Selective loss of RPGRIP1-dependent ciliary targeting of NPHP4, RPGR and SDCCAG8 underlies the degeneration of photoreceptor neurons

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The retinitis pigmentosa GTPase regulator (RPGR) and nephrocystin-4 (NPHP4) comprise two key partners of the assembly complex of the RPGR-interacting protein 1 (RPGRIP1). Mutations in RPGR and NPHP4 are linked to severe multisystemic diseases with strong retinal involvement of photoreceptor neurons, whereas those in RPGRIP1 cause the fulminating photoreceptor dystrophy, Leber congenital amaurosis (LCA). Further, mutations in Rpgrip1 and Nphp4 suppress the elaboration of the outer segment compartment of photoreceptor neurons by elusive mechanisms, the understanding of which has critical implications in uncovering the pathogenesis of syndromic retinal dystrophies. Here we show RPGRIP1 localizes to the photoreceptor connecting cilium (CC) distally to the centriole/basal body marker, centrin-2 and the ciliary marker, acetylated-α-tubulin. NPHP4 abuts proximally RPGRIP1, RPGR and the serologically defined colon cancer antigen-8 (SDCCAG8), a protein thought to partake in the RPGRIP1 interactome and implicated also in retinal–renal ciliopathies. Ultrastructurally, RPGRIP1 localizes exclusively throughout the photoreceptor CC and Rpgrip1nmf247 photoreceptors present shorter cilia with a ruffled membrane. Strikingly, Rpgrip1nmf247 mice without RPGRIP1 expression lack NPHP4 and RPGR in photoreceptor cilia, whereas the SDCCAG8 and acetylated-α-tubulin ciliary localizations are strongly decreased, even though the NPHP4 and SDCCAG8 expression levels are unaffected and those of acetylated-α-tubulin and γ-tubulin are upregulated. Further, RPGRIP1 loss in photoreceptors shifts the subcellular partitioning of SDCCAG8 and NPHP4 to the membrane fraction associated to the endoplasmic reticulum. Conversely, the ciliary localization of these proteins is unaffected in glomeruli or tubular kidney cells of Rpgrip1nmf247, but NPHP4 is downregulated developmentally and selectively in kidney cortex. Hence, RPGRIP1 presents cell type-dependent pathological effects crucial to the ciliary targeting and subcellular partitioning of NPHP4, RPGR and SDCCAG8, and acetylation of ciliary α-tubulin or its ciliary targeting, selectively in photoreceptors, but not kidney cells, and these pathological effects underlie photoreceptor degeneration and LCA.

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Abnormal biogenesis or deregulation of the function of primary cilia is thought to promote a wide variety of clinical manifestations commonly referred as ciliopathies.1-5 Yet, the molecular pathogeneses of these diseases are ill-defined, due to our lack of understanding of functional relationships between ciliary components and subcellular processes supporting ciliary targeting. Ciliated photoreceptor neurons present a primary cilium connecting two morphologically and functionally distinct subcellular compartments, the inner and outer segments.3 The connecting cilium (CC), which is thought to be equivalent to the transition zone (TZ) of other cilia, projects outward from the basal body (BB) with a membrane-coated bundle of double microtubules organized in a 9+0 pattern and becomes the axoneme (AX) upon entering the outer segment.3 The CC is likely the only route of cargoes’ trafficking between inner and outer segment compartments of photoreceptors and is vital to ciliogenesis, genesis of the outer segment and delivery of components for its maintenance. Yet, multiple lines evidence support that ciliary components shared by distinct cell types often perform cell type-specific function(s) for reasons that are not understood.4-11

The retinitis pigmentosa GTPase regulator interacting protein 1 (RPGRIP1) interactome was found to be comprised of at least three proteins, RPGR, nephrocystin-4/nephroretinin (NPHP4) and RPGRIP1 itself.12-16 Although human-recessive

