Usher syndrome and Leber congenital amaurosis are molecularly linked via a novel isoform of the centrosomal ninein-like protein

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Usher syndrome (USH) and Leber congenital amaurosis (LCA) are autosomal recessive disorders resulting in syndromic and non-syndromic forms of blindness. In order to gain insight into the pathogenic mechanisms underlying retinal degeneration, we searched for interacting proteins of USH2A isoform B (USH2A isoB) and the LCA5-encoded protein lebercilin. We identified a novel isoform of the centrosomal ninein-like protein, hereby named Nlp isoform B (Nlp isoB), as a common interactor. Although we identified the capacity of this protein to bind calcium with one of its three EF-hand domains, the interaciton with USH2A isoB did not depend on this. Upon expression in ARPE-19 cells, recombinant Nlp isoB, lebercilin and USH2A isoB were all found to co-localize at the centrosomes. Staining of retinal sections with specific antibodies against all three proteins revealed their co-localization at the basal bodies of the photoreceptor-connecting cilia. Based on this subcellular localization and the nature of their previously identified binding partners, we hypothesize that the pathogenic mechanisms for LCA and USH show significant overlap and involve defects in ciliogenesis, cilia maintenance and intraflagellar and/or microtubule-based transport. The direct association of Nlp isoB with USH2A isoB and lebercilin indicates that Nlp can be considered as a novel candidate gene for USH, LCA and allied retinal ciliopathies.

INTRODUCTION

Usher syndrome (USH) is the most common cause of combined deaf-blindness, with a prevalence of about one in 20 000 in the Caucasian population (1–3). It is a clinically and genetically heterogeneous disorder for which three different types can be distinguished based on the severity and progression of the hearing loss, the presence or absence of vestibular symptoms and the age of onset of retinitis pigmentosa (RP) (4). To date, six different genetic loci have been identified for USH type I (USH1), three loci for USH type II (USH2) and one locus for USH type III (USH3) (reviewed in 5). Mutations have been identified in the genes encoding myosin VIIa, harmonin, cadherin 23, protocadherin 15 and SANS as the underlying defect in USH1b, -c, -d, -f and -g, respectively (6–11). Defects in USH2A, GPR98 (VLGR1) and whirlin are causative for USH2a, -c and -d, respectively (12–15). USH3 is caused by mutations in the clarin-1 encoding USH3A gene (16).

Recently, we and others have shown that all known USH1 and USH2 proteins are present in a protein network, the Usher
interactome, in which harmonin and whirlin play the role of key organizers that interconnect the other USH proteins (17–22). Localization studies in rodents revealed that this interactome is present in the stereocilia and the synaptic region of hair cells in the cochlea and in the synaptic region of photoreceptor cells. Except for harmonin, all known USH1 and USH2 proteins are also present at the region of the connecting cilium of photoreceptor cells, a microtubule-based structure which separates the outer segments from the inner segments (21). This region encompasses the connecting cilium, its basal body and centriole as well as the periciliary region surrounding the connecting cilium. In the region of the connecting cilium, the Usher interactome may be recruited via PDZ (postsynaptic density 95, PSD-95; discs large, Dlg; zona occludens-1, ZO-1) domain-based interactions with whirlin (19,21,23). Based on the localization of the Usher protein network to the region of the connecting cilium and the identification of ciliary abnormalities in photoreceptor cells and sperm cells of several patients with USH (24), the syndrome can be considered as a (retinal) ciliopathy (25). Recent phenotypic data of patients with different Usher subtypes (USH 1B, 1F, 2A or 2C) also suggest that the primary pathogenic insult does not take place at the photoreceptor synapse (26).

Leber congenital amaurosis (LCA) is another clinically and genetically heterogeneous recessive disorder which has been described as a severe form of RP presenting before the age of 1 year with the absence of photoreceptor function (27). Patients present with profound visual loss, nystagmus, poorly reactive pupils and a markedly diminished or non-detectable electroretinogram (27,28). Defects in the centrosomal and cilia-associated proteins RPGRIP1, CEP290 and lebercilin (encoded by LCA5) have been identified to be associated with LCA (29–31), implicating a role of these proteins in ciliary processes.

At the moment, mutations in several different genes have been described to be the underlying cause of USH and LCA, but the molecular basis of the pathogenic defects of these syndromes still remains largely elusive. Based on similarities in clinical manifestations and the subcellular localization of proteins involved in USH and a number of proteins involved in LCA, the pathogenic mechanisms underlying both disorders might significantly overlap. In order to gain a better insight into the molecular basis of USH and LCA, we searched for novel interacting partners of the recently identified lebercilin (LCA5) and the intracellular region of USH2A isoform B (USH2A_tail) by using yeast two-hybrid screening.

In this study, we show that both lebercilin and the intracellular region of USH2A_tail directly interact with a novel isoform of the second member of the ninein family, the centrosomal ninein-like protein [Nlp isoform B (Nlp_isoB)], encoded by the Nlp (KIAA0980) gene. Nlp_isoB is a novel component of the Usher interactome, connecting USH and LCA at the molecular level, and the Nlp gene can therefore be considered as a novel candidate for USH, LCA, and related disorders.

