A Complex of BBS1 and NPHP7 Is Required for Cilia Motility in Zebrafish

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Abstract

Bardet-Biedl syndrome (BBS) and nephropathies (NPH) are hereditary autosomal recessive disorders, encoded by two families of diverse genes. BBS and NPH display several overlapping phenotypes including cystic kidney disease, retinitis pigmentosa, liver fibrosis, situs inversus and cerebellar defects. Since most of the BBS and NPH proteins localize to cilia and/or their appendages, BBS and NPH are considered ciliopathies. In this study, we characterized the function of the transcription factor Nphp7 in zebrafish, and addressed the molecular connection between BBS and NPH. The knockout of zebrafish bbs1 and nphp7.2 caused similar phenotypic changes including convergent extension defects, curvature of the body axis, hydrocephalus, abnormal heart looping and cystic pronephros, all consistent with an altered ciliary function. Immunoprecipitation assays revealed a physical interaction between BBS1 and NPHP7, and the simultaneous knockdown of zbbs1 and znphp7.2 enhanced the cystic pronephros phenotype synergistically, suggesting a genetic interaction between zbbs1 and znphp7.2 in vivo. Deletion of zbbs1 or znphp7.2 did not compromise cilia formation, but disrupted cilia motility. Although NPHP7 has been shown to act as transcriptional repressor, our studies suggest a crosstalk between BBS1 and NPHP7 in regulating normal function of the cilium.

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Introduction

Autosomal recessive cystic kidneys are typically part of complex syndromes involving multiple organs. Since most gene products mutated in these syndromes localize to the cilium, an abnormal function of the cilium has been implicated in their pathogenesis. Cilia are specialized microtubule-based organelles attached to most vertebrate cells. The primary cilium is thought to function as a mechano- and/or chemosensor, which detects signals from extracellular environment and transmits them to the inside of the cell. Mutations of numerous ciliary proteins lead to structurally or functionally abnormal cilia [1]. The resulting defects are closely related to developmental and degenerative disorders, collectively termed ‘ciliopathies’ [2]. In kidney epithelial cells, the primary cilium is involved in important signalling cascades such as Hedgehog, Wnt, planar cell polarity, and calcium-dependant signalling pathways [3]. Cilia-related cystic kidney diseases include autosomal dominant as well as recessive polycystic kidney disease, including the Bardet-Biedl syndrome (BBS) and nephronophthisis (NPH) [4].

Bardet-Biedl syndrome (BBS) is a genetically heterogeneous autosomal recessive disorder (BBS 1-17) [5], symptoms of which include obesity, renal degeneration, anosmia, post-axial polydactyly, cognitive impairment, hypogonitalism, renal and cardiovascular anomalies, diabetes mellitus and hypertension [6,7]. The BBS proteins have been reported to form a stable multiprotein complex called BBSome (BBS1, 2, 4, 5, 7, 8 and 9), and to play a role in ciliogenesis and cilia maintenance [8,9,10]. NPH is an autosomal recessive cystic kidney disease associated with retinal degeneration, cerebellar hypoplasia, liver fibrosis, situs inversus, and mental retardation [11,12,13]. Mutations in 15 genes have been linked to NPHP so far (OMIM 614845). The proteins encoded by NPHP genes are called nephrocystins, and are highly conserved among species. They are involved in various cellular signalling events, employing the non-canonical Wnt, sonic hedgehog and planar cell polarity signalling pathways [11,14]. Recent studies indicate that most of BBS and NPHP proteins localize to the cilia/basal body complex [15,16,17,18], providing a structural basis for the overlap of the symptoms exhibited by BBS and NPH patients, including renal cysts and retinitis pigmentosa, liver fibrosis, situs inversus and mental problems, and supporting the notion that BBS and NPHP protein family share functional pathways [19,20].

During the last decade zebrafish has emerged as a powerful animal system for studying ciliopathies [21,22]. In the present study, we analyzed the biochemical interaction between BBS1 and NPHP7, and investigated this interaction in zebrafish. BBS1 is one of the seven members of the BBSome, which regulates vesicular trafficking of proteins to the ciliary membrane [10,23]. Mutations
of BBS1 in humans have been reported as one of the most frequent causes of BBS, implying a significant function of BBS1 [6,24,25,26]. BBS1 is highly expressed in the kidney [24]. NPHP7/GLIS2 (Gli-similar 2) is a member of the Gli-related Krüppel-like zinc-finger (ZF) transcription factor subfamily [27,28]. Gli2 is essential for maintaining renal functions by regulating genes which are involved in epithelial-to-mesenchymal transition, fibrosis and apoptosis [29,30]. In adult mouse kidneys, Gli2 was detected in epithelial cells of the renal tubule and Bowman’s capsule [29], Gli2 lacZ/lacZ mutant mice developed renal atrophy, fibrosis and glomerular cysts, thus resembling some of the key features of NPH [29,30]. Both BBS1 and GLIS2 are present in cilia [23,29,31].