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mutations in RPGRIP1 are associated only to photoreceptor dystrophies typically with early onset and rampant progression, such as Leber congenital amaurosis (LCA6, (MIM 605446, 613826)), cone-rod dystrophy (CRD13 (MIM 608194)) and juvenile retinitis pigmentosa, heterozygous non-synonymous variants of RPGRIP1 and distinct from all photoreceptor dystrophies-causing mutations were found to be associated to various forms of glaucoma. The exact causes of the clinical and genetic heterogeneity of RPGRIP1 remain unclear, but they likely arise from the differential expression or function of distinct RPGRIP1 isoforms among retinal neurons and species, and loss-of-function (e.g., glaucoma) of mutations between distinct retinal cell types. In this regard, the largest isoform, RPGRIP1α (~175 kDa), is expressed specifically in the retina, where it is localized to photoreceptor neurons. Distinct domains of RPGRIP1α interact directly with RPGR, NPHP4 or selective-encoding domains of RPGRIP1β disrupt specifically the interaction between these partners and they cause retinal dystrophies with or without the involvement of extra-ocular organs (e.g. kidney) and with variable clinical manifestations, such as LCA (MIM 605446, 613826), cone-rod dystrophy (CRD13 (MIM 608194, 304020)) or linked to other systemic diseases, are additional candidates to partake in the genesis of such ciliary complexes, because they localize to the cilium and may participate dynamically and in a cell type-dependent manner in higher-order assembly complexes with RPGRIP1. To shed light into the selective role(s) of RPGRIP1 in photoreceptor function and degeneration, this report defines the relative ciliary localization of components of the RPGRIP1 interactome in photoreceptor and kidney cell types and it uncovers pivotal physiological roles of RPGRIP1 in the ciliary targeting of NPH4, RPGR and SDCCAG8 specifically in photoreceptor neurons, but not in various cell types of the kidney.

Results

RPGRIP1α determines distinct ciliary localizations of NPHP4, RPGR and SDCCAG8 in photoreceptors. The localizations of RPGRIP1α and its assembly components at various ciliary regions of photoreceptors (Figures 1a and b) were compared between wild-type and Rpgrip1α/− mice at P12 of age (Figures 1c–g), when the outer segments begin to develop in wild-type mice and importantly, before pathomorphological changes in the inner segments and cell death ensue in Rpgrip1α/− photoreceptors, because the outcome of these manifestations may lead to the expression of confounding (secondary) phenotypes. We employed high-resolution immunofluorescence microscopy with several antibodies against ciliary markers and components of the RPGRIP1 interactome. RPGRIP1α localized to the cilium distally to the BB marker, centrin-2, with which it colocalizes partially (Figure 1c). RPGRIP1α was absent from the cilium of Rpgrip1α/− mice, but centrin-2 immunolocalization was not affected by the loss of RPGRIP1α (Figure 1c). RPGRIP1α also abuts distally another ciliary marker, acetylated α-tubulin, but its immunostaining was strongly decreased in Rpgrip1α/− photoreceptors (Figure 1d). NPH4, a direct partner of RPGRIP1α, is reported to localize, like other NPHPs, to the TZ of cilia in Caenorhabditis elegans, whereas RPGR, another direct partner of RPGRIP1α and SDCCAG8, also localize to

Figure 1 RPGRIP1α determines distinct ciliary localizations of proteins in mouse photoreceptor neurons. (a) Schematic diagram of a rod photoreceptor with its ciliary region connecting the inner (IS) and outer segment (OS) compartments depicted within the circle (left). Structural organization of the schematic and amplified ciliary region is noted on the right. (b) Overall primary structure and secondary structure with its domains, protein kinase C conserved region 2 (C2) and RPGR-interacting domain (RID), and its interacting partners, NPH4 and RPGR, and examined by this study. (c–g) Left and right panels, respectively, are low- and high-magnification images of distal regions of P12 retinas showing: localization of RPGRIP1α in the connecting cilium and juxtaposed distally to the centriolar/basal body marker, centrin-2, whose localization in the basal body is not affected by loss of RPGRIP1α in Rpgrip1α/− (c); ciliary localization of RPGRIP1β distal of and abutting acetylated α-tubulin and whose ciliary signal is strongly decreased in Rpgrip1α/− (d); ciliary localization of RPGRIP1α distal to NPH4, whose ciliary localization is abolished in Rpgrip1α/− (e); localization of RPGR at the connecting cilium distal to NPH4, the ciliary localization of both proteins are abolished in Rpgrip1α/− (f); NPH4 ciliary localization abutting proximally SDCCAG8, whose ciliary localization is strongly reduced in Rpgrip1α/− (g). Bars represent the mean ± s.d. (n = 4), P < 0.05 is considered significant. Low- and high-magnification scale bars: 10 and 0.5 μm; +/−, wild type; −/−, Rpgrip1α/−. AU, arbitrary units; AX, axoneme; BB, basal body; CC/TZ, connecting cilium/transition zone; DC, daughter centriole; Nuc, nucleus; SR, striated rootlet; Syn, synapse. Dash arrows point to locations of schematic regions in photoreceptor neurons in the retinal sections.
the mouse cilia. In photoreceptors, we found that RPGRIP1z (Figure 1e), RPGR (Figure 1f) and SDCCAG8 (Figure 1g) abut distally NPHP4 and with RPGRIP1z presenting partial and lateral ciliary colocalization with NPHP4 at their interface. In contrast, the Rpgrip1m247cilia were conspicuously void of immunostaining of NPHP4 (Figures 1e–g), whereas immunostaining of SDCCAG8 was extremely weak (Figure 1g). Despite the strong decrease and lack of ciliary staining of SDCCAG8 and NPHP4, respectively, in Rpgrip1m247photoreceptors, the loss of ciliary

