RPGR ORF15 isoform co-localizes with RPGRIP1 at centrioles and basal bodies and interacts with nucleophosmin

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The ORF15 isoform of RPGR (RPGRORF15) and RPGR interacting protein 1 (RPGRIP1) are mutated in a variety of retinal dystrophies but their functions are poorly understood. Here, we show that in cultured mammalian cells both RPGRORF15 and RPGRIP1 localize to centrioles. These localizations are resistant to the microtubule destabilizing drug nocodazole and persist throughout the cell cycle. RPGR and RPGRIP1 also co-localize at basal bodies in cells with primary cilia. The C-terminal (C2) domain of RPGRORF15 (ORF15C2) is highly conserved across 13 mammalian species, suggesting that it is a functionally important domain. Using matrix-assisted laser desorption ionization time-of-flight mass spectrometry, we show that this domain interacts with a 40 kDa shuttling protein nucleophosmin (NPM). The RPGRORF15–NPM interaction was confirmed by (i) yeast two-hybrid analyses; (ii) binding of both recombinant and native HeLa cell NPM to RPGRORF15 fusion proteins in vitro; (iii) co-immunoprecipitation of native NPM, RPGRORF15 and RPGRIP1 from bovine retinal extracts and of native HeLa cell NPM and transfected RPGRORF15 from cultured cells and (iv) co-localization of NPM and RPGRORF15 at metaphase centrosomes in cultured cells. NPM is a multifunctional protein chaperone that shuttles between the nucleoli and the cytoplasm and has been associated with licensing of centrosomal division. RPGR and RPGRIP1 join a growing number of centrosomal proteins involved in human disease.

INTRODUCTION

The ORF15-containing isoform of the RPGR protein (RPGRORF15) is mutated in a variety of retinal dystrophies, including X-linked forms of retinitis pigmentosa (1–6), cone dystrophy (1,7), cone–rod dystrophy (8,9), atrophic macular dystrophy (10) and a syndromal form of retinitis pigmentosa (RP) and recurrent respiratory infections with or without sensorineural hearing loss (11,12). Naturally occurring mutations in RPGRORF15 are also found in two independent breeds of dog with progressive retinal atrophy (13). ORF15 is an alternatively spliced C-terminal exon of RPGR, which is expressed in a variety of human tissues but is most abundant in the retina in all species examined (1,13–18).

In the mouse retina, RPGR protein is associated with the connecting cilium of rod and cone photoreceptors, which connects the inner and outer segments (15); this is also the case in humans and other mammalian species, but it is unclear whether it is also expressed in photoreceptor outer segments (17,18). An Rpgr knockout mouse shows a progressive degeneration of cone and rod photoreceptors and is associated with mislocalization of opsin-containing vesicles (15). These
vesicles are normally transported to the base of the connecting cilium by means of the minus-end directed microtubule-associated motor protein cytoplasmic dynein (19). Here, they fuse with the plasma membrane before being transported along the ciliary plasma membrane by plus-end directed microtubule motors such as kinesin-II to the photoreceptor outer segment (20).

The function of RPGR is unclear although structural alignment of the N-terminal half of RPGR (exons 1–11) with RCC1, a guanine nucleotide exchange factor (GEF) for the small GTPase Ran (21–23), suggests a role in the regulation of small GTPases. However, RPGR has not been shown to have GEF activity with Ran. Yeast two-hybrid screening of retinal libraries has shown that the RCC1-like domain of RPGR (RPGRRLD) binds a novel protein of unknown function, RPGR interacting protein 1 (RPGRIP1) (24–27). Loss-of-function mutations in RPGRIP1 cause a form of congenital retinal blindness, Leber’s congenital amaurosis (28–29) and a later onset cone–rod dystrophy (30). RPGRIP1 also localizes to the connecting cilium of rod and cone photoreceptors and an Rpgr1p knockout mouse shows progressive degeneration of rod and cone photoreceptors, with abnormal extended outer segment discs and an inability to localize RPGR to the connecting cilium (27).

The C-terminal half of the RPGR-ORF15 isoform contains a highly repetitive glutamic acid-rich plaid domain, which is a mutation hotspot, plus a non-repetitive and basic ORF15 C-terminal domain (ORF15C2) that is reported to be conserved among human, mouse, bovine and Fugu fish (1). The latter observation suggested that ORF15C2 may have an important functional role. Here we show that the ORF15C2 domain of RPGR interacts and co-localizes with nucleophosmin (NPM or B23). Furthermore, RPGR co-localizes with RPGRIP1 at centrioles and basal bodies of cultured mammalian cells.

RESULTS

Conservation of ORF15C2

We first determined the extent of cross-species conservation of the C-terminal (C2) region of RPGR (ORF15C2), defined by a clone containing the C-terminal 83 amino acids of human RPGR (residues 1070–1152; GenBank accession nos AF286472 and AAC50481), which marks a relatively abrupt change in sequence composition from the acidic glutamic acid-rich ‘plaid’ domain to the basic C-terminal domain (1) (Fig. 1A and B). This was investigated by sequencing the ORF15C2 region in 13 different species, comprising primates (human, gorilla, rhesus monkey, marmoset), rodents (mouse, rat, hamster), carnivores (cat, dog), artiodactyls (cow, pig, sheep) and fish (Fugu rubripes). The results (Fig. 1B) show that amino acid sequence identity within the C2 domain is high across all species, with the exception of F. rubripes, in which identity was confined to the most C-terminal 30 amino acids. The identity of human RPGR was highest with gorilla (96% identity) followed by marmoset (90%), rhesus monkey (87%), cow and cat (64%), sheep (63%), dog and pig (62%), hamster (55%), rat (51%), mouse (45%) and F. rubripes (21%). The high degree of conservation supports the view that ORF15C2 is a functional domain.