RESULTS

A novel isoform, isoform B, of the centrosome-associated ninein-like protein interacts with USH2A and lebercilin

Yeast two-hybrid screens of an oligo-d(T) primed human retina cDNA library were performed to identify proteins interacting with lebercilin or with the intracellular region of USH2A_tail. Analysis of positive clones that activated all reporter genes revealed a common interactor for both bait proteins. In total, 11 overlapping clones of a novel splice variant of the centrosomal ninein-like protein (Nlp), hereby named Nlp_isoB, were identified. By using the intracellular region of USH2A_isoB as a bait, two clones for Nlp_isoB were identified. These encode amino acids 598–1033 and 658–1033, respectively. For lebercilin, four Nlp clones encoding amino acids 598–1033 and five clones encoding amino acids 658–1033 were identified. The transcript encoding Nlp_isoB lacks exon 17 from the originally described Nlp gene, resulting in in-frame skipping of 349 amino acids after residue 734. The specificity of the interaction in the yeast two-hybrid assay was determined by co-transforming pAD-Nlp_isoB (amino acids 658–1033) with the pBD-Gal4 vector expressing the non-related p63 protein. No interaction was observed performing this control experiment (data not shown).

Bioinformatic analysis of the Nlp_isoB protein (SMART database; http://smart.embl-heidelberg.de/) identified three regions that are predicted to form EF-hand domains, potentially Ca²⁺-binding, which are known to be involved in the interaction with γ-tubulin (32). In addition, the C-terminal region was predicted to form an intermediate filament (IF) domain (amino acids 656–925), likely to be involved in protein–protein interactions (Pfam Home Page; http://pfam.sanger.ac.uk/) (Fig. 1A). The IF domain, however, was predicted with a low significance (E-value: 5.2 × 10⁻¹). We analysed the spatial and temporal expression pattern of transcripts encoding Nlp_isoA and Nlp_isoB by semi-quantitative RT–PCR, using several human fetal and adult tissues. No major differences in expression were observed for isoforms A and B, indicating that both isoforms function in the same tissues (Fig. 1B). However, differences at the cellular level cannot be excluded.

Deletion constructs were made of the intracellular region of USH2A_isoB and Nlp_isoB and tested in yeast two-hybrid analysis to determine the regions that are involved in the interaction. By using these deletion constructs, we were able to show that in USH2A, the interacting region is located in the fragment containing amino acids 5124–5196 and in Nlp_isoB in the fragment containing the predicted IF domain (amino acids 656–925) (Fig. 2A). In order to further pinpoint the domains involved in the interaction between Nlp_isoB and lebercilin, parts of the lebercilin protein comprising the amino acids 1–95, 96–305, 306–490 and 491–697 were tested in a yeast two-hybrid assay with different parts of Nlp_isoB. This revealed that the predicted IF domain of Nlp_isoB specifically interacts with the fragment encompassing the first two coiled-coil domains of lebercilin (amino acids 96–305) (Fig. 2B). To test whether the interactions of USH2A_isoB and lebercilin with Nlp are isoform-specific, a liquid β-galactosidase assay was performed using both Nlp_isoA and Nlp_isoB together with lebercilin and the intracellular region of USH2A_isoB. A specific interaction was observed between both proteins and Nlp_isoB whereas no interaction with Nlp_isoA could be detected (Fig. 2C and D).

Co-immunoprecipitations were performed to confirm the interaction between USH2A and Nlp_isoB. For this purpose, COS-1 cells were co-transfected with constructs encoding HA-tagged Nlp_isoB and the flag-tagged intracellular region of USH2A (USH2A_tail). From the COS-1 cell lysates we were able to co-immunoprecipitate HA-tagged Nlp_isoB and flag-tagged...
USH2A with antibodies against the flag-tag. As a negative control, unrelated flag-tagged STRAD was co-expressed with HA-tagged NlpisoB. As expected, HA-tagged Nlp isoB and flag-tagged STRAD did not co-immunoprecipitate (Fig. 3A).

The interaction between lebercilin and NlpisoB could be confirmed in a glutathion-S-transferase (GST) pull-down experiment and in co-immunoprecipitation assays. We were able to pull down flag-tagged lebercilin from a COS-1 cell lysate with GST-fused NlpisoB but not with GST alone (Fig. 3B). In addition, we were able to co-immunoprecipitate flag-tagged NlpisoB and HA-tagged lebercilin with antibodies against the HA-tag. As a negative control, unrelated flag-tagged LRRK2 was co-expressed with HA-tagged lebercilin. No LRRK2 was co-immunoprecipitated with lebercilin (Fig. 3C). Also, co-immunoprecipitation assays of endogenous proteins were performed from bovine retinal extracts by using antibodies against lebercilin and Nlp. We were able to co-immunoprecipitate Nlp with an antibody against lebercilin. As a negative control, rabbit IgGs were used (Fig. 3D).