![Diagram of photoreceptor structure](image)

![Immunostaining images](image)

![Western blot analysis](image)
localization of NPHP4 and SDCCAG8 was not accompanied by a decrease of expression of these proteins in \textit{Rpgrip1}^{tm247} retinas, even though photoreceptor neurons comprise 70–80% of all retinal cells (Figure 1h). Conversely, the expression level of acetylated \(\gamma\)-tubulin was upregulated significantly in \textit{Rpgrip1}^{tm247} compared with the wild-type retinas, despite of its strong decrease at the photoreceptor cillum of \textit{Rpgrip1}^{tm247} (Figures 1d and h). Likewise, \(\gamma\)-tubulin, another microtubule/centriolar marker, was significantly upregulated in \textit{Rpgrip1}^{tm247} retinas (Figure 1h).

Loss of RPGRIP1\(_{1x}\) expression promotes changes in the subcellular partitioning of SDCCAG8 and NPHP4. The ciliary loss of SDCCAG8 and NPHP4 in \textit{Rpgrip1}^{tm247} photoreceptors without a parallel decrease of the levels of these proteins in retinal homogenates was surprising, as photoreceptors comprise 80–70% of all retinal neurons.\(^{44}\) We have previously reported also the expression of a still unknown RPGRIP1 isoform distinct from the photoreceptorspecific RPGRIP1\(_{1x}\) isoform in uncharacterized class(es) of amacrine neurons,\(^{26,28}\) which collectively account for 10% of all murine retinal neurons.\(^{45,46}\) Hence, the relative contribution of amacrine neurons to changes in the total level of retinal proteins, if any, upon loss of RPGRIP1, is negligible. We hypothesized that the loss of SDCCAG8 and NPHP4 immunolabeling of \textit{Rpgrip1}^{tm247} photoreceptors reflect a loss of antigenicity upon impairment of their RPGRIP1\(_{1x}\)-dependent chaperoning, sorting or targeting that may mask or promote the proteolysis of epitopes. This notion is supported by previous findings with transfected cultured cells, where expression of the LCA-causing mutation, D1114G, in the C-terminal RPGR-interacting domain (RID) of RPGRIP1\(_{1x}\), promotes the loss of antigenicity of its N-terminal coiled-coil domain toward antibodies against this domain.\(^{47}\) Further, RPGRIP1\(_{1x}\) has a high propensity to self-aggregate in the absence of its partners, which are critical for the subcellular targeting of RPGRIP1\(_{1x}\) to the endoplasmic reticulum (ER) or cytosol.\(^{27}\) Hence, to overcome the potential immunohistochecmical limitations described, we examined whether the loss of RPGRIP1\(_{1x}\) expression promotes a change of the levels of SDCCAG8 and NPHP4 between retinal subcellular fractions (Figure 2). In agreement with our prior studies,\(^{26}\) we found that RPGRIP1\(_{1x}\) was predominantly found in the cytoskeletal fraction of retinas of wild-type mice (Figures 2a and b). Although predominant in the cytosolic fraction (cytosolic \textit{versus} any other fraction, \(P<0.016\)), SDCCAG8 was partitioned between the cytosolic, membrane and nuclear fractions (Figures 2a and c). NPHP4 was equally distributed between the cytosolic and membrane fractions (\(P = 0.064\)), where it was largely present (cytosolic and membrane \textit{versus} any other fraction, \(P<0.005\)). Loss of RPGRIP1\(_{1x}\) expression in \textit{Rpgrip1}^{tm247} photoreceptors significantly shifts the localizations of SDCCAG8 and NPHP4 to the ER-associated membrane fraction (Figures 2a, c, d). These and the observations that RPGR and NPHP4 are critical to the targeting or retrieval of RPGRIP1\(_{1x}\) to or from the ER (data not shown),\(^{27}\) prompted us to investigate whether the loss of RPGRIP1\(_{1x}\) expression promotes ER stress. In comparison to wild-type retinas, the levels of GRP78 (glucose-regulated protein of 78 kDa), a stress marker for the ER and cytosol,\(^{48,49}\) were significantly increased in \textit{Rpgrip1}^{tm247} retinas (Figures 2a and e). By contrast, the level of another ER chaperone, calreticulin,\(^{50}\) was decreased in \textit{Rpgrip1}^{tm247} retinas, albeit the difference between wild type and \textit{Rpgrip1}^{tm247} narrowly missed significance (\(P = 0.053\); Figure 2a).