Identification of NPM in protein pull-down experiments

A fusion protein containing the C-terminal 81 amino acids of bovine RPGR (ORF15C2) fused to glutathione-S-transferase (GST) was immobilized onto glutathione–Sepharose beads and used to pull down ORF15C2-interacting proteins from bovine retinal and kidney extracts. Eluted proteins were separated by sodium dodecyl sulphate–polyacrylamide gel electrophoresis (SDS–PAGE), and retinal extract bands that were not present in the GST vector-only control were excised, and following in-gel digestion with trypsin, peptide mass mapping was performed using matrix-assisted laser desorption ionization time-of-flight (MALDI-TOF) mass spectrometry. One of the excised bands of mass 40 kDa from the retinal pull-down was identified as NPM by MALDI-TOF (Fig. 1C). The same band was prominent (but not analysed by mass spectrometry) in kidney. Other excised bands were identified as cleavage products of GST, bovine adenine nucleotide translocase, actin, β-tubulin, eukaryotic translation elongation factor 1 gamma, pre-procollagen and glutamate dehydrogenase. Further work was carried out on the putative interaction of ORF15C2 with NPM. The interaction of ORF15 with NPM was initially confirmed by immunoblotting using anti-NPM antibody following GST–bORF15C2 pull down from bovine retinal extracts (Fig. 1D).

Human RPGR ORF15C2 was used as ‘bait’ (pAS vector) and human NPM as ‘prey’ (pACT vector) to test for interaction in a yeast two-hybrid analysis. A positive interaction between NPM and hORF15C2 was shown by the ability to grow on triple selection plates lacking tryptophan (~Trp), leucine (~Leu) and histidine (~His) (Fig. 1E). This was confirmed by testing the ability of putative interacting clones to metabolize X-gal with β-galactosidase, as shown by formation of blue colour (data not shown).

Additional pull-down experiments were carried out using both native NPM in HeLa cell extracts and recombinant NPM. HeLa cell extracts were incubated in vitro with a purified GST human ORF15C2 fusion protein (GST–hORF15C2) and GST control. After washing and elution from glutathione–Sepharose beads, interacting proteins were analysed by SDS–PAGE and western blots. This confirmed that native NPM from crude HeLa supernatant binds to hORF15C2 in vitro (Supplementary Material, Fig. S1A). Recombinant human NPM was also incubated with GST–hORF15C2 in vitro and, after affinity purification and washing, analysis of the eluted products showed that recombinant NPM also binds hORF15C2 (Supplementary Material, Fig. S1B).

The possibility of non-specific interaction between the basic ORF15C2 peptide (pl = 9.64) and the acidic NPM (pl = 4.64) was excluded by testing for interaction between the protein 14-3-3 (pl = 4.63) and GST-tagged ORF15C2 by protein pull-down experiments. Cell extracts from the human embryonic kidney (HEK) 293 cell line, which express both 14-3-3 and NPM as native proteins, were incubated with GST–ORF15C2 and GST control beads. The bound proteins were screened by western blotting using anti-14-3-3 antibody (full-length isoform) and anti-NPM. This showed no interaction between 14-3-3 and ORF15C2 but confirmed the interaction with NPM (Supplementary Material, Fig. S1C). A similar experiment with recombinant enhanced green fluorescent protein
Figure 1. (A) Schematic structure of RPGR\textsuperscript{ORF15\textsubscript{C2}} protein (drawn to scale). The N- and C-termini and RCC1-like, glutamic acid-rich and ORF15\textsuperscript{C2} (C2) domains are shown (with their residue numbers) and the product of exon ORF15, which includes over half of the final protein, is indicated by a line (based on RPGR accession nos AF286472 and AAC50481). (B) Conservation of RPGR\textsuperscript{ORF15} C-terminal (ORF15\textsuperscript{C2}) region in 13 different species. The amino acid sequence of each species is shown below the C-terminal 83 amino acids (residues 1070–1152) of human RPGR\textsuperscript{ORF15} (which defines C2) after alignment using CLUSTALW. Identical residues are shown as boxed in black. Accession nos used for RPGR ORF15 were: AF286472 (human); AY855163 (gorilla); AY855162 (rhesus); AY855164 (marmoset); AF286474 (bovine); Y855169 (sheep); AY855167 (pig); AF385629 (dog); AF286473 (mouse); AY855168 (rat); AY855166 (hamster); AY855167 (cat) and AF286475 (Fugu). (C) Protein pull-down using GST–bORF15\textsuperscript{C2} (GST–C2) with bovine retinal and kidney extracts leading to the identification of NPM, as an interacting protein. Coomassie Blue-stained PAGE gel is shown containing retinal and kidney extract proteins prior to (5% of input extract) and after pull down with GST–bORF15\textsuperscript{C2} (GST–C2) when compared with GST-only controls. The 40 kDa NPM band is indicated by an arrowhead. The major bands are GST and GST–C2. (D) The ORF15\textsuperscript{C2}-interacting band is further identified as NPM (arrowed) with anti-NPM antibody after a protein pull-down experiments using GST–bORF15\textsuperscript{C2} (GST–C2) and bovine retinal extract, compared with GST-only control (GST). The results are compared with anti-GST antibody. Molecular weight size standards (in kilodalton) and antibody dilutions are shown beside each gel. (E) Interaction of ORF15\textsuperscript{C2} (C2) and NPM shown by yeast two-hybrid analysis (in quadruplicate), based on triple nutrient selection (medium lacking tryptophan, leucine and histidine), when compared with positive (POS) and negative (NEG) control plasmids.
anti-NPM antibody did not reveal RPGR ORF15 upon immuno-
cooprecipitation when normal rabbit IgG was used as control. Anti-EGFP antibody was used to co-immunoprecipitate native NPM with EGFP–hORF15C2 (Fig. 2A). A minor NPM band with slightly lower molecular weight was also identified, perhaps representing a post-translational modification or degradation product.