Morpholino oligonucleotide (MO)-mediated knockdown of zbbs1 and znphp7.2 in zebrafish embryos caused phenotypic changes characteristic for “ciliopathies”. Detailed analysis of cilia revealed that despite normal morphology, the cilia in the pronephric tubule and the nasal pit of zBbs1- and zNphp7.2-depleted embryos showed aberrant motility, suggesting that the observed phenotypes are due to the disruption of the normal ciliary beating pattern.

**Results**

**BBS1 and NPHP7 interact with each other**

To investigate the interconnection between BBS and NPH protein families, we tested the interaction between NPHP7 and the BBS family members BBS1-12, using tagged versions of human proteins overexpressed in human embryonic kidney (HEK) 293T cells. We took interest in NPHP7 for the reason of being a transcription factor not having been very well characterized so far. Precipitation of BBS1 immobilized NPHP7 most extensively, suggesting that this interaction may be relevant in vivo. Since BBS1 has been characterized as a component of the BBSome, involved in ciliary transport processes, we decided to clarify the role of the BBS1/NPHP7 complex. Fig. 1A shows the interaction between human BBS1 and NPHP7. While BBS1 lacks apparent conserved functional domains, NPHP7 is a transcription factor with 5 zinc finger (ZF) domains responsible for DNA binding and transcriptional regulation [32]. To identify the domain of NPHP7 responsible for the interaction with BBS1, we generated three truncation constructs, Flag-NPHP7 1–156, Flag-NPHP7 141–359 and Flag-NPHP7 346–525 (Fig. 1B). We found that the N-terminus (Flag-NPHP7 1–156) of NPHP7 interacts with V5-BBS1 (Fig. 1A). Unfortunately, the lack of suitable antibodies for NPHP7 prevented us from confirming this interaction in vitro. Therefore, we turned to zebrafish, a genetically tractable animal model, to search for a potential genetic interaction between these two gene products.

**zbbs1 and znphp7 are expressed in various tissues in the zebrafish**

The zebrafish homologue of human BBS1 has been previously described [33]. Homology blast searches with the human protein as a query identified 2 potential zebrafish Nphp7 homologues. The genes encoding these proteins are located on chromosome 22 and 3, which we named znphp7.1 (Genbank KF054060, Zv9: ENSDARG00000078388) and znphp7.2 (Genbank KF054061, Zv9: ENSDARG0000007836), respectively; subsequent genome annotations (genome assembly ZV9) confirmed our findings. Protein alignments showed that zNphp7.1 sequence had 43.9% identity and 50.8% similarity to the human NPHP7/GLIS2; zNphp7.2 was 51.4% identical and 60.2% similar to the human gene product (Fig. 2A). As for the Gli family, the domain with the highest identity was the ZF domain: zNphp7.1 and zNphp7.2 were 89.3% and 91.3% identical with the human ZF domain of human GLIS2, respectively. Previous studies showed ubiquitous expression of BBS1 in human organs including fetal tissue, testis, retina and adipose tissue with the highest expression in the kidney [24]. Gli2 mRNA was also most abundantly expressed in mouse kidney with low levels in heart, lung, placenta, prostate, colon, and brain [27,28]. In zebrafish whole mount in situ hybridization with various antisense RNA probes of zzb1 and znphp7.1 and znphp7.2 did not show consistently reproducible results. Semi-quantitative RT-PCR revealed that znphp7.1 and znphp7.2 are expressed maternally, at 6 hpf and 24 hpf whereas zbb1 is expressed at 24 hpf. A second maternal splice variant of znphp7.2 (Transcript 2) was observed and confirmed by sequencing (Fig. 2B and Fig. S1). The sequence alignment showed that 1 amino acid is substituted (S99R) and 18 amino acids (aa 101–118) are not coded for in transcript 2 of znphp7.2 (Fig. S1). In addition, semi-quantitative RT-PCR from various organs of adult zebrafish showed that zbb1, znphp7.1 and znphp7.2 are expressed in kidney, eye and testis. However, znphp7.1 and znphp7.2 are additionally expressed in the heart, gut and muscle (Fig. 2C).