Heterogeneous subcellular localizations of NPHP4, RPGR and SDCCAG8 are not affected in \textit{Rpgrip1}^{tm247} kidney. To determine whether RPGRIP1\(_{1x}\) role in the ciliary targeting of NPHP4, RPGR, SDCCAG8 or acetylated \(\gamma\)-tubulin is cell type and tissue dependent, we examined the ciliary localization of these proteins in the kidney of P21 wild-type and \textit{Rpgrip1}^{tm247} mice when retinal degeneration is already very advanced.\(^{35}\) Acetylated \(\gamma\)-tubulin and SDCCAG8 partially colocalize to perinuclear centrioles of glomerular and tubular cells of the kidney and such colocalization was not impaired in \textit{Rpgrip1}^{tm247} mice (Figure 3a). However, there were glomerular and tubular cells in wild-type mice where the centriolar localizations of acetylated \(\gamma\)-tubulin and SDCCAG8 were mutually exclusive. NPHP4 and RPGR subcellular distributions were heterogeneous between various kidney cell types. NPHP4 localization in the kidney glomerulus was distinct from RPGR (Figure 3b) and SDCCAG8 (Figure 3c). NPHP4 and RPGR had pan-intracellular or centriolar distribution, whereas SDCCAG8 had restricted centriolar localization (Figures 3b and c, upper panels). In tubular cells, RPGR was present at discrete foci throughout the cell, the base of the cilium or centrioles (Figure 3b, middle and lower panels), whereas NPHP4 immunostaining extended from the foot and cap of the BB distally into the AX and proximally into the striated rootlet (SR; Figure 3c, lower panel; see the schema in Figure 3d). SDCCAG8 partially colocalized with and flanked NPHP4 at the appendages of the BB in \textit{Rpgrip1}^{tm247} mice (Figure 3c, lower panel; Figure 3d), but RPGR did not colocalize with NPHP4 (Figure 3b). Further, no changes of NPHP4, SDCCAG8 and \(\gamma\)-tubulin levels were detected in the cortex and medulla of P12 kidney, but by P21 of age, NPHP4 and \(\gamma\)-tubulin were downregulated and upregulated, respectively, in the cortex, but not medulla of \textit{Rpgrip1}^{tm247} (Figure 3e).