To further corroborate the interaction between NPM and RPGRORF15, HeLa cells were transfected with constructs expressing EGFP fused to hORF15C2 (EGFP–hORF15C2) or with an EGFP control plasmid. Anti-EGFP antibody was used to co-immunoprecipitate native NPM with EGFP–hORF15C2. The results (Fig. 2B) show that RPGRORF15, RPGRIP1 and NPM are each co-immunoprecipitated from retina by anti-ORF15C2 antibody (but not by pre-immune serum or protein A–Sepharose beads).

NPM is predominantly present in nucleoli (31). We therefore tried to determine whether RPGRORF15 forms part of a nuclear complex with NPM in vivo. Two different antibodies (anti-ORF15C2 and anti-ORF15CP) were used in co-immunoprecipitation experiments using bovine retinal nuclear extract (RNE). Immunoblot analysis of immunoprecipitates detected anti-NPM immunoreactive bands of the expected size, ~40 kDa (Fig. 2C). NPM was not immunoprecipitated when normal rabbit IgG was used as control. However, reverse co-immunoprecipitation experiments with anti-NPM antibody did not reveal RPGRORF15 upon immunoblot analysis using ORF15 antibodies (data not shown). This is probably because of low amounts of RPGRORF15 in the NPM-containing multiprotein complex(es) in the retina. Similar sized RPGRORF15 protein bands were detected in RNE with both anti-ORF15 antibodies but the nuclear band was slightly smaller than the major bands seen in whole retinal extracts, suggesting the presence of a distinct nuclear isoform, which showed substantial (50–100-fold) enrichment in nuclei (Fig. 2E). We also found a punctate pattern of nuclear labeling in cell lines using anti-RPGRORF15 antibody in cultured cells, which was absent from cells labelled with pre-immune serum (Fig. 2D).

Co-localization of RPGRORF15 and RPGRIP1 at centrosomes in cultured mammalian cells

Antibodies raised against the bovine ORF15C2 domain (anti-bORF15C2) and against a synthetic peptide (1878) from a non-repetitive region of human ORF15 (anti-hORF151878) were used in indirect immunofluorescence microscopy experiments to determine the subcellular localization of both the transfected and the native RPGRORF15 isoform in a variety of cultured cells. The expression of RPGRORF15 was first examined in cultured mammalian cells using reverse transcriptase–polymerase chain reaction (RT–PCR) (Supplementary Material, Fig. S1D). This confirmed equivalent expression of RPGRORF15 transcripts across cell lines, when compared with the constitutively expressed enzyme glyceraldehyde-3-phosphate dehydrogenase (GAPD).

COS7 cells were transfected with a vector expressing EGFP–bORF15C2 fusion protein and analysed by fluorescence microscopy. Transfected cells could be identified by green fluorescence which co-localized with anti-NPM antibody in nucleoli (Fig. 3A). Constructs expressing the entire exon ORF15 were not detected in the nucleus, presumably because their size exceeds the nuclear pore diffusion limit and RPGRORF15 does not have appear to have a nuclear localization signal (NLS).

The localization of native RPGRORF15 was examined in cultured cells expressing RPGRORF15. Two different anti-ORF15 antibodies (anti-1878 and anti-ORF15C2) co-localized with γ-tubulin in U2OS and HeLa cell lines, consistent with a centrosomal localization (Fig. 3B). In certain views, it appeared that the RPGR labelling of the centrosomes formed a crescent or annular appearance (Fig. 3B, inset). Co-localization of RPGRORF15 with centrosomes was observed in all five cell lines examined—HeLa, U2OS, NIH 3T3, ARPE-19 and COS7—using both anti-ORF15 antibodies. High-resolution imaging revealed two pairs of fluorescent dots in cells in G2 phase of the cell cycle, which was consistent with a specific centriolar localization (Fig. 3C). Localization of RPGRORF15 in mitotic cells revealed spindle pole labelling from prophase through to telophase (Fig. 3D).

The dependence of RPGRORF15 centrosomal localization on microtubule polymerization was examined by culturing cells in the presence and absence of the microtubule destabilizing agent nocodazole. Staining with anti-bORF15C2 and α-tubulin antibodies showed that ORF15 staining of centrosomes was resistant to treatment with nocodazole, hence it was not dependent on the presence of an intact microtubule network (Fig. 4).

Since RPGR interacts with RPGRIP1, and the ciliary localization of RPGR is dependent on the presence of RPGRIP1, two different antibodies to RPGRIP1 were tested in the same cell lines for co-localization with RPGR by indirect immunofluorescence microscopy. The results show that both RPGRIP1 antibodies co-localized with γ-tubulin in U2OS cells consistent with a centrosomal localization (Fig. 4). High resolution imaging again revealed two pairs of fluorescent dots in prophase cells consistent with centriolar localization (Fig. 4). Staining with anti-RPGRIP1 and α-tubulin antibodies showed that RPGRIP1 staining of centrosomes was again resistant to treatment with nocodazole and not dependent on microtubule polymerization (Fig. 4).