**Interaction between USH2A and Nlp is calcium-independent**

Because of the predicted presence of three EF-hand domains in the N-terminal half of Nlp, we hypothesized that the interaction between USH2A isoB and Nlp isoB might be dependent on the binding of Ca^{2+} ions to these EF hands, as was shown for the interaction between the EF-hand domain containing proteins centrin and transducin (33). Therefore, we first determined the Ca^{2+}-binding capacity of Nlp. At residue 12 of EF-hand domains, an invariant Glu or Asp provides two oxygens for liganding Ca^{2+} (bidentate ligand) (34). Based on this, only EF hand 3 would be able to efficiently bind Ca^{2+} (Fig. 4A). To test this hypothesis, GST-fusion proteins were made of the three predicted EF hands of Nlp. As a positive control, the GST-fused calmodulin 28K subunit was used. We indeed were able to show binding of Ca^{2+} to Nlp EF hand 3 and not to EF hands 1 and 2 (Fig. 4B). Subsequently, we performed GST pull-down assays in the presence or absence of 2mM CaCl₂ in which HA-tagged full-length Nlp isoB was pulled down from a COS-1 cell lysate with GST-USH2A tail but not with GST alone. Also, no HA-tagged Nlp isoB was pulled down with GST-fused NBC3 tail. NBC3 was used as an additional negative control and is a previously described member of the Usher interactome (17). Similar results were obtained for pull-down experiments performed in the presence or absence of calcium, and therefore, there are no indications for calcium-dependence of the interaction (Fig. 4C).

**Nlp co-localizes with USH2A isoB and lebercilin at the photoreceptor cell basal body and centriole**

We performed immunohistochemistry for Nlp and USH2A isoB to determine whether both proteins co-localize in the retina.

Figure 1. Schematic representation of the protein structure of Nlp isoA and Nlp isoB. (A) The predicted EF hands 1, 2 and 3 are formed by amino acids 11–39, 200–228 and 237–265, respectively. Coiled-coil domains (CC) 1, 2, 3 and 4 are formed by amino acids 384–425, 470–579, 621–699 and 1046–1375, respectively, and the predicted IF domain is formed by amino acids 656–925. (B) Semi-quantitative RT–PCR for transcripts encoding Nlp isoA and Nlp isoB in human fetal and adult tissues. Shown are the samples that were taken after 35 cycles. The results are representative for the samples taken after 25 and 30 cycles. As a control, RT–PCR analysis of the housekeeping gene GAPDH was performed. The transcripts for Nlp isoA and Nlp isoB show a similar distribution, with the strongest expression in fetal cochlea and adult brain, testis and kidney.
Monoclonal anti-pan-centrin antibodies (20H5) were used as a marker for the basal body and connecting cilium. With affinity-purified antibodies against Nlp, the presence of Nlp was detected in the inner segment and at the basal body, shown as a partial co-localization with centrins (Fig. 5B). On retinal cryosections, Nlp and USH2AisoB co-localized in the inner segments and at the region of the connecting cilium (Fig. 5A). In addition, USH2AisoB localized to the outer plexiform layer as was already described (17,19).

To determine the exact subcellular localization of Nlp in the retina, immunoelectron microscopy was performed. We detected Nlp in the basal body and the centriole of the photoreceptor-connecting cilium as well as in the periciliary region of the apical inner segments of mouse photoreceptor
cells (Fig. 5C). Immunoelectron microscopy for USH2AisoB on the retina showed the presence of USH2AisoB in the periciliary region, the connecting cilium, basal body and the centriole of photoreceptor cells (Fig. 5D) (21). Immunohistochemistry on mouse retinal cryosections revealed lebercilin as a component of the photoreceptor-connecting cilium and basal body (29). Thus, our results indicate that Nlp co-localizes with USH2AisoB and also with lebercilin at the photoreceptor cell basal body.

**Nlp, lebercilin and USH2A co-localize at the centrosome of ARPE-19 cells**

Casenghi et al. (32) have shown that Nlp localizes at the mother centriole of the centrosome in cells during interphase. In order to visualize the interaction between the intracellular domain of USH2A and NlpisoB, we fused these proteins at their N-terminus to enhanced cyan fluorescent protein (eCFP) and monomeric red fluorescent protein (mRFP), respectively. In single transfected human retinal pigment epithelium cells (ARPE-19) expressing NlpisoB, this protein shows a centrosomal localization specifically at one centriole, most probably the mother cilentriole (Fig. 6A). In mitotic cells, a punctate localization in the cell periphery was observed for NlpisoB (data not shown). In single transfected cells, a nuclear localization (data not shown) (17,19) or a centrosomal localization was observed for eCFP-USH2A (Fig. 6B). In cells co-expressing NlpisoB and USH2A, co-localization was observed at both the mother and the daughter centriole of the cell (Fig. 6D–D’). In addition, overexpression assays in ARPE-19 cells were performed by co-expressing N-terminally fused mRFP-NlpisoB and C-terminally fused lebercilin-eYFP. In single transfected cells, lebercilin-eYFP was localized to the centrosome and microtubule network of the cell, as previously described (29) (Fig. 6C). In cells co-expressing Nlp isoB and lebercilin-eYFP, co-localization at the centrosome and non-centrosomal microtubule organization centres (MTOCs) was observed (Fig. 6E–E’).