**Depletion of zbb1 and znphp7 in zebrafish embryos causes ciliopathic phenotypes including pronephric cyst formation**

To address the role of zBbs1 and zNphp7 in zebrafish kidney development, we performed knockdown experiments by injection of MO, targeting zbb1 translational start codon (AUG MO) and exon 2 splice donor site (SP MO) of zbb1 to deplete zBbs1 [33] (Fig. S2A). To test the efficacy of zbb1 AUG MO, we performed an *in vitro* transcription and translation (TNT) assay. This showed that zbb1 AUG MO efficiently interfered with zBbs1 protein expression (Fig. S2B). Both MOs showed similar phenotypes, which implies that the observed phenotypes were knockdown-specific. However, the AUG MO caused less general developmental side effects, and was therefore used in all subsequent experiments. To examine the effect of depletion of zBbs1 on kidney development in *zivo*, we used the transgenic fish line Tg(wt1b:EGFP) [34]. At 55 hpf, zbb1 morphants displayed cysts with distension of the neck segment of the pronephric tubule (Fig. 3E and G) indicating that zBbs1 is required for normal zebrafish pronephros development. The overall appearance of the morphant embryos was categorized by the level of dysmorphology ranging from wild-type-like (WT) to increasingly severe dysmorphology (D1 to D3) (Fig. 3G and Fig. S3). The grade of dysmorphology and the number of cysic morphants showed a MO dose dependency (Fig. 3G). In addition, we observed the previously described effects of zBbs1-depletion manifesting in hydrocephalus and curvature of the body axis (Fig. 3B and Fig. S3), in convergent extension (CE) defects (Fig. 4), and in defects of left-right (L-R) asymmetry, i.e. organ laterality, such as heart looping (Fig. 5A and B; Fig. S5A and B), and situs of liver and pancreas (Fig. S5C and D) [35,36,37]. Moreover, zbb1 deficient embryos showed defects in eye development displaying reduced eye size and impaired layer formation of the retina (Fig. S8) similar to the knockdown of the orthologue of retinitis pigmentosa gene 2 [38,39]. Altogether, the phenotypes detected in zbb1 morphants were reminiscent of ciliopathy-associated phenotypic changes previously described in zebrafish [33,33].

Depletion of zNphp7.1 by either translational initiation site blocking AUG MO, or by a MO binding to the exon 5 splice donor site did not result in any specific phenotype readily distinguishable from wild-type (data not shown). For knockdown of znphp7.2, 2 different SP MOs, one targeting the exon 3 splice
donor site (SP1) and the other targeting the exon 6 splice acceptor site (SP2) of znphp7.2 gene, were used (Fig. S4A). Both MOs caused similar phenotypes, confirming the specificity of the knockdown. However, the SP1 MO targeting exon 3 showed higher efficiency at lower concentrations and was used subsequently.

znphp7.2 MO injections in 1-cell embryos caused aberrant splicing of intron 3 in znphp7.2 pre-mRNA leading to 2 splice products both of which cause an early truncation within the first ZF domain eliminating all ZF domains together with the C-terminus of zNphp7.2 (Fig. S4B–D).

The phenotype of znphp7.2 depletion includes curvature of the body axis and hydrocephalus (Fig. 3), retinal degeneration with disturbed retinal layering and reduced eye size (Fig. S8) and cystic pronephros (Fig. 3C, F and H), which are very similar to the phenotype of zbb1 morphants pointing to defects in cilia function.

Co-injection of in vitro transcribed zbb1 or znphp7.2 mRNA together with the corresponding MO partially rescued the morphant phenotype (Fig. S6), confirming the specificity of the MO-mediated knockdown.

Figure 1. BBS1 interacts with N-terminus (amino acids 1–156) of NPHP7. (A) Full-length or FLAG-tagged truncation of NPHP7 (NP_115964.2) was co-expressed with V5-tagged BBS1 (V5.BBS1). Flag-tagged CD2AP (Flag.CD2AP) was used as a negative control. After precipitation using anti-Flag M2 agarose beads, the precipitates were analyzed for the presence of V5-tagged BBS1 with anti-V5 antibodies by western blotting. V5.BBS1 was precipitated by Flag.NPHP7 1–156. Arrows indicate precipitated Flag-tagged proteins. In the lane of CD2AP, there was an additional non-specific band at the position of 55 kD. The short black lines in the dashed box indicate protein size markers (M). (B) Schematic description of V5-tagged full-length BBS1, Flag-tagged full-length (Flag.NPHP7) and 3 different fragments (Flag.NPHP7 1–156, Flag.NPHP7 141–359, Flag.NPHP7 346–524) of NPHP7. Flag.NPHP7 141–359 contains 5 zinc finger (ZF) domains.

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To exclude that pronephric cyst formation was attributable to cloaca malformation, FITC-conjugated fluorescent dextran (70 kD) was injected into the common cardinal vein of control and morphant embryos with pronephric cysts at 96 hpf. The excretion of fluorescent dextran with the urine was observed shortly after the injection in 22 control embryos, 16 z*bb1* and 21 *znphp7.2* morphants (Fig. S7). Only, 2 *znphp7.2* morphants showed cloaca blockage. Therefore, we conclude that pronephric cyst