The co-localization of native RPGRORF15 and NPM was investigated in cultured mammalian cells using indirect immunofluorescence microscopy (Fig. 5). In both HeLa and U2OS cells, NPM was detected exclusively in nucleoli during interphase, but co-localized with spindle poles in mitosis. This falls in line with previous studies that NPM is a nucleolar protein (31), but that at mitosis it can associate with spindle poles (32). However, in contrast to this latter study (32), we could not detect NPM at interphase centrosomes in an asynchronous population of cells. The spindle pole association of NPM precisely overlapped with the RPGRORF15 signal during...
metaphase but not during anaphase, when the two signals were slightly separated (Fig. 5A). During metaphase, the NPM signal extended outside of the RPGRORF15 signal, in a triangular pattern of labelling, suggesting labelling of centrosomal satellites (33). In contrast, γ-tubulin and RPGRORF15 co-localized precisely at all stages of mitosis (Fig. 5B).

NPM immunofluorescence in the retina

In order to examine the localization of NPM in retina, bovine retinal sections were examined for NPM immunofluorescence, when compared with the centriolar and basal body protein centrin (34), and with RPGR, which was previously reported.
to label connecting cilia but not basal bodies (18). The results (Fig. 6) showed strong centrin and RPGR signals, but no NPM signal, in the region of photoreceptor connecting cilia. Photoreceptor outer segments are also labelled with anti-RPGR ORF15 antibody (Fig. 6B). NPM immunofluorescence was detected in a punctate distribution within each nuclear layer, consistent with nucleolar localization, which was most marked in ganglion cells. A nuclear localization for RPGR was evident in cultured cells (Fig. 2D) but not in retinal sections unless antigen retrieval methods were used (unpublished data). Failure to detect NPM in photoreceptor connecting cilia or basal bodies is probably because it is only present in the centrosomes of mitotic cells.

**RPGR and RPGRIP1 co-localize in basal bodies**

Quiescent mouse inner medullary collecting ductal epithelial (IMCD3) cells possess primary cilia growing from basal

**Figure 3.** RPGR\(^{ORF15}\) localization in human U2OS and HeLa cells. (A) Transfected EGFP–hORF15\(^{C2}\) co-localizes with NPM in nucleoli of cultured HeLa cells. Larger EGFP–ORF15 constructs did not show nuclear labelling. (B) In U2OS cells, antibodies raised against ORF15\(^{C2}\) (green) co-localize with antibodies directed against the centrosome-specific component, γ-tubulin (red). An enlarged view of the region containing the centrosome is shown in the inset. Note the apparent crescent-shaped labelling with anti-ORF15\(^{C2}\) antibody. (C) High-resolution images of centrosomes in U2OS (first three panels) or HeLa (fourth panel) cells labelled with anti-bORF15\(^{C2}\) antibodies. The cells shown have already undergone centrosome duplication and two pairs of centrioles are detected by the anti-bORF15\(^{C2}\) antibodies indicating that RPGR\(^{ORF15}\) is a centriolar, rather than PCM, component. (D) RPGR\(^{ORF15}\) is present at the centrosome at all stages of mitosis in U2OS cells. Prophase, metaphase, anaphase and telophase cells are shown in which spindle poles are detected with anti-bORF15\(^{C2}\) antibodies. No staining of other spindle structures or mitotic chromosomes is detected using anti-ORF15\(^{C2}\) antibody. Scale bars are indicated for b–d.
Figure 4. Staining of centrioles with anti-ORF15\textsuperscript{C2} and anti-RPGRIP1 antibodies is independent of microtubule polymerization. (A) Immunolocalization of RPGRIP1 at centrosomes in a microtubule-independent manner. Human U2OS cells were either untreated (+MTs) or treated (−MTs) with 6 mg/ml nocodazole to depolymerize microtubules. Following fixation, cells were labelled with antibodies against RPGRIP1 (green) and α-tubulin (red) and nuclei stained with DAPI (blue). Merged images including nuclei (DAPI, blue) are shown. (B) An interphase U2OS cell is labelled with anti-α-tubulin (red) and anti-ORF15\textsuperscript{C2} (green) in the absence (+MTs) and presence (−MTs) of the microtubule depolymerizing agent nocodazole. Labelling of centrioles is seen to be resistant to nocodazole. The DNA is stained blue by DAPI. (C). An enlarged view of a prophase cell labelled as described earlier. Note that RPGRIP1 forms two discrete dots representative of centrioles (yellow) within each spindle pole.
Figure 5. Co-localization of NPM and RPGR\textsuperscript{ORF15} at centrosomes in mitotic cells during metaphase. (A) Cell cycle stages in ARPE-19 cells analysed by indirect immunofluorescence, using anti-NPM (red) and anti-ORF15\textsuperscript{52} (green) antibodies. The NPM signal is located at centrosomes in metaphase (closed arrows), in nucleolar granules during interphase (open arrows) and adjacent to centrosomes in anaphase (closed arrows). RPGR\textsuperscript{ORF15} is located at centrosomes in interphase (which are duplicated in G2), and at the spindle poles during mitosis (metaphase and anaphase) (closed arrows). NPM co-localizes with RPGR\textsuperscript{ORF15} in metaphase but extends beyond the RPGR\textsuperscript{ORF15} signal. (B) Co-localization of RPGR\textsuperscript{ORF15} with \(\gamma\)-tubulin (closed arrows) during cell cycle stages in ARPE-19 cells by indirect immunofluorescence, using anti-\(\gamma\)-tubulin (red) and anti-RPGR\textsuperscript{ORF15} (green). RPGR\textsuperscript{ORF15} overlaps with \(\gamma\)-tubulin at all stages of the cell cycle.
bodies which label with anti-acetylated γ-tubulin antibodies. These cells were fixed and used for immunocytochemistry with anti-acetylated γ-tubulin, anti-RPGR ORF15 and anti-RPGRIP1 antibodies. The results show that both RPGR and RPGRIP1 co-localize at the basal bodies in IMCD3 cells (Fig. 7).