To determine the subcellular localization of endogenous Nlp and lebercilin, ARPE-19 cells were induced to form primary cilia by serum starvation and subsequently used for
immunohistochemical stainings with antibodies directed against Nlp and lebercilin. Both proteins could be detected at the base of the cilium, most probably the basal body (Fig. 7A and B). Interestingly, we detected both lebercilin and Nlp at the midbody during cytokinesis (Fig. 7C and D). After telophase, the mother centrioles which are present at the midbody develop into the basal bodies of the newly formed cells (35). No endogenous USH2A was detected in ARPE-19 cells.

**Lebercilin mutations affect the interaction with Nlp isoB**

Recently, we have identified truncating lebercilin mutations in LCA patients (29). We addressed the biologic relevance of the interaction between lebercilin and Nlp isoB by testing the effect of two of these mutations, p.Q279X and p.P493TfsX1, on the association with Nlp isoB. In a quantitative yeast two-hybrid interaction assay, the p.Q279X mutation significantly enhanced the interaction of Nlp isoB and lebercilin, whereas the p.P493TfsX1 mutant was found to be completely abolished (Fig. 8A).

The subcellular localization of the lebercilin mutants was studied upon expression in ciliated ARPE-19 cells. When expressed alone, mRFP-tagged lebercilin localizes to the basal body, the primary cilium and the microtubule network of the cell (Fig. 8B) (29). The mRFP-lebercilin Q279X mutant does not associate with the microtubule network any longer but is present in the primary cilium and its basal body (Fig. 8C). In contrast, mRFP-lebercilin P493TfsX1 only localizes to the microtubule network in the cell periphery but no longer to the cilium and the basal body (Fig. 8D). Co-expression of Nlp isoB and lebercilin showed a clear co-localization, specifically at the basal body (Fig. 8E–E'). In the primary cilium and the microtubule network, only lebercilin was present. As indicated by the yeast two-hybrid assay, the interaction between Nlp isoB and lebercilin Q279X is enhanced. When co-expressed with Nlp isoB, lebercilin P493TfsX1 is recruited to the basal body, but is no longer observed in the primary cilium (Fig. 8F–F'). Upon co-expression of Nlp isoB and lebercilin P493TfsX1, no co-localization was observed, confirming the loss of interaction between the two proteins (Fig. 8G–G').

These data indicate that truncating mutations in lebercilin severely affect the association with Nlp at the basal body. However, downregulation of both endogenous Nlp and LCA5 by RNAi in ciliated ARPE-19 cells does not result in altered protein localization of Nlp or lebercilin (Supplementary Material, Fig. S2), suggesting that additional binding partners are involved in their localization at the basal body.

**DISCUSSION**

In this study, we demonstrate that a novel isoform of the centrosome-associated ninein-like protein, Nlp isoB, specifically interacts with lebercilin and the cytoplasmic region of USH2A, thereby linking USH and LCA at the molecular level. Previous analysis of *Xenopus laevis* Nlp already suggested the presence of two Nlp isoforms, which correspond with human Nlp isoA and Nlp isoB (36). Nlp is the second member of the ninein protein family and has a 37% sequence identity with ninein (32). Nlp is predominantly present in the mother centriole of the centrosome and in the basal body of primary cilia in cultured cells and is involved in microtubule nucleation, anchoring and outgrowth (32,36,37). However, it remains to be elucidated which isoform of Nlp is present and functions at these subcellular structures. The interaction between USH2A isoB, lebercilin and Nlp isoB and their co-localization in the basal bodies, the centrioles and the periciliary compartments of photoreceptor cells show that at least Nlp isoB molecules are present there.

Basal bodies and associated centrioles are found at the base of cilia and serve as a nucleation site for the axonemal and cytoplasmic microtubules, respectively (38,39). The photoreceptor cell outer segment is regarded as a highly specialized cilium corresponding to the cilary shaft of a prototypic cilium (38,39). The connecting cilium then correlates with the short junction between the basal body and the axoneme of a prototypic cilium, the transition zone (38). The presence of Nlp at the basal bodies and centrioles of the ciliary apparatus of photoreceptor cells matches with the previously determined localization of Nlp in the basal body of ciliated cells (36). Based on the knowledge on the Nlp function in mitotic cells (32,36,37), we propose that Nlp functions in the development of
and maintenance of the connecting cilium and the outer segment and in the establishment of a microtubule network in (the apical part of) the inner segment in differentiated photoreceptor cells.