**Figure 2. Expression of *zbb1* and *znphp7.** (A) Identification of 2 NPHP7 homologues in zebrafish: Zebrafish Nphp7.1 (*znphp7.1*) and zebrafish Nphp7.2 (*znphp7.2*) consist of 446 and 489 amino acids respectively. Amino acid sequence alignment showed that *znphp7.1* shares 43.9% identity and 50.8% similarity with the human NPHP7/GLIS2 (*hNPHP7*); *znphp7.2* was 51.4% identical and 60.2% similar to the human homologue. The ZF domains of *znphp7.1* and *znphp7.2* were 89.3% and 91.3% identical with those of the human homologue, respectively. (*, completely conserved; •, identical in 2 sequences or belonging to same type of amino acid group in 2 or 3 sequences) (B) Semi-quantitative RT-PCR reveals maternal transcript expression for *znphp7.1* and *znphp7.2* whereas *zbb1* is not expressed maternally nor at 6 hpf. 2 maternal splice products were identified for *znphp7.2* (open arrowhead: Transcript 1; filled arrowhead: Transcript 2). The transcript 2 of *znphp7.2* is expressed only maternally. Sequencing of the lower splice product revealed an excision of 18 bp corresponding to amino acid (aa 101–118) (Fig. S1). (C) Semi-quantitative RT-PCR with organ specific cDNA from adult zebrafish indicated that *zbb1* is expressed in kidney, eye and testis. *znphp7.1* and *znphp7.2* are expressed in other organs including kidney, eye, heart, testis, gut and muscle.

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formation of zbbs1 and znphp7.2 morphants was not caused by disruption of cloaca development (Fig. S7) [40].

Knockdown of znphp7.2 also caused CE defects judged by a wider body gap angle in relation to the stage of somitogenesis which translates into a shorter body axis and broader somites on in situ hybridization with probes for krx20 and myod1 (Fig. 4). Cilia-driven fluid flow in the Kupffer’s vesicle (KV) triggers symmetry breaking events during zebrafish development [37,41,42,43,44]. Defects in L-R asymmetry were observed in znphp7.2 morphant embryos for early laterality markers like lefty2 (Fig. S5A and B) and in MO dose dependent manner for heart looping (Fig. 5A and C) and the situs of liver and pancreas (Fig. S5C and D). This shows the involvement of zbbs1 and znphp7.2 in establishing organ laterality most likely via cilia function in KV.

zbbs1 and znphp7.2 show a genetic interaction in cystogenesis

The co-immunoprecipitation suggested a link between BBS1 and NPHP7 (Fig. 1). To confirm this interaction in vivo, we
performed combined knockdown of \( \text{z} \text{bbs1} \) and \( \text{znphp7.2} \) in zebrafish. Individual injections of a suboptimal dose of MO, \( \text{z} \text{bbs1} \) (\( \text{z} \text{bbs1} \) MO 0.2 mM+Cont MO 0.1 mM) and \( \text{znphp7.2} \) (\( \text{znphp7.2} \) MO 0.1 mM+Cont MO 0.2 mM), displayed 10% and 17% of pronephric cyst formation, respectively. However, combined knockdown of \( \text{z} \text{bbs1} \) and \( \text{znphp7.2} \) (\( \text{z} \text{bbs1} \) MO 0.2 mM+\( \text{znphp7.2} \) MO 0.1 mM) synergistically increased the number of embryos with cystic pronephros to 59%, suggesting a genetic interaction between \( \text{z} \text{bbs1} \) and \( \text{znphp7.2} \) (Fig. 6). To keep the total amount of injected MO the same for all injections, the suboptimal doses of \( \text{z} \text{bbs1} \) and \( \text{znphp7.2} \) were combined with Cont MO to achieve 0.3 mM as the final MO concentration for injection.

**Knockdown of \( \text{z} \text{bbs1} \) and \( \text{znphp7.2} \) influences cilia motility**

As indicated above, knockdown of either \( \text{z} \text{bbs1} \) or \( \text{znphp7.2} \) led to defects in cardiac looping in zebrafish embryos. One potential cause of this defect could be the defective “anlage” of the KV [45]. Indeed, our examinations revealed that depletion of \( \text{z} \text{Bbs1} \) causes defects in KV formation in a substantial number of embryos.
consistent with previous findings [33], while morphological examination of znphp7.2-depleted zebrafish embryos showed normal formation of KV (Fig. 7A and B). Further examinations of the ciliogenesis in KV of znphp7.2 knockdown embryos were performed using immunostaining for acetylated tubulin. This result indicated that development of cilia in the KV appeared shorter compared to control, however, the overall distribution of cilia in KV was unchanged (Fig. 7C and D). In addition, cilia formation in the pronephric tubule of zBbs1- and zNphp7.2-depleted embryos was not disturbed even though the cilia in zBbs1-depleted embryos were longer, and the cilia in zNPHP7.2-depleted embryos were shorter at significant level compared to control (Fig. 7E and F). The ciliary morphology and length in the pronephric tubule were also assessed in the combined knockdown condition of both zBbs1 and zNphp7.2 which showed the synergistic enhancement of cystic pronephros phenotype. In this condition, the experimental results showed that single and simultaneous knockdown of zBbs1 and zNphp7.2 does not affect cilia length in pronephric tubule (Fig. 7G and H). This data raised the question, whether the motility of cilia in morphant zebrafish embryos was impaired. Therefore, we assessed cilia motility by live embryo confocal imaging using the Tg(bact:zbt136-GFP) transgenic line which labels cilia with GFP in various organs [46]. We used this fish line to be able to detect subtle changes in motility within the population of cilia of one tubule. The movies were obtained along the whole pronephric tubule. For better comparison, three regions were defined as anterior, middle, and posterior (Fig. S9A). Regular beating movements of cilia in the lumen of the pronephric tubule in downstream direction, towards the cloaca, were observed in control zebrafish embryos (Movie S1, S2 and S3) as described previously [37]. The movies were recorded from morphant embryos with clear pronephric cysts. Strikingly, the cilia in zBbs1 (0.5 mM)- and zNphp7.2 (0.2 mM)-depleted zebrafish embryos exhibited greatly reduced motility and abnormal beating pattern (Movie S4 and S5, S6 for zBbs1 MO; Movie S7 and S8, S9 for zNphp7.2 MO). These results suggest that zBbs1 and zNphp7.2 are required for motility of cilia in pronephric tubule of zebrafish.