DISCUSSION
Here, we show for the first time that both RPGR ORF15 and RPGRIP1 localize to centrosomes and spindle poles in cultured dividing cells. Centrosomes are complex organelles, which act as the major microtubule organizing centre for the cell (33,35). They consist of two orthogonally aligned, barrel-shaped centrioles, each with a 9+0 microtubule triplet array surrounded by a protein matrix, the pericentriolar material, which is involved in nucleating microtubules. During S/G2 phase, centrosomes duplicate, generating two pairs of centrioles and during mitosis, these pairs separate to form the two spindle poles. In certain post-mitotic differentiated cells, such as photoreceptors, centrioles are repositioned within the cytoplasm and contribute to the growth of cilia. Here, the centrioles are referred to as basal bodies. Although not structurally distinct from centrioles, basal bodies do not appear to recruit pericentriolar material. The older centriole/basal body of the pair sits at the base of the cilium and nucleates the ciliary microtubules from its distal end.
Both RPGR\textsuperscript{ORF15} and RPGRIP1 localize to centrosomes throughout the cell cycle in all of the non-ciliated cell lines examined, including HeLa, HEK 293, COS7, NIH 3T3 and ARPE-19. Importantly, the detection of both RPGR\textsuperscript{ORF15} and RPGRIP1 in four small punctate structures late in the cell cycle supports the notion that both these proteins are components of centrioles themselves rather than the surrounding pericentriolar material. RPGRIP1 may serve to anchor RPGR\textsuperscript{ORF15} to the centriole in cultured cells, as it does to the ciliary axonemes of photoreceptors. Localization of both RPGR\textsuperscript{ORF15} and RPGRIP1 to centrosomes is not dependent on microtubules, strengthening the hypothesis that these are core centriolar proteins and not proteins localizing to the minus-ends of cytoplasmic microtubules. In the renal medullary cell line IMCD3, which contains primary cilia nucleated by basal bodies, both RPGR\textsuperscript{ORF15} and RPGRIP1 co-localize at the basal bodies. This is consistent with the close relationship between basal bodies and centrioles. Although both proteins have previously been reported to be present in photoreceptor connecting cilia, neither was known to be a component of basal bodies (18).

Interestingly, we also show that RPGR\textsuperscript{ORF15} isoform interacts with the multifunctional molecular chaperone NPM. NPM is a 40 kDa phosphoprotein, which is present in nucleoli and centrosomes at different stages of the cell cycle. In both these locations, protein ‘crowding’ is significant, and NPM may facilitate specific interactions or prevent protein aggregation and denaturation (31,36–38). It is a member of the nucleoplasmin family of chaperones and has a number of apparently distinct functions, including association with pre-ribosomal particles, binding and cleavage of pre-rRNA, shuttling of ribonucleoprotein and other proteins between nucleus and cytoplasm, binding to NLS-containing proteins and interaction with p53, RB, the nucleolar protein p120, nucleolin and a number of viral proteins (36). It is able to oligomerize, and is found as a hexamer in the nucleolus, although phosphorylated NPM may be monomeric (36).

The interaction of RPGR\textsuperscript{ORF15} and NPM and the co-localization of these two proteins at centrosomes raise the question as to the physiological significance of the interaction. NPM has been suggested to be present at centrosomes from mitosis through to late G1/S (32). We were only able to confirm the presence of NPM at centrosomes during mitosis and specifically during metaphase (Fig. 5A). At this stage, the NPM signal overlaps with RPGR\textsuperscript{ORF15} but extends beyond it, whereas RPGR\textsuperscript{ORF15} is confined to the centrioles. In interphase cells, there was no clear evidence of NPM at centrosomes, whereas there was a strong nucleolar signal. At anaphase, the NPM and RPGR\textsuperscript{ORF15} signals were narrowly separated, suggesting that NPM may be localized to the distal centrosomal satellites (33) or a neighbouring structure. It is not known what proteins anchor NPM to centrosomes but the evidence presented suggests that it is not RPGR\textsuperscript{ORF15}, which overlaps with but does not co-localize with NPM at all stages of the cell cycle.

NPM has been proposed to ‘licence’ centrosomal duplication in late G1/S phase, in co-ordination with DNA duplication, and is then released from centrosomes in response to
Cdk2/cyclin E-mediated phosphorylation of Thr$^{199}$ (which may be maintained by Cdk2/cyclin A during G2) (36). NPM is thought to re-associate with centrosomes during mitosis, possibly in response to phosphorylation of Thr$^{234}$ and Thr$^{237}$ residues by Cdk1/cyclin B (32). One possibility is that RPGR$^{ORF15}$ and RPGRIP1 associate together at the centrosomes and are involved in the unloading of microtubule transport cargoes at various stages of the cell cycle, one of which includes NPM. In this context, the recent demonstration that Ran is a core centrosomal component which can form Ran–GTP from Ran–GDP at this site (38), raises the question as to whether the RCC1-like domain of RPGR$^{ORF15}$ could act as a GEF for centrosomal Ran, perhaps facilitating the unloading of NPM, which has a NLS and is capable of binding to importin. It has been shown that a number of core centrosomal proteins shuttle between the nucleus and the centrosomes, which is blocked by leptomycin (39). These shuttling proteins are proposed to be released from importin in the presence of centrosomal Ran–GTP, by analogy with the unloading of importin cargoes in the nucleus (39). This would explain the transient nature of the interaction and subsequent separation of RPGR$^{ORF15}$ and NPM at later stages of the cell cycle.