Lack of missorting and mistargeting of rhodopsin and M-opsin in \textit{Rpgrip1}^{tm247} photoreceptors. Ciliary trafficking is thought to have a critical role in the transport of opsins. Even though no evidence supports a direct role of RPGRIP1\(_{1x}\) in the subcellular targeting of opsins, two prior studies reported the mislocalization of rhodopsin in \textit{Rpgrip1}^{tm1Tili} mice,\(^{36}\) which still expresses RPGRIP1, but without its C2 and RID domains,\(^{35}\) and \textit{Rpgrip1}^{tm247} mice, which lack RPGRIP1 expression.\(^{35}\) One report suggested that opsin mislocalization may arise from the degeneration of the photoreceptors,\(^{36}\) regardless of the primary defect as demonstrated by others with unrelated disease mouse models,\(^{51}\) whereas the rhodopsin antibody (RET-P1) used by the other study\(^{35}\) is known to detect rhodopsin across multiple subcellular compartments of wild-type photoreceptors, including their cell bodies.\(^{52}\) Regardless, to distinguish primary from potential secondary phenotypes as they relate
to opsin (mis)localization, we re-examined whether the lack of RPGRIP1 expression promotes the mistrafficking of rhodopsin and M-opsin in rod and cone photoreceptors, respectively. First, we employed the same antibodies used by the other studies (e.g., 1D4 and RET-P1) to analyze rhodopsin localization in photoreceptors of P12 and P14 of age when outer segments are beginning to emerge and their growth is well under way, respectively, and importantly, before pathomorphological changes and secondary phenotypes in the inner segments ensue from the absence of RPGRIP1 expression or cell death. No evidence of prominent mislocalization of rhodopsin in the inner segments and cell bodies of rod photoreceptors of \textit{Rpgrip1}\textsuperscript{mm427} mice of P12 and P14 of age were found with any antibody against rhodopsin (Figure 4a, upper and middle panels). Further, rhodopsin was correctly targeted to the apical end of the inner segments of rod photoreceptors (Figure 4a, upper and middle panels). However, the lack of RPGRIP1 expression was accompanied by a twofold reduction of rhodopsin levels in P12.5 retinas of \textit{Rpgrip1}\textsuperscript{mm427} compared with the wild-type mice (Figure 4b), an effect that reflects the absence of developing outer segments in \textit{Rpgrip1}\textsuperscript{mm427} mice. In M-cone photoreceptors, we also did not observe mislocalization and mistargeting of M-opsin and they already appeared much reduced in \textit{Rpgrip1}\textsuperscript{mm427} mice by P14 of age (Figure 4c).
Rpgrip1nmf247 photoreceptors exhibit ruffling of the ciliary membrane and shorter cilia. The extent by which disease mutations impair the interplay between ciliary components and the molecular organization underpinning the ultrastructure of the cilium remains unclear. Hence, we examined qualitatively and quantitatively the effects of lack of RPGRIP1α expression in the ultrastructure of cilia at P12.5 of age, as such effects have not been determined. The developing outer segments of photoreceptors of wild-type mice present disks well stacked and cilia well developed and structured (Figure 5a). Photoreceptors reveal normal BB, CC and AX ultrastructures (Figures 5a–c). In contrast, Rpgrip1nmf247 photoreceptors present naked cilia without outer segments and scarce formation of few nascent and...
The apical targeting of rhodopsin and M-opsin is unaffected in Rpgrip1<sup>nmf247</sup> photoreceptors. (a) Localization of rhodopsin in rod photoreceptors of wild-type (+/+; left panels) and Rpgrip1<sup>nmf247</sup> (−/−; right panels) at P12 and P14 of age with two different antibodies against rhodopsin, 1D4 (upper panels) and RET-P1 antibodies (lower panels). Note, the lack of mislocalization or accumulation of rhodopsin in the cell bodies of photoreceptors with any of the antibodies and the correct polarized targeting of rhodopsin to the apical end of the inner segments of rod photoreceptors of Rpgrip1<sup>nmf247</sup> mice. In comparison to the 1D4 antibody, the RET-P1 antibody detects rhodopsin also in the photoreceptor nuclear layer (ONL) of wild-type and Rpgrip1<sup>nmf247</sup> mice. Images were acquired under the same exact acquisition parameters. Duplicated regions stained with DAPI of retinal sections shown are also displayed for reference to the localization of the cell bodies (ONL) of photoreceptors. (b) Qualitative (upper panel) and quantitative (lower panel) immunoblot analyses of rhodopsin levels in retinal homogenates of wild-type and Rpgrip1<sup>nmf247</sup> mice of P12.5 of age. Bars represent the mean ± S.D. (n = 4). P < 0.05 is considered significant. (c) Localization of M-opsin in cone photoreceptors of wild-type (left panel) and Rpgrip1<sup>nmf247</sup> mice (right panels) at P14 of age. Note, the lack of mislocalization or accumulation of M-opsin in the cell bodies of cone photoreceptors and the correct polarized targeting of M-opsin to the apical end of the inner segments of cone photoreceptors of Rpgrip1<sup>nmf247</sup> mice. The number of M-cone photoreceptors is also decreased in Rpgrip1<sup>nmf247</sup> mice. Images were acquired under the same exact acquisition parameters. Duplicated regions with DAPI of retinal sections shown are also displayed for reference to the localization of the cell bodies (ONL) of photoreceptors. Scale bars: 20 μm in all panels, except in panels of P14 of age with 1D4 staining, where scale bar is 10 μm. +/+; Wild type; −/−, Rpgrip1<sup>nmf247</sup>; AU, arbitrary units; DAPI, 4',6-diamidino-2-phenylindole; IS, inner segment of photoreceptors; ONL, outer nuclear layer; OS, outer segment of photoreceptors.

Figure 4

very irregular disks (Figures 5d–f). Although the overall morphology and relative positioning of accessory ciliary structures, such as the orthogonal positioning between the BB and daughter centrioles, SR and organization of microtubule doublets, appeared normal, there were two significant ultrastructural abnormalities found in ciliary structures of photoreceptors of Rpgrip1<sup>nmf247</sup>. First, there was a decrease of the length of the BB and CC by at least by ~25% compared with age-matched wild-type mice (Figures 5a, e, i). In light of the lack of outer segment disks and a well-defined boundary between the CC and AX, the decrease in ciliary length of photoreceptors in Rpgrip1<sup>nmf247</sup> is very likely underestimated. Our computations of the length of Rpgrip1<sup>nmf247</sup> cilia likely include part of the AX (Figure 1a) as they reflect the distances from the base of the BB up to the longest ciliary tips observed among a large sample of photoreceptors examined. In contrast, the ciliary length of wild-type mice was calculated from the BB to the end of the CC defined at the level of the base the first nascent disk. Second, Rpgrip1<sup>nmf247</sup> photoreceptors exhibited the ruffling of the ciliary membrane throughout the cilium (Figures 5g–k), suggesting the weakening of cytoskeletal bridges between the ciliary membrane and cortical cytoskeleton.