Furthermore, disturbed cilia movement was detected in the nasal pit of zBbs1 and zNphp7.2 morphants at 72 hpf, confirming that zBbs1 and zNphp7.2 play a role in cilia motility. This was monitored by ordinary videomicroscopy with a high-speed camera system (Movie S10, S11 and S12). Immuno-staining of cilia in the nasal pit with acetylated tubulin antibody shows that formation of cilia in the nasal pit is unimpaired in zBbs1 and zNphp7.2 morphant embryos (Fig. S9B).

To scrutinize the cause of the impaired motility of cilia, electron microscopy was performed to examine ultrastructure of cilia in the pronephric tubule of zBbs1 and zNphp7.2 morphants: ultrastructural...
defects, e.g. deficiency of dynein arms, could not be detected in pronephric cilia of both morphant embryos (Fig. 7I). Hence, we conclude that the disturbed motility of zbs1 and znphp7.2 morphant cilia is not necessarily associated with the axonemal ultrastructure of cilia [47].

To examine the interaction between zbs1 and znphp7.2, cilia beating in the pronephric tubule of morphants with combined knockdown condition was analyzed. Interestingly, the simultaneously zBbs1- and zNphp7.2-depleted embryos displayed severely impaired motility of cilia compared to zBbs1- or znphp7.2-deficient embryos (Movie S13, S14, S15, S16, S17, S18, S19, S20, S21, S22). In the movies of combined knockdown embryos, the arrows point to the individual cilia showing impaired movement. The movies were recorded from the single knockdown morphant embryos without pronephric cysts, whereas the movies were recorded from the combined injected morphant embryos showing clear pronephric cysts. This result supports the genetic interaction between zbs1 and znphp7.2 which was shown for pronephric cystogenesis in this study.

Figure 6. Genetic interaction between zbs1 and znphp7.2 in vivo. Injected zebrafish embryos were assessed for the incidence of pronephric cysts. (A) Cont MO injected embryos with normal pronephros. While the majority of zbs1 or znphp7.2 morphants (suboptimal dose) showed pronephros of normal morphology, those of combined knockdown exhibited pronephros with cysts. Cysts are marked by asterisks. (B) Pronephric cysts were detectable after individual injections of znphp7.2 MO at 0.1 mM and zbs1 MO at 0.2 mM with the level of 17% and 10% respectively whereas the combined knockdown caused pronephric cysts in 59% of microinjected embryos. The final MO concentration for injection was 0.3 mM in all conditions to keep total MO dose constant. Therefore suboptimal doses of zbs1 and znphp7.2 MO were combined with Cont MO to obtain this final concentration of 0.3 mM. The numbers in the brackets (n) are the numbers of total embryos which were examined.

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Therefore, we conclude that the cystic pronephros phenotype in the knockdown of zbb1 and znphp7.2 is due to the aberrant motility of cilia and disturbed fluid flow in the pronephric tubule [37,48].

**Discussion**

Multiple ciliopathy-associated genes have been identified that are involved in syndromic cystic kidney disease. Despite phenotypic overlap, the functional relationship and interactions are not well understood yet [4]. The clinical features and the severity of phenotypic changes vary with different identified mutations within one gene, but also with the mutational load and the combination of mutations in two or more genes of ciliopathy-associated gene families [49,50,51,52]. Based on the clinical overlap, we speculated that the similarity in phenotypes is reflected by functional interaction[s] between BBS and NPHP gene products. Immuno-precipitation assays identified an interaction between BBS1 and NPHP7. The interacting domain was located to the N-terminus of NPHP7. To examine the in vivo relevance of this interaction, we identified the zebrafish homologues of NPHP7, zNphp7.1 and zNphp7.2, likely the result of zebrafish genome duplication events [53]. The protein sequence identity between zebrafish and human NPHP7.2 and NPHP7.1 remains unclear. The phenotypic changes (curved body axis, hydrocephalus, retinal degeneration and pronephric cyst formation) induced by knockdown of znphp7.2 were comparable with depletion of other ciliopathy gene products [37]. We excluded cyst formation on the grounds of cloaca malformation by fluorescent dye injection and urinary excretion experiments. There is no detailed description of the clinical phenotype available in the Canadian Oji-Cree kindred with the first known human NPHP7/GLIS2 mutation (homozygous splice-site mutation, c.755+1G>T) and recently found second mutation in the Turkish patient (c.523T>C, p.E175R) [29,54]. In zebrafish, knockdown of znphp7.2 induced defects in left-right asymmetry of organ situs consistent with the concept that mutations of NPHP7 belong to the growing family of ciliopathies. Kupffer’s vesicle revealed a normal size and shape in zphp7.2 compared to control, there was no change in ciliary length observed in the set of experiments with combined MO injection. Furthermore, the general distribution and electron microscopic ultrastructure revealed no difference between morphant and control. However, our results of video-microscopy and confocal microscopy revealed an astonishingly impaired ciliary motility in pronephric tubule and nasal pit of znphp7.2 and zbb1 morphants, suggesting that the ciliopathy-associated phenotypes caused by the knockdown of either zbb1 or znphp7.2 are due to the disruption of ciliary motility [37].