In addition to the physical interaction of both USH2A isoB and lebercilin with Nlp isoB, similarities in clinical manifestations between USH and LCA suggest a significant overlap in the pathogenic mechanisms underlying both disorders. Based on the subcellular localization and the current knowledge on USH2A isoB and lebercilin, these pathogenic mechanisms are likely to include ciliary dysfunction. Post-mortem observation of connecting cilium defects in the retina and the identification of sperm abnormalities in some USH patients (reviewed in 40), as well as the involvement of other cilia-associated proteins in non-syndromic forms of retinal degeneration and USH, contribute to this hypothesis (49–52). In photoreceptor cells, IFT20, IFT52, IFT57 and IFT88 molecules and both kinesin motors are thought to be present in an anterograde transport complex (53–55). Dysfunction of the components of the IFT transport complex lead to mislocalization of structural proteins and proteins involved in the phototransduction and subsequently lead to retinal degeneration (53,56). This is corroborated by the development of both polycystic kidney disease in addition to retinal degeneration in mice after the introduction of a hypermorphic mutation in IFT88 (53,57). Also, photoreceptor-specific silencing of KIF3A leads to mislocalization of opsin and arrestin, proteins involved in phototransduction (56). Similar defects were observed in mice lacking BBS4, a gene involved in the Bardet–Biedl syndrome (58). BBS4 is a member of the recently discovered BBSome, which is required for ciliogenesis and hypothesized to function in IFT and vesicular transport to the cilium (59). The evaluation of the cellular function of USH proteins in photoreceptors suggests the participation of these proteins in the transport through the connecting cilium (21,48,60,61). Also, there is growing evidence that BBS molecules, IFT proteins and USH proteins participate in microtubule-based vesicular transport through the cytoplasm (21,53,59,62). The molecular links of Nlp isoB with the...
Figure 6. Centrosomal localization of Nlp isoB, USH2A_tail and lebercilin in ARPE-19 cells. When expressed alone, mRFP–Nlp isoB (red signal) was localized to the mother centriole of the centrosome (A), eCFP-USH2A_tail (green signal) was localized to the nucleus and the centrosome (indicated by an arrow and in the inlay) (B), whereas eYFP-lebercilin (green signal) was localized at the centrosome and the microtubule network of the cell (C). After co-expression of Nlp isoB and USH2A, both proteins were localized at the mother and daughter centriole of the centrosome, confirming the interaction between both proteins (D–D"). Co-expression of Nlp isoB and lebercilin showed co-localization of both proteins at the centrosome and non-centrosomal MTOCs (yellow signal; E–E"). Lebercilin in addition partly localized at the microtubule network (E–E"). Nuclei were stained with DAPI (blue signal).

Figure 7. Localization of endogenously expressed Nlp and lebercilin in ARPE-19 cells by immunocytochemistry using the anti-Nlp and anti-lebercilin antibodies (green signals) and anti-acetylated tubulin antibodies as an axoneme and microtubule marker (red signal). Nlp (A) and lebercilin (B) were present at the basal body of primary cilia and both proteins were found at the midbody region of ARPE-19 cells during telophase (C and D). Nuclei were stained with DAPI (blue signal).
Figure 8. Mutations in lebercilin affect the interaction with Nlp isoB. A quantitative liquid β-galactosidase assay shows an enhanced interaction of Nlp isoB and lebercilin Q279X when compared with wild-type lebercilin and a reduced interaction of Nlp isoB and lebercilin P493TfsX1 (A). When expressed alone in ARPE-19 cells, mRFP–lebercilin wild-type (red signal) was localized to the basal body, the primary cilium and the microtubule network of the cell (B), mRFP-lebercilin Q279X was localized to the basal body and primary cilium (C) and mRFP-lebercilin P493TfsX1 was localized to the microtubule network of the cell (D). Co-expression of Nlp isoB and lebercilin showed co-localization of both proteins at the basal body (yellow signal; E, inset). In addition, lebercilin localized to the primary cilium and the microtubule network in the cell periphery (red signal; E’–E”, inset). Upon co-expression of Nlp isoB and lebercilin Q279X, both proteins co-localize at the basal body (yellow signal; E”, inset). The mutant lebercilin was not found in the primary cilium (F–F”, inset). After co-expression of Nlp isoB and lebercilin P493TfsX1, no co-localization of these proteins was observed (G–G”, inset). Nuclei were stained with DAPI (blue signal).
USH2AisoB and lebercilin proteins qualified these as common denominators in the associated retinal protein network, suggesting that common molecular processes are disrupted in the retinas of USH2A and LCA5 patients. We indeed were able to determine that protein truncating mutations in lebercilin affect the interaction with Nlp, but as downregulation of expression of any of these proteins by RNAi had no effect on the localization of the interacting partner, the physiological defect is more complex and remains to be identified. The USH2A-associated protein network, the lebercilin interactome (29) and the Nlp interactors p150glued and Polo-like kinase 1 (Plk1) (32,36,37) provide multiple links with centrosomal and ciliary processes and pathways that are now potentially connected through Nlp.