Vertebrate photoreceptors contain an outer segment, which is a modified sensory cilium connected to the cell body by a narrow connecting cilium. Both the RPGR$^{ORF15}$ and the interacting protein RPGRIP1 have been consistently localized to ciliary axonemes in all species examined (17–18). The axoneme of connecting cilium is non-motile and projects above the basal body as a 9+0 array of microtubule doublets, corresponding to the transitional zone of motile cilia (40), to which RPGR also localizes (18). All phototransduction and structural proteins that are distributed to the outer segment are transported to the peri-ciliary ridge at the base of the connecting cilium (41), which has been proposed to correspond to the transitional fibre intra-flagellar transport docking site (42). Proteins destined for the outer segment, such as the visual pigment rhodopsin, are carried by the minus-ended microtubule-associated molecular motor cytoplasmic dynein (19), docked in the vicinity of basal bodies and further transported into the outer segments by means of plus-end directed microtubule-associated motors such as kinesin-II (20). Although centrioles are the equivalent structures to basal bodies in photoreceptors, it may be that RPGR$^{ORF15}$ plays a role in the docking of proteins at the base of the connecting cilia, as suggested by the mislocalization of opsin-containing vesicles in the $^{R}$pgr knockout mouse (15), and is a component of a transport complex that shuttles between the inner and outer segments along the connecting cilium. The co-localization of RPGR$^{ORF15}$ and RPGRIP1 to basal bodies in ciliated cells is consistent with this possibility.

What is the significance of the RPGR$^{ORF15}$–NPM interaction in a post-mitotic tissue such as retina? NPM was not seen in photoreceptor connecting cilia by immunohistochemistry, whereas strong labelling of nucleoli was evident. NPM was co-immunoprecipitated from extracts of total retina as well as both nuclear and ROS fractions. When whole retina and crude ROS fractions were examined for nuclear contamination with anti-RCC1 antibody, a weak band was present in ROS (Supplementary Material, Fig. S1E), suggesting that in post-mitotic retinal cells, the major RPGR–NPM interaction occurs within nuclei. There are two possible explanations. The first is that different isoforms of RPGR$^{ORF15}$ (Fig. 2E) serve different roles in nuclei and centrioles/basal bodies. NPM may be common to both by acting as a chaperone for assembling protein complexes including RPGR$^{ORF15}$. The second possibility is that both proteins are involved in the assembly of centrosomal protein complexes within nuclei, which are subsequently exported to centrosomes, as proposed for other core centrosomal proteins, because leptomycin resulted in their nuclear retention (39). Further work is required to establish the precise functions of RPGR and RPGRIP1 both in nuclei and in centrosomes but they join a growing number of centrosomal proteins implicated in human disease (43–46).

**MATERIALS AND METHODS**

**Generation of anti-RPGR$^{ORF15}$ antibodies**

Polyclonal antibodies were raised in rabbits against the bovine ORF15$^{C2}$ domain (anti-$^{b}$ORF15$^{C2}$) (18) and against a synthetic peptide (1097–1107) from a non-repetitive region of human ORF15$^{C2}$ (anti-hORF15$^{1878}$) (GenBank accession nos AF286472 and AAC50481). Another peptide from the C-terminus of human ORF15 protein (1, 1108–1117), was used to generate rabbit polyclonal antibody anti-ORF15$^{CP}$. The antibody was affinity-purified against the cognate peptide. Specificity of the antibodies was checked against *Escherichia coli* expressed fusion proteins and controls (Supplementary Material, Fig. S1F). The ORF15$^{CP}$ antibody identified five to six bands with apparent molecular weight range of 100–250 kDa in mammalian retinas. The bands could be abolished by pre-incubation with molar excess of the relevant peptide, but not with an unrelated peptide. In addition, the immunoreactive bands were not detected in the Rpgr knockout mouse retina (15) (Khanna et al., unpublished data). This antibody gave the same sized bands in fractionated nuclei as anti-ORF15$^{C2}$ antibody (Fig. 2E). The anti-RPGRIP1 antibody was kindly provided by Dr Tiansen Li (26,27) and used at 1:1000 dilution.

**Protein alignment**

The sequences of ORF15$^{C2}$ from gorilla, rhesus monkey and marmoset, cat, rat, hamster, pig and sheep were cloned by PCR using two degenerate primers (forward: 5′-ATM CCA GAR MAG GAA; reverse: 5′-CTT CAA TTC CAR RTA AWG TGG YAA. M = A(C, R = A(G, W = A(T, Y = C(T). PCR products were ligated into TOPO-TA vector (Invitrogen) and sequenced with T3 primer. The sequences of ORF15 C terminal from other species were available at NCBI (http://www.ncbi.nih.gov/). Alignment of these sequences with human ORF15$^{C2}$ was performed using CLUSTALW tool available at the European Bioinformatics Institute (http://www.ebi.ac.uk).