The limited resolution of confocal microscopy (~230 nm) left some ambiguity as to whether the RGRGIP1<sub>α</sub> localization to the photoreceptor CC overlapped partially with that of centrin-2 at the BB (Figure 1c). Hence, we carried out post-embedding ultrastructural immunocytochemistry to define precisely the ciliary boundaries of RGRGIP1<sub>α</sub>. This approach is comparable with procedures employed for confocal microscopy, as it prevented the loss of RGRGIP1<sub>α</sub> antigenicity caused by strong fixation conditions required for transmission electron microscopy. As shown by multiple ultrathin sections of photoreceptors, RGRGIP1 localization was solely restricted to the CC, where it was largely localized throughout its ciliary submembrane region (Figures 5m–p). RGRGIP1 was excluded from the BB and outer segments (AX) of photoreceptors (Figures 5m–p). The accumulation of RGRGIP1 was typically found at the proximal and distal ends
of the CC, such as where newly nascent disks emerge (Figures 5m–p). Quantitation of these observations confirmed that the immunogold particles labeled mostly the CC (Figure 5q).

**Discussion**

This work provides compelling evidence that the RPGRIP1 assembly complex presents a strong degree of subcellular...
and physiological plasticity. Several conclusions can be drawn from our work. First, SDCCAG8, RPGR and RPGRIP1α share similar ciliary localization at the CC by abutting distally NPHP4, centrin-2 and acetylated α-tubulin. Second, RPGRIP1α determines the ciliary localization of NPHP4, RPGR and SDCCAG8 in photoreceptors, because its absence suppresses their ciliary targeting without affecting the overall expression levels of NPHP4 and SDCCAG8, while causing the accumulation of NPHP4 and SDCCAG8 in the ER-associated membrane subcellular fraction. Third, the RPGRIP1α-dependent impairment of the ciliary targeting of NPHP4, RPGR and SDCCAG8 does not affect the apical targeting or cause the missorting of abundant proteins of the outer segment, such as rhodopsin and M-opsin. This observation contradicts a previous report56 and the notion that somehow RPGRIP1 is directly or indirectly implicated in the sorting or targeting of opsins. The reason for this discrepancy arises likely from the broader specificity of the RET-P1 antibody used by Won et al.56 In contrast to the 1D4 antibody, RET-P1 is known also to label the cell bodies, fibers and synaptic pedicles of rod photoreceptors and such labeling may cause easily the misinterpretation of data of mutant mouse strains, such as Rpgrip1−mt/m+ photoreceptors. Instead, RPGRIP1α determines the ciliary targeting of critical photoreceptor disease-related proteins (e.g., NPHP4, RPGR, SDCCAG8) that are pivotal to the formation or maintenance of the outer segments of photoreceptors. Fourth, the effects of RPGRIP1α loss on the ciliary localization of NPHP4, RPGR and SDCCAG8 is highly cell-context dependent, because they are restricted to photoreceptors, but not kidney cells. Further, the ciliary localization of the RPGRIP1 interactome varied between cell types. For example, NPHP4 was excluded from the photoreceptor CC and AX, whereas its localization varied in centrioles, AX and/or TZ of the cilium among cell types of the kidney. This underscores the notion set forth by our prior work that not only the tethering and targeting of the RPGRIP1 interactome harbors structural and functional plasticity and it is highly dynamic at biological level27 but such plasticity occurs also physiologically and likely developmentally as shown by this report. Further, these properties of the RPGRIP1 interactome underpin likely the multiple clinical presentations of disease mutations impairing its components.

