domain present in LRIT3 interacts with an as yet unknown scaffolding protein to bring TRPM1 to the surface (Zeitz et al., 2013).

**LRIT3 may play a role in cone synapse formation**
Surprisingly, we found that most of the known components of the ON-bipolar cell signaling cascade including mGluR6, GPR179, RGS7, RGS11 and Gβ5 are nearly absent at the dendritic tips of cone ON-bipolar cells in $Lrit^3_{nob6/nob6}$ mouse, whereas their staining at the dendritic tips of rod bipolar cells appears normal. In the $Lrit^3_{nob6/nob6}$ mouse, some punctate mGluR6 labeling was present in the INL, possibly indicating a defect in trafficking mGluR6 in cone ON-bipolar cells. We also revealed a dramatic decrease of PNA staining at the synaptic clefts between cones pedicles and corresponding cone ON-bipolar cells in $Lrit^3_{nob6/nob6}$ mouse, whereas cone pedicles are still present. This decrease was qualitatively evident and was well reflected in the quantification of the number of PNA stained synaptic clefts per mm of OPL, while PNA staining was unchanged in IS and OS of cones. Together, these results suggest that LRIT3 may have an additional function in cone synapse formation. To our knowledge, this is the first report of absence of PNA in the OPL in a mouse model of cCSNB. Indeed, PNA staining in the OPL on Grm6 knock-out, nob (Nyx), Trpm1 knock-out and nob5 (Gpr179) retinal sections is not different from wild-type (Ray et al., 2014). However, alterations in ribbon synapse formation or bipolar dendrite invagination have been described previously in Grm6 mutants and Gβ5 knock-out mice (Rao et al., 2007; Cao et al., 2009; Ishii et al., 2009; Tsukamoto & Omi, 2014). Electron microscopy studies are needed to confirm if there is indeed an ultrastructural defect in $Lrit^3_{nob6/nob6}$ retina, which might also explain the slight but significant thinning of inner retinal layers in $Lrit^3_{nob6/nob6}$ mouse that we previously highlighted (Neuille et al., 2014). We quantified LRIT3 staining at the dendritic tips of rod versus cone ON-bipolar cells and showed that the staining is similar at the dendritic tips of both rod and cone ON-bipolar cells. These findings are in accordance with our previous LRIT3 staining on human retina, which revealed equal intensities at the dendritic tips of both rod and cone ON-bipolar cells (Figure 2 in (Zeitz et al., 2013). Thus, the additional function of LRIT3 in cone ON-bipolar cells does not seem to be correlated with a
difference in the amount of LRIT3 protein in these cells compared to rod bipolar cells. Rather, the mGluR6 signaling cascade may be differently regulated in rod and cone ON-bipolar cells.

To conclude, this study reveals that several of the known components of the ON-bipolar cell signaling cascade display a disrupted localization in \( Lrit3^{nob6/nob6} \) mouse, a mouse model of cCSNB lacking \( Lrit3 \). In particular, TRPM1 is reduced at the dendritic tips of ON-bipolar cells, explaining the \emph{no b-wave} phenotype and supporting our hypothesis that LRIT3 might bring TRPM1 to the cell surface. Moreover, LRIT3 may have an additional function in synapse formation as most of the known components of the cascade and PNA are absent at the cone to cone ON-bipolar cell synapse. Putative interactions between LRIT3 and partners of the cascade highlighted here have now to be confirmed to clarify the exact role of LRIT3 in the mGluR6 signaling cascade and to elucidate the pathogenic mechanism(s) of cCSNB.
Acknowledgments

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Abbreviations

AIS: antibody incubation solution; BCA: bicinehinoninic acid; cCSNB: complete form of congenital stationary night blindness; CO₂: carbon dioxide; DAPI: 4’,6-diamidino-2-phenylindole; G₆: heterotrimeric G-protein; GCL: ganglion cells layer; Gpr179/GPR179: G-protein receptor 179; Grm6/GRM6/mGluR6: metabotropic glutamate receptor 6; Gβ5/Gβ5: G-protein beta 5; HRP: horseradish peroxidase; INL: inner nuclear layer; IPL: inner plexiform
layer; IS: inner segments; KLH: keyhole limpet haemocyanin; Lrit3/LRIT3/LRIT3: leucine-rich repeat, immunoglobulin-like and transmembrane domains 3 protein; MAR: melanoma-associated retinopathy; NA: numerical aperture; NaCl: sodium chloride; NaN₃: sodium azide; nob: no b-wave; Nyx: nyctalopin; ONL: outer nuclear layer; OPL: outer plexiform layer; OS: outer segments; PAGE: polyacrylamide gel electrophoresis; PBS: phosphate buffer saline; PDZ: PSD95, DLG1 and ZO1; PKCα: protein kinase C alpha; PNA: peanut agglutinin; R9AP: RGS9 anchor protein; Rgs11/RGS11: regulator of G-protein signaling 11; Rgs7/RGS7: regulator of G-protein signaling 7; ROI: region of interest; SDS: sodium dodecylsulfate; Trmp1/TRPM1: transient receptor potential melastatin 1 cation channel.