** Constructs**

GST and EGFP fusion constructs were made using the C-terminal 81 and 83 amino acids of bovine and human RPGR$^{ORF15}$ in pGEX 4T3 or pGEX 4T1 (GST tag, Amersham
Pharmacia Biotech) or EGFP-C1 (Clontech), respectively. Bacterial expression was induced with IPTG at 25°C and the supernatant purified with 50% slurry of glutathione–Sepharose 4B beads. Human NPM was expressed in pBAD-TOPO (His tag, Invitrogen) and expression induced with 0.002% arabinose at 25°C.

**Yeast two-hybrid analysis**

The human ORF15C2 construct was inserted into the pAS vector (Clontech) and transformed into the Y190 yeast strain (Yeast protocols Handbook PT3024-1). Human NPM was inserted into the pACT vector (Clontech). A colour assay for β-galactosidase activity with chlorophenol red β-D-galactopyranoside was carried out as described (Clontech Yeast Protocols Handbook PT3024-1).

**Preparation of ROS extracts**

Bovine retinal fractions enriched in ROS were prepared using modifications to the method of Papermaster and Dreyer (47). Briefly, one bovine retina was suspended in 1.0 ml sucrose homogenizing medium (34% sucrose, 65 mM NaCl, 2 mM MgCl2, 5 mM Tris–acetate, pH 7.4) and shaken for 1 min. The resulting suspension was centrifuged at 1800 g for 4 min and the supernatant removed into two volumes 10 mM Tris–acetate, pH 7.4. The pellet was resuspended in 1.0 ml sucrose homogenizing medium and homogenized with a loose fitting pestle, four to five passes on ice. The suspension was centrifuged at 1800g for 4 min, the supernatant removed into two volumes of 10 mM Tris–acetate, pH 7.4 and pooled with the first supernatant. The pooled supernatants were centrifuged at 2600g for 4 min and the resulting pellet was designated the crude ROS extract.

**Pull-down assays**

The GST–ORF15C2 (bovine) protein was used to pull down interacting proteins from bovine retina extracts after overnight incubation at 4°C. One bovine retina (250 mg) and 250 mg bovine kidney were homogenized on ice using a Dounce homogenizer in phosphate buffered saline (PBS) containing 2% Triton X-100 and protease inhibitors and incubated by rotating at room temperature for 30 min. Tissues were further lysed by several rounds of freeze/thaw and sonication and the large debris removed by centrifugation at 10 000g for 20 min. The soluble supernatant was divided into two equal aliquots, one of which was incubated with bovineGST–ORF15C2 and the other with the GST-only vector control, each immobilized on glutathione–Sepharose 4B beads. Bound proteins were washed and removed by boiling, then separated by SDS–PAGE. Excised bands that were absent from GST-only control pull-downs were digested in situ with trypsin and identified by MALDI-TOF mass spectrometry or anti-NPM antibody by western blots.

In other experiments, sonicated HeLa cell extract supernatant was incubated at 4°C overnight with human and bovine EGFP–ORF15C2, washed with PBS and purified on glutathione–Sepharose 4B beads prior to analysis by SDS–PAGE and western blotting with anti-NPM. Recombinant human NPM was also incubated with human GST–ORF15C2 and analysed as described earlier. In SDS–PAGE gels, 10 μl input extract was loaded and 200 μl used for each of the ORF15C2 and control pull downs.

**Protein identification by mass spectrometry**

Identification of NPM by peptide mass mapping utilized methods described in detail elsewhere (48,49). Briefly, gel bands were excised, destained and digested in-gel with trypsin and the resulting peptides extracted for analysis with a Voyager DE Pro MALDI-TOF mass spectrometer (PE Biosystems, Framingham, MA, USA). Measured peptide masses were used to query the Swiss-Protein, TrEMBL and NCBI sequence databases for matches using MS-Fit and Profound search programs.

**Co-immunoprecipitation from cells transfected with EGFP–hORF15C2**

Human ORF15C2, subcloned in the pEGFP vector (Clontech), and pEGFP vector control were transfected into HeLa cells using FuGENE 6 (Roche). After 3 days, the transfected HeLa cells were lysed, and anti-EGFP antibody and protein A beads were added to the lysed supernatant and incubated at 4°C overnight then washed with PBS. After boiling, the extract was subjected to SDS–PAGE followed by immunoblotting with antibody (Supplementary Material, Fig. S1G).

**Co-immunoprecipitation from bovine RNE**

Nuclear extracts were prepared from bovine retina and used for co-immunoprecipitation experiments as described previously (50,51). The immunoprecipitates were subjected to SDS–PAGE followed by immunoblot analysis with appropriate antibodies. The immunoblots were developed using the enhanced chemiluminescence kit (Pierce, New York, NY, USA).

**Immunoblot analysis of the bovine RNE with ORF15CP and ORF15C2 antibodies**

Bovine retinal whole cell extract and nuclear extract (100–200 μg each) were subjected to SDS–PAGE followed by immunoblotting using anti-ORF15CP or anti-ORF15C2 antibody. Equal amounts of protein were run on 7.5% (for ORF15 analysis) or 10% (all other antibodies) SDS–PAGE gels and transferred to nitrocellulose membrane. Membranes were blocked in 5% non-fat dried milk in PBS. Primary antibodies were used at 1:100 [anti-ORF15 and anti-RCC1 (BD Biosciences)] or 1:2000 for anti-NPM (a gift from Dr P.K. Chan, Chinese University of Hong Kong). Anti-rabbit (BioRad) or anti-mouse (Scottish Antibody Production Unit) HRP-conjugated secondary antibodies were used at a 1:5000 dilution. Bound antibody was visualized by ECL (Amersham Biosciences).