Finally, it has been unclear whether the RPGRIP1 interactome presents any primary structural ciliary role(s). We have shown previously that the photoreceptor-specific RPGRIP1α isoform7 is predominantly localized to the detergent-insoluble cytoskeleton fraction of outer segments of bovine photoreceptors.26 Such cytoskeleton localization is consistent with the observations herein reported of the localization of the mouse RPGRIP1α isoform (RPGRIP1α29) also in the cytoskeletal subcellular fraction and that the loss of ciliary localization of the RPGRIP1 interactome promotes the ruffling of the cilary membrane likely through subtle changes and weakening of the submembrane cytoskeletal matrix of the CC. Another significant impact of such ciliary loss is the focal abatement of acetylated α-tubulin in the proximal cilium, a post-translational modification thought to contribute to the stability of microtubules without affecting the regulation of the length of the cilium.53,54 The decrease of acetylated α-tubulin at the cilium of Rpgrip1−mt/m+ photoreceptors may contribute directly to a decrease of the length of the BB and CC by affecting the transport of crucial cargoes (e.g., SDCCAG8, NPHP4, RPGR) required for the formation of outer segments. Indeed, acetylation of α-tubulin is critical to the regulation of transport processes mediated by microtubule-based motor proteins and its misregulation underlies neurodegenerative diseases.55–57 It is noteworthy also that the loss of NPHP4 at the TZ is tolerated by developing cilia in C. elegans, but these cilia present ultrastructural cilary defects, such as B-tubules seam breaks, mislocalization of selective ciliary components and deficits in intraflagellar transport.58 Yet, our study did not find any abnormal ciliary microtubule structures caused by the lack of RPGRIP1-dependent ciliary localization of NPHP4 in the mouse, another evidence of the high degree of molecular plasticity of the RPGRIP1 complex in ciliary targeting, function or between species. This observation is also in agreement with data that mice lacking NPHP4 do not present abnormal microtubule structures in cilia of photoreceptors and spermatogenesis.59 Hence, parsing the molecular bases of the degree of functional plasticity conferred by the remodeling of the RPGRIP1 complex and interplay between its partners will provide insights into shared and unique roles the RPGRIP1 components in sorting and targeting processes and pathways among ciliated cell types (e.g., photoreceptors and kidney), species and development of cell type-restricted clinical manifestations.

Figure 5 Ultrastructural ciliary changes and localization of RPGRIP1α in photoreceptors. Transmission electron micrographs of photoreceptors and their cilia in P12.5 wild type (+/−) (a–c) and Rpgrip1−mt/m+ (−/−) mice (d–k). (a) Longitudinal section of ciliary regions of rod photoreceptors. (b–d) Transverse sections of the connecting cilium at the distal level of transition zone with an amorphous disk structure and ciliary necklace. (d–f) Low and high magnifications of longitudinal sections of photoreceptors of Rpgrip1−mt/m+ mice. Note, Rpgrip1−mt/m+ cilia are naked of outer segments. (g–k) Proximal to distal transverse sections of the connecting cilium depicting the ruffling of the ciliary membrane. (l) Quantitative analyses of the length of basal bodies (BB) and connecting cilium (CC) between wild-type and Rpgrip1−mt/m+ photoreceptors with the latter having shorter cilia. Bars represent the mean ± S.D. (n = 21, −/−; n = 11, +/+) from three mice of each genotype, P < 0.05 was considered significant (Student’s t-test). Only the longest cilia observed in Rpgrip1−mt/m+ mice from the sampling of a large number of cilia were included for quantitation analysis. (m–p) Representative ultrastructural images of the restricted immunogold localization of RPGRIP1α throughout the connecting cilium (CC) of photoreceptors. RPGRIP1α was excluded from the basal body (arrow) and outer segment/taxonem of photoreceptors. RPGRIP1α localization was prominent beneath the ciliary membrane (e.g., m, p) and at the proximal and distal regions of the connecting cilium (e.g., n–p). (q) Quantitation of immunogold particles of RPGRIP1α in the connecting cilium (CC), inner (IS) and outer segments (OS) of photoreceptors. The majority of immunogold particles are restricted to the connecting cilium (CC). Bars represent the mean ± S.D. (n = 22) P < 0.05 was considered significant (Student’s t-test). White arrow, connecting cilium; black arrow, basal body; black arrowhead, proximal (daughter) centriole; white arrowhead, striated rootlets. OC, outer segments of rod photoreceptors; PC, periciliary ridge; RPE, retinal pigment epithelium.
Materials and Methods

Mice. Rpgrip1<sup>pto/d</sup> mice were kindly provided by Patsy Nishina at the Jackson Laboratory (Bar Harbor, ME, USA).<sup>39</sup> Wild-type mice were on a C57Bl6 background and housed in a temperature-controlled and pathogen-free transgenic barrier facility under a 12:12 light-dark cycle. Animal protocols were approved by the IACUC of the Duke University.