Table 1: Primary antibodies used in immunohistochemistry in this study

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Species</th>
<th>Dilution</th>
<th>Reference</th>
<th>Retinal section preparation</th>
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<tr>
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**Figure 1: Validation of mouse LRIT3 antibody** (A) Representative confocal images of cross sections of Lrit3<sup>+/+</sup> and Lrit3<sup>nob6/nob6</sup> retina stained with mouse LRIT3 antibody (green) with or without DAPI (grey). Scale bar, 50 µm. (B) 4x zoom of (A) focused on outer plexiform layer (OPL). Arrows represent putative dendritic tips of rod bipolar cells, arrowheads represent putative dendritic tips of cone ON-bipolar cells. Scale bar, 10 µm.

**Figure 2: Localization of LRIT3 in mouse retina.** Representative confocal images of cross sections centered on the OPL of Lrit3<sup>+/+</sup> retina stained with antibodies against LRIT3 (green) and (A) ribeye (red) (B) calbindin (red) (C) PKCα (red) or (D) mGluR6 (red) and merge (yellow). Scale bar, 10 µm.

**Figure 3: Localization of TRPM1 and mGluR6 as well as TRPM1 protein quantification in Lrit3<sup>nob6/nob6</sup>** Representative confocal images of cross sections centered on OPL of Lrit3<sup>+/+</sup> and Lrit3<sup>nob6/nob6</sup> retina stained with antibodies against (A) TRPM1 (red) and (B) mGluR6 (green) and (C) merge (yellow). Arrows represent putative dendritic tips of rod bipolar cells, arrowheads represent putative dendritic tips of cone ON-bipolar cells, asterisks represent putative somas of ON-bipolar cells. Scale bar, 10 µm. (D) Western blots of 3 Lrit3<sup>+/+</sup> and 3 Lrit3<sup>nob6/nob6</sup> total retinal lysates probed with antibodies to TRPM1 (top) and β-actin (bottom). Bars on the left side of blots represent molecular weight in kDa. Arrows represent bands of interest. (E) Quantification of TRPM1 Western blotting data in panel (D). For each lane, TRPM1 staining intensity was normalized to β-actin staining intensity. Values were plotted as the percentage of the level in Lrit3<sup>+/+</sup>. Error bars represent standard deviation values.

**Figure 4: Localization of GPR179, RGS11, RGS7 and Gβ5 in Lrit3<sup>nob6/nob6</sup> retina** Representative confocal images of cross sections centered on OPL of Lrit3<sup>+/+</sup> and Lrit3<sup>nob6/nob6</sup> retina stained with antibodies against (A) GPR179 (green) (B) RGS11 (red) (C) RGS7 (red) or (D) Gβ5 (green). Arrowheads represent putative dendritic tips of cone ON-bipolar cells. Scale bar, 10 µm.

**Figure 5: Localization of mGluR6, G<sub>α</sub> and cone arrestin in Lrit3<sup>nob6/nob6</sup> retina** Representative confocal images of cross sections centered on OPL of Lrit3<sup>+/+</sup> and Lrit3<sup>nob6/nob6</sup> retina stained with antibodies against mGluR6 (red) and (A) G<sub>α</sub> (red) and merge (yellow) or (B) cone arrestin (green) and merge (yellow).
Arrows represent putative dendritic tips of rod bipolar cells, arrowheads represent putative dendritic tips of cone ON-bipolar cells. Scale bar, 10 µm.

**Figure 6: Localization and quantification of PNA in Lrit3^nob6/nob6 retina** Representative confocal images of cross sections centered on OPL of Lrit3^+/+ and Lrit3^nob6/nob6 retina stained with antibodies against (A) PNA (green) and (B) mGluR6 (red) and (C) merge (yellow). Arrowheads represent putative dendritic tips of cone ON-bipolar cells. Scale bar, 10 µm. (D) Representative confocal images of cross sections of Lrit3^+/+ and Lrit3^nob6/nob6 retina stained with PNA. Scale bar, 50 µm. (E) Quantification of the intensity of PNA staining in the OPL and in the IS/OS complex in 3 independent Lrit3^+/+/Lrit3^nob6/nob6 pairs of retina. Error bars represent standard deviation values. Asterisks represent results that are significantly different (p<0.05). (F) Quantification of the number of PNA stained synaptic clefts between cone pedicles and corresponding cone ON-bipolar cells per mm of OPL in 3 independent Lrit3^+/+/Lrit3^nob6/nob6 pairs of retina. Error bars represent standard deviation values. Asterisks represent results that are significantly different (p<0.05).

**Figure S1: Method for the quantification of LRIT3 immunolabeling in the mouse retina OPL** Representative 3D reconstructions of cross sections centered on OPL of Lrit3^+/+ retina stained with antibodies against LRIT3 (green), PNA (red) and DAPI (white). (A) The yellow box represents the ROI drawn to isolate the OPL in which quantification was made. (B) Zoom on a region of the selected OPL. The dendritic tips of cone ON-bipolar cells appear yellow because of the colocalization of PNA and LRIT3 staining at this localization, whereas the dendritic tips of rod bipolar cells appear only green. (C) The red surface represents the ROI drawn to surround PNA staining and to quantify LRIT3 staining at the dendritic tips of cone ON-bipolar cells. (D) The green spots represent the ROI drawn to isolate each dendritic tip of rod bipolar cells and to quantify LRIT3 staining at this localization. Red and green surfaces do not overlay.
Fig. 1
Fig. 2
Fig. 3
Fig. 4
Fig. 5
Fig. 6