In primary cilia dyskinesia (PCD) the impaired ciliary motility often goes along with changes in ultrastructure, e.g. deficiency of dynein arms, central pair complex or radial spokes, but there have been reports of unchanged ciliary ultrastructure with DANA11 [56]. With DANA11 one possible explanation is that the mutant DNAH11 protein retains its N-terminal domain, which is required for the correct incorporation of outer dynein arms [57]. Another key mechanism for regulation of ciliary motility is phosphorylation, i.e. phosphorylation/dephosphorylation of the dynein arms, through kinases and phosphatases that are anchored immediately adjacent to their axonemal substrates [58]. These kinases and phosphatases could be candidates regulated by events downstream of znphp7.2 and zbb1 without causing ultrastructural defects on electron microscopic level.

Previous reports localized BBS1 and GLIS2 to cilia [23,29,31]; this is particularly remarkable for the transcription factor NPHP7/ GLIS2 preferably found in the cell nucleus. Recently, the BBSome, structurally related to coat complexes, has been reported to mediate the transport of cargo to the ciliary compartment [23]. BBS1 interacts with other proteins such as BBS3, LepRb, Rabin8, and has been implicated in protein and membrane trafficking to cilia [10,59,60]. Thus, BBS1 might be involved in the transport of GLIS2 to the cilium, where posttranslational modifications of GLIS2 could take place similar to the mandatory processing of GLI transcription factors in the cilium [61,62]. Cleavage of GLIS2 has been described to be induced for example by p120 catenin [63]. Alternatively BBS1 could also facilitate the transport of GLIS2 from the cytosol to the nucleus. Curiously, other NPHP gene products such as NPHP2/Inversin [64] and NPHP15/CEP164 [65] have been shown to localize to both the cilium as well as the nucleus.

In conclusion, we demonstrate that depletion of Nphp7/GLis2 leads to a NPH-typical phenotype in zebrafish including defects in...
left-right organ asymmetry, further validating Glis2 as a legitimate member of the NPH gene family in spite of the limited description of human mutations. We also show that Bbs1 and Nphp7 are both required for normal ciliary motility, a pathway responsible for normal body asymmetry. Although the presence of BBS1/NPHP7 in the cilium suggests a cilia-specific function, this module could also regulate the expression of other gene products required for ciliary motility.

Methods

Zebrafish lines

Wild-type TL or TUAB zebrafish were maintained and raised as described (Westerfield, 1995). Dechorionated embryos were kept at 28.5 °C in E3 solution with or without 0.003% 1-Phenyl-2-thiourea (PTU, Sigma) to suppress pigmentation and staged according to somite number (som) or hours post-fertilization (hpf) (Westerfield, 1995).

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The transgenic fish line Tg(sp1:eGFP) was a kind gift of Christoph Englert [34]. The Tg(flact:Arrl3h–GFP) transgenic line we received thankfully from Brian Ciruna’s lab [46].