**RT–PCR of RPGRORF15**

Total RNA from cultured cells was isolated using the Microto-midi total RNA purification system (Invitrogen). RT–PCR was performed with the SuperScript™ one-step RT–PCR kit.
(Invitrogen). The following primers were used: RPGR<sup>ORF15</sup>, 5'-GAT TCT TTT TCA ATG AGA ACA -3' and 5'-ATT TCC TTT TGA ATC CTC TGC TCC-3'; glyceraldehyde-3-phosphate dehydrogenase (GAPD) 5'-ATG GGG AAG GTG AAG GTC GGA-3' and 5'-TTA CTC CTT GGA GGC CAT GTG-3'. The forward primer for RPGR is in exon 11, the reverse primer is near the start of exon ORF15, which results in an RT–PCR product of ~498 bp.

**Cells and transfection**

COS7 cells were maintained in Dulbecco's modified Eagle medium (DMEM) supplemented with 5% fetal calf serum, penicillin (100 U/ml) and streptomycin (130 μg/ml) at 37°C in an atmosphere of 5% CO₂. Transient transfection was performed using FuGENE 6 (Roche) (6 μl) added to DNA (2.0 μg) following the manufacturer's instructions. Cells were transfected with an enhanced green fluorescent protein (EGFP) vector containing the ORF15<sup>C2</sup> DNA sequence fused to the C-terminus of EGFP. After incubation for 48 h, cells were washed in PBS, fixed with ice cold methanol for 2 min at −20°C, washed in PBS, blocked with 10% normal sheep serum in PBS, then incubated with anti-NPM at 1:100 in 10% normal sheep serum for 1 h at room temperature, in a moist chamber. Cells were washed in PBS, blocked again in 10% normal sheep serum for 1 h at room temperature, in methanol for 10 min. The cells were rehydrated with PBS and incubated with the anti-ORF15 C2 antibody (1:50, for 10 min then blocked in PBS–1% BSA for 5 min. Cells were then incubated with the anti-ORF15<sup>C2</sup> antibody (1:50, rabbit) plus anti-γ-tubulin (1:2000; Sigma, St Louis, MO, USA). The mouse primary antibodies were detected using a Alexafluor two fluorescence microscope and analysed using IPLab software.

Mouse internal IMCD3 were purchased from American Type Culture Collection (Rockville, MD, USA) and cultured to confluence in DMEM/HAM's F12 (Gibco-BRL, Rockville, MD, USA) and supplemented with 10% fetal bovine serum (FBS) and 100 U/ml of penicillin/streptomycin in a humidified 5% CO₂ incubator, maintained at 37°C.

**Immunocytochemistry**

COS7 cells were grown at 37°C in a 5% CO₂ atmosphere in DMEM supplemented with 10% FBS and penicillin–streptomycin (100 IU/ml and 100 μg/ml, respectively). For immunofluorescence, cells were grown on acid treated coverslips overnight before being fixed and permeabilized with −20°C methanol for 10 min. The cells were rehydrated with PBS for 10 min then blocked in PBS–1% BSA for 5 min. Cells were then incubated with the anti-ORF15<sup>C2</sup> antibody (1:50, rabbit) plus anti-γ-tubulin (1:2000; Sigma, St Louis, MO, USA). The mouse primary antibodies were detected using biotin anti-mouse (Amersham) followed by incubation with Streptavidin Texas Red to detect the biotin (Amersham). The rabbit primary antibodies were detected using Alexa 488 (Molecular probes, Eugene, OR, USA) and the DNA was stained using Hoechst 33258 dye (Calbiochem, 0.2 μg/ml) for 1 min. All antibodies were diluted in PBS–3% BSA and all antibody incubations were for 45 min at room temperature. The cells were washed 3 × 5 min between each antibody incubation. Following the completion of labelling, the coverslips were mounted onto slides in 80% glycerol 3% propylgallate to prevent fading. Images were captured on a Nikon TE300 inverted microscope using an ORCA ER Hamamatsu, Shizuoka, Japan) using Openlab 3.09 software (Improvement, Coventry, UK) and processed using Adobe Photoshop (Adobe Systems, San Jose, CA, USA).

**Fluorescence staining of retinal cryosections**

Immunofluorescence studies were essentially performed as previously described (52,53). Briefly, eyes from adult C57BL/6J mice were prefixed in 4% paraformaldehyde in PBS for 1 h at room temperature, washed, soaked with 30% sucrose in PBS overnight, and cryofixed in melting isopentane. Cryosections were placed on poly-L-lysine-precoated coverslips. Specimens were incubated with 50 mM NiCl₂ and 0.1% Tween-20 in PBS and blocked with blocking solution [0.5% cold-water fish gelatin (Sigma) plus 0.1% ovalbumin (Sigma) in PBS]. For double labelling, sections were incubated with a mixture of antibodies in blocking solution overnight at 4°C. In these experiments, anti-centrin antibodies were used as markers for the connecting cillum (54). The specimens were washed and subsequently incubated with secondary antibodies conjugated to Alexa<sup>488</sup> or Alexa<sup>546</sup> (Molecular Probes) in blocking solution for 1 h at room temperature in the dark. Washed sections were mounted in Mowiol 4.88 (Hoechst, Frankfurt, Germany) containing 2% propylgallate to prevent fading. Images were captured on a Leica DMRP microscope. Images were obtained with a Hamamatsu Orca ER CCD camera (Hamamatsu city, Japan) and processed with Adobe Photoshop (Adobe Systems).

**SUPPLEMENTARY MATERIAL**

Supplementary Material is available at HMG Online.

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**REFERENCES**