Immunohistochemistry reagents. Rabbit antibodies (RPG no 1, RPG no 2) against the recombinant RCC1-homologous domain (RHD) of human RPGR (N-RPGR, a.a.: 1–401) were produced upon four booster shots with recombinant protein (≤ 100 μg) and Hunter’s TiterMax Gold adjuvant (CytRx Corp., Norcross, GA, USA) followed by affinity-purification against the same antigen under non-denaturing conditions according to the manufacturer’s instructions (Sterogene, Arcadia, CA, USA). RPGR no. 2 antibody was employed at 5 μg/ml for immunofluorescence. Other antibodies used for immunofluorescence (IF) or immunoblots (IB) were as following: anti-RPGRIP1 (Ab no. 38, 10 μg/ml);<sup>39</sup> anti-NPHP4 (1:100 (IF), 1:1000 (IB)) was kindly donated by Ronald Roepman; monoclonal mouse anti-acetylated α-tubulin (1:200 (IF), 1:25000 (IB), Sigma-Aldrich, St. Louis, MO, USA); anti-SDCACG8 (1:1000 (IF), IB), ProteinIntech Group, Chicago, IL, USA; anti-centrin-2 (Cetn-2; 1:50) was kindly donated by Jeffrey Salisbury; anti-red opsin p1709 (1:500, (IF),<sup>40</sup> anti-rhodopsin 1D4 (1:500 (IF), 1:40000 (IB); Millipore, Temecula, CA, USA); anti-rhodopsin RET-P1 (1 μg/ml (IF)); Leiko Technologies, St. Louis, MO, USA); monoclonal rabbit anti-γ-tubulin antibody (1:1000 (IF), Abcam, Cambridge, MA, USA); monoclonal rabbit anti-GAPDH (glyceraldehyde 3-phosphate dehydrogenase) antibody (1:500 (IB), Santa Cruz Biotechnology, Inc., Santa Cruz, CA, USA); polyclonal rabbit anti-RPGR (1:1000 (IB), Thermo Scientific); monoclonal rabbit anti-calretilcin (1:1000 (IB), Thermo Scientific); monoclonal mouse mAb414 against nuclear pore complex proteins (e.g., nucleoporin 62 (Nup62; 400 ng/ml IB), Covance, Emeryville, CA, USA). Alexa Fluor-conjugated secondary antibodies and Hoechst 33342 were from Invitrogen (Grand Island, NY, USA).

Immunohistochemistry and confocal microscopy. Fresh eyes and kidneys of P12.5 and P21 mice were employed for ciliary localizations of proteins, whereas rhodopsin and M-opsin were localized in fixed retinas. Fixation of eyes and kidneys was carried in 2% paraformaldehyde, phosphate buffer, pH 7.4, at 4 °C for 4 hours by equilibration in 5% sucrose, 1 × PBS, pH 7.4 for 1 h and then 30% sucrose, 1 × PBS, pH 7.4, overnight at 4 °C. Fresh and fixed tissues were frozen in OCT medium (Sakura, Torrance, CA, USA) on dry ice and stored at −80 °C. Thick sections of 12 μm were cut with a microtome (Microm HM550, Walldorf, Germany). Fixed sections were incubated first with 5% normal goat serum in 1 × PBS, pH 7.4, 1 h, 4 °C for 1 h, then with primary antibodies in the same buffer followed by secondary antibodies (2.5 μg/ml in 1 × PBS, pH 7.4 for 1 h). For fresh sections, primary and secondary antibodies were incubated with 0.1% saponin, 1 × PBS, pH 7.4. Sections were counter-stained with Hoechst 33342 (10 μg/ml). Z-stacks confocal images were imaged with a Nikon C1-laser scanning confocal/Nikon Eclipse 90i microscope controlled by the Nikon EZ-C1 v3.10 software (Nikon, Melville, NY, USA), collapsed, imported to and analyzed in MetaMorph v7.7 (Molecular Devices, Sunnyvale, CA, USA) by computing the integrated density values (ivd) of non-saturating bands corrected against the background. Corrected ivd of each protein was normalized against the corrected ivd of GAPDH. Relative protein optical densities were plotted with Excel (Microsoft). Statistical analysis was determined by the two-sample Student’s t-Test (assuming unequal variances) with Origin 8.5 (OriginLab, Northampton, MA, USA). The cystosolic, membrane, nuclear and cytoskeleton subcellular fractions of two frozen retinas of a mouse of P14 of age were prepared with the Qproteome Cell Compartment kit and exactly as described by the manufacturer instructions (Qiagen, Valencia, CA, USA) with the exception that the cytosolic fraction was collected from retinal lysate upon centrifugation at 1000 r.p.m. All subcellular fractions were solubilized in SDS-sampler buffer, resolved by SDS-PAGE and immunoblots were carried out with antibodies against proteins of interest and markers to each subcellular fraction. The ivd of the immunoblots bands were first corrected by subtracting the ivd background of the same area in the corresponding lane. Then, they were normalized to the ivd of Nup62 of the same fraction, a nuclear pore protein that copurifies with all subcellular fractions<sup>26</sup> and without any significant changes between wild type and Rpgrip1<sup>pto/d</sup>. Upon normalization, the ivd of all four fractions for each of protein in wild type and Rpgrip1<sup>pto/d</sup> were transformed into a percentage scale (total protein = 100%). Average values obtained for each fraction of wild type and Rpgrip1<sup>pto/d</sup> were compared using two-sample t-Test with assumption of unequal variance at the minimum significance level of 0.05.

Conflict of Interest

The authors declare no conflict of interest.

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