Zebrafish embryo manipulations

Embryos at 1-cell stage were microinjected with 4 nl of solution containing MO (MO; Gene Tools LLC) or in vitro transcribed capped mRNA, diluted in 100 mM KCl, 0.1% Phenol Red and 10 mM HEPES, pH 7.5. 2 different MOs targeting zebrafish bbs1 were used, a translation-blocking MO (zbb1 AUG MO; 5’-GGGAAACAGATGACATGTTGGTTTTG-3’) and an MO targeting against the splice donor site of exon 2 (zbb1 Ex2 MO; 5’-TGAAACTCAGCACAATGCATTAGGAAGA-3’) [33]. We designed the 2 independent MOs against znphp7.1 (XM_001340674.2 which was identified in Ensembl Zv7), targeted to translation initiation site (znphp7.1 AUG MO; 5’-GGTCCGTTCGACGATGCTGTTT-3’) and exon 5 splice donor site (znphp7.2 SP MO; 5’-TATAATATCCACAGTCTGGCACTGGCC-3’). 2 separate MOs against znphp7.2 were also designed and ordered, targeted to exon 3 splice donor site (znphp7.2 SPI MO; 5’-TATCCCTTACTAATCTGCTAC-3’) and exon 6 splice acceptor site (znphp7.2 Ex6 MO; 5’-CTCACGTTAAACAGTTACAGAGA-3’). In order to decrease the nonspecific effects of the reagents to zebrafish embryos, all MOs were co-injected with p53 MO (0.1 mM), p53 MO (0.1 mM) was also injected into all wild-type embryos as controls [66]. For combined knockdown experiment with 2 different MOs simultaneously, all MO injection samples were balanced with Cont MO (standard control MO, Gene Tools LLC). MO to have 0.3 mM in order to inject the same amount of MOs in total. The clone of zebrafish bbs1 homologue was obtained from ImaGenes GmbH. The full-length zbb1 was isolated by PCR and cloned into the vector pCS2+ for in vitro transcription/translation assay. To produce the zbb1 mRNA for rescue experiment, we modified the sequence of zbb1 with sequence-directed mutagenesis and generate a construct with 3 silent mutations in zbb1 AUG MO binding site (original sequence from start codon, atg tca tct gtt; modified sequence, atg tct tca gta). The full-length of znphp7.2 obtained by PCR using zebrafish embryonic cDNA was also inserted into pCS2+ (znphp7.1 (Genbank KF054060, Zv9: ENSDARG00000078388) and znphp7.2 (Genbank KF054061, Zv9: ENSDARG00000073861)). Phenotype rescue experiments were performed with in vitro transcribed 20 pg of full-length MO-resistant zbb1 mRNA or 10 pg of wild-type znphp7.2 mRNA (mMessage Machine kit, Ambion) which was co-injected with zbb1 AUG MO (0.4 mM) or znphp7.2 SPI (0.3 mM) respectively.

The body gap angle of embryos at 8–10 somite stage was measured to assess convergent extension movement [36]. KV areas of control and morphant embryos were measured at 8–10 somite stage using imageJ [http://rsb.info.nih.gov/ij/]. The average area measured in control embryos was set as 1.00 to determine other measurement of morphants.

All animal studies were approved by the Committee on Research Animal Care, Regierungsparkhülim Freiburg (55-9185.01/G-11/41).

Cell culture and transfection

For co-immunoprecipitations, HEK 293T cells (received from American Type Culture Collection, ATCC, Manassas, VA) were grown in DMEM with 10% FBS. The 70–90% confluent cells were split at a ratio of 1:5 and transfected the next day with plasmid DNA applying the calcium phosphate method as described previously [67]. The transfection was stopped after 6–8 h by replacing the media.

Co-immunoprecipitation

To construct Flag-tagged BBS1, NPHP7 and truncations of NPHP7/GLIS2, the cDNAs of full-length BBS1 and NPHP7 were amplified from the EST clone by PCR and cloned between MluI and NotI site of the Flag-pcDNA6. HEK (Human embryonic kidney) 293T cells were transiently transfected with plasmids by the calcium phosphate method, and lysed after incubation for 24 h in immunoprecipitation buffer containing 20 mM Tris-HCl (pH 7.5), 1% Triton-X 100, 25 mM NaF, 12.5 mM Na4P2O7, 0.1 mM EDTA, 50 mM NaCl, 2 mM Na3VO4, and protease inhibitors. After lysates were cleared by centrifugation at 15 000 g for 30 min at 4°C, the supernatants were incubated with 30 μl of anti-Flag-M2 beads for 3 h [68]. The beads were washed extensively with immunoprecipitation buffer, and the retained proteins were analyzed with western blot analysis. Antibodies used in this western blot analysis, included FLAG® monoclonal antibody produced in rabbit (Sigma, Hamburg, Germany), rabbit antibody to V5 (Serotec).

Eye size measurement and HE staining for eye sections

Eye size measurement and HE staining for eye sections were performed with 80 hpf and 96 hpf embryos, respectively. Image J (http://rsb.info.nih.gov/ij/) was used for measurement of the area of the eye. The embryos at 96 hpf that were sectioned for Hematoxylin and Eosin staining were fixed in BT-fix (4% PFA, 0.1 M Na2HPO4 buffer (pH 7.5), 3% sucrose, 0.12 mM CaCl2) at 4°C overnight. After being washed in PBS and taken through an ethanol dehydration series, the embryos were embedded in JB-4 resin (Polyscience Inc.) and sectioned at 3–5 μm using a microtome (Leica). Slides were stained for Hematoxylin and Eosin.

In situ hybridization and immunohistochemistry

In situ Hybridisation (ISH) was carried out as described previously [69]. Dig-labelled antisense RNA was transcribed with T3 or T7 RNA polymerase (Roche). The npy17 (mesyn, light polypeptide 7) antisense probes were generated by linearizing template plasmids with NotI and transcription with T7 RNA polymerase. krox20 (krox20) antisense probe was produced by linearizing template plasmids (pBS KS vector) with PstI and transcription with T3 RNA polymerase. myod1 (myogenic differen-
In pCRII-Topo, the vector was linearized by BamHI and transcribed by T7. β2 (β2f) in pCS2+ was linearized by ClaI and transcribed by T7; ins (insulin) and fosA3 (forkhead box A3) probes in pBS SK vector were linearized with XbaI and XhoI respectively and transcribed with T7 and T3 RNA polymerase respectively.