Recently, we and others showed the presence of the Usher protein network in a periciliary collar-like structure at the apical part of the photoreceptor inner segments, analogous to the periciliary ridge complex of Xenopus photoreceptor cells, and the Usher proteins were hypothesized to function in the transport and docking of vesicles containing proteins for the outer segment (21,23,63). These cargo vesicles originate from the trans-Golgi network of the photoreceptor cells and are thought to be transported along microtubules by cytoplasmic dynein through the inner segment to the apical membrane (21,64). Nlp might function in this vesicular transport by the direct association with the dynactin p150glued subunit of the dynactin—dynactin motor complex (37), explaining the presence of Nlp in the inner segments of photoreceptor cells. The periciliary region and also the basal body and centriole are thought to serve as a docking site from where ciliary proteins (e.g. IFT proteins and proteins involved in signalling) are further distributed (65). In such a model, Nlp would function as the molecular switch between intracellular and IFT transport. The effect of mutations in lebercilin on its interaction with NlpisoB and on its localization in the cilium is in line with this model.

Interestingly, Nlp is known to be phosphorylated by Plk1 (32,36,37), which is a key regulator in centrosome function. Plk1 also phosphorylates nucleophosmin (66), a centrosome-associated protein that acts in nucleocytoplasmic shuttling (67) and is a member of the previously identified lebercilin interactome (29). In addition, nucleophosmin associates with RPGR-ORF15, a protein involved in X-linked retinal degeneration (68). The Nlp-interacting protein p150glued (37) was found in a protein complex with RPGR-ORF15 and with lebercilin (29,69). The direct or indirect association of Nlp with several protein complexes in the region of the photoreceptor-connecting cilium stresses the importance of this protein in photoreceptor cell function.

In conclusion, our data show that USH and LCA are molecularly linked by the direct association of USH2AisoB and lebercilin with a novel isoform of Nlp, NlpisoB. Co-localization of these proteins at the basal body of photoreceptor cells and the current knowledge on the function of the existing Usher protein network(s) point towards an interdependent function for NlpisoB, USH2AisoB and lebercilin, possibly in cytoplasmic trafficking and ciliary transport of proteins involved in photoreception. We propose that at the basal body, NlpisoB could function as a molecular hinge connecting cytoplasmic transport mechanisms to the IFT transport machinery, suggesting an important role for Nlp in photoreceptor cell function.

The central position of Nlp in the protein network indicates that Nlp can be considered as an excellent candidate gene for USH, LCA or other (retinal) ciliopathies. However, no pathogenic mutations have been identified so far.

MATERIAL AND METHODS

Animals and tissues

In this study, mature Wistar rats and C57BL/6J mice housed in standard cages and receiving water and food ad libitum were used. Animal experiments were conducted in accordance with international and institutional guidelines. Bovine retinas were dissected from eye balls obtained from the local slaughterhouse (21).

Plasmids and antibodies

Affinity-purified polyclonal antibodies directed against Nlp were described earlier (32). Monoclonal antibodies recognizing centrins 1–4 (20H5), polyclonal antibodies directed against the cytoplasmic region of USH2A and affinity-purified polyclonal antibodies directed against lebercilin were described previously (17,19,29). The immunohistochemical stainings of Nlp and USH2A were specific. No staining was observed after pre-adsorption of the primary antibodies with the corresponding peptide epitope. In addition, primary Nlp antibodies did not recognize GFP-tagged ninein after western blot analysis (Supplementary data, S1). Anti-flag, anti-HA, anti-acetylated tubulin and anti-gamma tubulin antibodies were purchased from Sigma (Germany). Anti-HA beads were purchased from Roche (Germany). Secondary antibodies for immunohistochemistry and western blot analysis were purchased from Molecular Probes—Invitrogen (USA), Rockland (USA) and Jackson ImmunoResearch Laboratories (USA). cDNA encoding (parts of) the cytoplasmic region of human USH2A (GenBank NP_996816) (amino acids 5064–5202, 5064–5168, 5064–5196, 5124–5196, 5124–5202 and 5169–5202) were cloned in the pDONR201 vector using the Gateway cloning system (Invitrogen) according to manufacturers’ instructions. cDNA fragments of the human Nlp gene were amplified by using IMAGE clone IRATp970C1131D (RZPD, Germany) as a template. By using the Gateway technology (Invitrogen), cDNAs encoding human full-length NlpisoB (amino acids 1–1033) (GenBank EU718622), NlpisoA (amino acids 1–1382) (GenBank NM_025176), the predicted EF-hand domains (amino acids 11–39, 200–228 and 237–265), the predicted IF domain (amino acids 656–925) of NlpisoB and deletion constructs for NlpisoB encoding amino acids 599–1033, 656–1033 and 656–825 were cloned in the pDONR201 vector. Lebercilin fragments (GenBank NP_859065) encoding amino acids 1–95, 96–305, 306–490 and 491–697 were amplified by PCR, using pDONR201-lebercilin as a template, and cloned in pDONR201 (29).

Yeast two-hybrid analysis

A GAL4-based yeast two-hybrid system (HybriZAP, Stratagene, USA) was used to identify proteins that interact with the
cytoplasmic region of USH2AisoB (amino acids 5064–5202) and proteins that interact with lebercilin (amino acids 1–697), using methods previously described (70) with minor variations. Yeast strain PJ69-4α (71) was used as a host, which carried the HIS3 (histidine), ADE2 (adényne), MEL1 (α-galactosidase) and LacZ (β-galactosidase) reporter genes. The DNA-binding domain fused to the human USH2A cytoplasmic region (pBD-USH2A_tail) and to the full-length lebercilin (pBD-lebercilinβ) was used as a bait for screening a human oligo-d(T) primed retina cDNA library containing 1.9 × 10^8 primary clones fused to the activation domain (pAD). In total, 1.1 × 10^7 clones (USH2A) and 5.0 × 10^7 clones (lebercilin) were plated on amino acid dropout plates lacking Trp, Leu and His (SD -WLH plates), containing 1 mM 3-aminotriazol, and selected for growth. Clones were then patched on medium additionally lacking adenine (SD-WLHA plates) and selected for growth and α-galactosidase activity by the activation of the MEL1 reporter gene. The latter was done using 20 µg/ml of the chromogenic substrate 5-bromo-4-chloro-3-indolyl α-D-galactopyranoside (X-α-Gal) in the dropout plates and selecting the yeast cells that developed a blue-green colour due to hydrolysis of X-α-Gal by the secreted α-galactosidase enzyme. Further selection of positive clones was based on β-galactosidase activity by the activation of the LacZ reporter gene, which was detected by a filter lift assay as previously described (70).

Liquid β-galactosidase assay

Three independent clones of PJ69-4α yeast cells co-transformed with pAD-NlpisoA full length or NlpisoB full length and pBD-lebercilin full length were cultured in selective medium lacking Trp and Leu (SD-WL). After two overnight incubation steps at 30°C, the optical density of the cultures was determined at a wavelength of 600 nm. Cell lysis and subsequent colourimetric reactions were performed by using the Yeast β-galactosidase assay kit (Pierce, USA) according to the manufacturer’s instructions. Absorbance was measured at a wavelength of 420 nm.

Expression analysis

The expression of NlpisoA and NlpisoB was examined by performing semi-quantitative RT–PCR analysis on RNA from human fetal and adult tissues as described before (72). Primers used for the detection of the transcripts encoding NlpisoA are 5'-GAGGGGGAGACAAAAATAGC-3' and 5'-TCTGAAATGGTCACACAGTGC-3'. For detection of NlpisoB, the following primers were used: 5'-ACCTGCAACGACATCA GACTG-3' and 5'-TGATTGTCACACTGGCTG-3'. Samples were taken after 25, 30 and 35 cycles.

Calcium-binding assay

GST-fusion proteins of the predicted Nlp EF hands 1 (amino acids 11–39), 2 (amino acids 200–228) and 3 (amino acids 237–265) were produced by transforming BL21-DE3 cells with pDEST15-Nlp EF hands 1, 2 and 3. Pre-cleared lysates were separated on pre-casted 4–12% NuPage gradient gels (Invitrogen) and subsequently blotted onto nitrocellulose membranes. The membranes were incubated for 10 minutes at room temperature with 10–20 µCi/1 45CaCl2 (New England Nuclear, USA) in 10 mM imidazole, pH 6.8, and 60 mM KCl. The blots were washed twice for 5 minutes with 50% ethanol, dried and subsequently exposed to a radiation sensitive film (Kodak, Germany) (73).

GST pull-down assay

In order to produce GST-fusion proteins, BL21-DE3 cells were transformed with pDEST15-USH2A_tail (amino acids 5064–5202), pDEST15-NBC3_tail (amino acids 1119–1214) and pDEST15-NlpisoB (amino acids 1–1033). Cells were induced at 30°C for 4 h with 0.5 mM IPTG and subsequently lysed with STE buffer (1% Sarkosyl, 1% Triton X-100) supplemented with protease inhibitor cocktail (Roche). Lysates were pre-cleared and incubated at 4°C for 16 h with glutathione Sepharose 4B beads (Amersham Biosciences, USA). Beads with bound fusion proteins were washed twice with lysis buffer and three times with TBS with 1% Triton-X-100 and 2 mM DTT. During each washing step, samples were incubated on a rotating wheel at 4°C for 5 min. The amount of bound GST-fusion protein was verified on a 10% SDS–PAGE gel stained with Gelcode Blue Stain Reagent (Pierce). HA-tagged NlpisoB, 3 × flag-tagged lebercilin, 3 × flag-tagged USH2A_tail and 3 × flag-tagged NBC3_tail were produced by transfecting COS-1 cells with, respectively, pcDNA3-HA-NlpisoB, p3xflag-lebercilin, p3xflag-USH2A_tail and p3xflag-NBC3_tail using Nucleofector kit V (Amaxa, USA) and program A-24 according to the manufacturer’s instructions. The pre-cleared supernatants were incubated overnight at 4°C in the presence or absence of 2 mM CaCl2 with equal amounts of blocked beads with GST or beads with GST-fusion proteins. After three washes with lysis buffer, the beads were boiled in 1 × SDS loading buffer. Protein complexes were resolved on 4–12% NuPage gradient gels (Invitrogen). For western blot analysis, proteins were electrophoretically transferred onto nitrocellulose membranes, blocked with 5% non-fat dry milk (Biorad, Germany) in PBST (0.1% Tween) and analysed with the appropriate primary and secondary antibodies in 0.5% milk in PBST. Bands were visualized using the Odyssey Infrared Imaging System (LI-COR, USA). Tagged molecules were detected by anti-HA or anti-flag mono- or polyclonal antibodies. As secondary antibodies IRDye800 goat-anti-mouse IgG (Rockland) and Alexa Fluor 680 goat-anti-rabbit IgG were used (Molecular Probes, USA).

Co-immunoprecipitation from COS-1 cells

HA-tagged NlpisoB and HA-tagged lebercilin were expressed by using the mammalian expression vector pcDNA3-HA/DEST. Flag-tagged intracellular region of USH2A, flag-tagged NlpisoB and flag-tagged LRRK2 were expressed by using p3xflag-CMV/DEST from the Gateway cloning system (Invitrogen). Both plasmids contain a CMV promoter. COS-1 cells were co-transfected by using Effectene (Qiagen, Germany) according to the manufacturer’s instructions. Thirty hours after transfection, cells were washed with PBS and subsequently lysed on ice in lysis buffer (50 mM Tris–HCl pH 7.5, 150 mM NaCl and 0.5% Triton X-100) supplemented
with complete protease inhibitor cocktail (Roche). HA-tagged NlpisoB and lebercilin were immunoprecipitated from cleared lysates overnight at 4°C by using rat monoclonal anti-HA beads (Roche), whereas flag-tagged USH2A_tail, NlpisoB and LRRK2 were immunoprecipitated by using polyclonal anti-flag antibodies (Sigma) and Protein A/G PLUS-Sepharose (Santa Cruz Biotechnology, USA). After four washes in lysis buffer, the protein complexes were resolved on 4–12% NuPage gradient gels (Invitrogen) and subsequently analysed on immunoblots as described for the GST pull-down assay. Bands were visualized by using the Odyssey Infrared Imaging System (LI-COR). Tagged molecules were detected by anti-HA or anti-flag mono- or polyclonal antibodies. As secondary antibodies IRDye800 goat-anti-mouse IgG (Rockland) and Alexa Fluor 680 goat-anti-rabbit IgG were used (Molecular Probes).

Co-immunoprecipitations from bovine retinal extracts

For immunoprecipitations, bovine retinas from a local slaughterhouse were used. Retinas were lysed by sonification for 10 s in lysis buffer (50 mM Tris–HCl pH 7.4, 150 mM NaCl, 0.5% Nonidet P-40 and 1 mM sodium-orthovanadate) supplemented with complete protease inhibitor cocktail (Roche). Lysates were pre-cleared and incubated for 16 h at 4°C with mouse monoclonal anti-rabbit IgGs, rabbit polyclonal Nlp antibodies or polyclonal rabbit lebercilin antibodies. Protein-antibody complexes were coupled to Protein A/G PLUS-Sepharose beads (Santa Cruz) for 2 h at 4°C. After incubations, the beads were pelleted and washed three times with lysis buffer. Beads were boiled and proteins were resolved on 4–12% NuPage gradient gels (Invitrogen) and subsequently analysed on immunoblots as described for GST pull-down. Bands were visualized using the Odyssey Infrared Imaging System (LI-COR).

Co-localization analyses in ARPE-19 cells

To determine the cellular localization of the cytoplasmic region of human USH2A, full-length lebercilin and full-length NlpisoB in ARPE-19 cells, cDNAs encoding the region of USH2A in pDEST501 were cloned by using the Gateway cloning technology (Invitrogen), resulting in N-terminally fused eCFP-USH2A. Nlp was cloned in pDEST733, resulting in N-terminally fused mRFP-NlpisoB. Lebercilin was cloned in pDEST504, resulting in C-terminally fused lebercilin-eYFP. ARPE-19 cells were co-transfected with pDEST733-NlpisoB and pDEST501-USH2A_tail or pDEST504-lebercilin by using Effectene (Qiagen) according to the manufacturer’s instructions. Twenty hours after transfection, cells were washed with PBS, fixed with 4% paraformaldehyde and mounted with Vectashield containing DAPI (Vector Laboratories, Inc., UK). Images were taken with an Axioplan2 Imaging fluorescence microscope (Zeiss, Germany) equipped with a DC350FX camera (Zeiss) and processed using Adobe Photoshop (Adobe Systems, USA).

Supplementary material

Pre-embedding immunoelectron microscopy

Immunoelectron microscopy was performed on isolated mouse retinas as previously described (21). In short, vibratome sections through mouse retina were stained by primary antibodies against Nlp and visualized by appropriate secondary antibodies (Vectastain ABC-Kit, Vector, UK). After fixation with 0.5% OsO4, specimens were embedded in araldite and ultrathin sections were analysed with an FEI Tecnai 12 Biotwin transmission electron microscope (FEI, The Netherlands).

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