For immunohistochemistry, primary mouse monoclonal antibodies, anti-acetylated tubulin (Sigma) were used in a 1:500 dilution. Secondary anti-mouse antibodies labeled with Cy3 (Jackson ImmunoResearch and Molecular Probes) were used. Embryos were fixed in 1% DMSO and 4% PFA in PBS and stained with primary and secondary antibodies by conventional whole-mount antibody staining method. The length of stained cilia was measured with imageJ.

**RT PCR analysis**

The total RNA was extracted from entire zebrafish embryos at desired stage or from adult zebrafish organs (kidney, eye, heart, testis, gut and muscle) with RNeasy® MiniKit (Qiagen). cDNA was synthesized from the total RNA using ProtoScript® First Strand cDNA Synthesis Kit (Biolabs Inc.). These cDNAs were used for further PCR analyses. zebrafish eif2α (translation elongation factor 1 alpha) was amplified as a positive control. The following primers were used for corresponding PCR analysis: zbb1 (forward: 5’-tgtcctcagctggaag-3’; reverse: 5’-gcaacacggctgcgaaata-3’), znp7p7.1 (forward: 5’-ggtctctcgccgtagttg-3’; reverse: 5’-ccatggttaggctgcaat-3’), znp7p7.2 (forward: 5’-ggtctccagagctctc-3’; reverse: 5’-ggcaacagctgcaactc-3’), ztaf1 (forward: 5’-attcatacctgctgcaat-3’; reverse: 5’-atccagctctcaactc-3’)

**Fluorescent dextran injection**

For urine excretion assays, 5% FITC-conjugated dextran solution (70 kD) (Molecular Probes) was injected into the common cardinal vein of 96 hpf embryos anesthetized with 0.2 mg/ml tricaine (3-aminobenzoic acid ethylster, Sigma) in egg water.

**Transmission electron microscopy**

Zebrafish embryos at 35 hpf were fixed with PBS containing 1% PFA/2% glutaraldehyde for 1 h at RT and post-fixed with 1% osmium tetroxide (Polyscience) for 30 min at RT. The embryos were dehydrated in ethanol (50% and 60% for 10 min each) and transferred into 1% uranyl acetate (Polyscience) in 70% ethanol overnight at 4°C.

The embryos were subjected to further dehydration through a series of 80%, 90%, 98% and 100% ethanol. After washing the embryos with propylene oxide, they were embedded in Durcupan (Fluka). Ultra-thin sectioning was performed with a Leica EM UC6. Serial sagittal sections (60 nm) were conducted from rostral to caudal (from the beginning of the yolk extensions to copper-grids). Sections were observed and imaged in an electron microscope (LEO 906E; Carl Zeiss). All images were exported as TIFF files.

**Confocal microscopy with Tg(factAr113b-GFP) transgenic line**

All embryos were taken with LSM-Live Duo ZEISS LSM 510 DUO using ZEN 2010 software (Carl Zeiss, Inc.) and laser diode 488. Image acquisition for the moving cilia was 30 frames per second and the recorded movies are replayed in real-time. The movement of cilia in pronephric tubule was imaged with 55 hpf embryos of the Tg(factAr113b-GFP) transgenic line [43]. The embryos were incubated with 40 mM BDM (2,3-butanediol monoxime, Sigma-Aldrich) for 5 min to eliminate the interfering signal of blood flow, and then the mounted embryos in 1% low melting temperature agarose were covered with egg water (20 mM BDM) for imaging. The movies were taken with LD LCI Plan-Apochromat 25×/0.8 (zoom: 2.0).

**High-speed videomicroscopy**

The 55 hpf phenthiouric (PTU)-treated embryos were put in E3 egg water containing tricaine. The imaging of beating cilia in the nasal pit was conducted on a Zeiss AxioPlan microscope (Carl Zeiss, Jena, Germany) using a 63×/0.55 water immersion lens installed with a high-speed Photron FastCAM-PCI 500 videocamera (Photron LTD) [37]. Image acquisition was 250 frames per second by Photron FastCAM version 1.20.7 (Photron LTD). The movies are replayed in 30 frames per second.

**Protein blotting and in vitro translation**

1 μg of plasmids DNA (wild-type full-length zbb1 in pCS2+) were mixed with or without MOs and in vitro transcription/translation reactions were performed in the presence of biotinylated lysine according to manufacturer’s manual (TNT® Quick Coupled Transcription/Translation Systems, Transcend™ Non-Radioactive Translation Detection Systems, Promega). 1 μl of the reaction was diluted in 14 μl of 1× protein sample loading buffer. The reaction mixture was analyzed by SDS-PAGE and protein blotting using HRP-conjugated streptavidin.

**Statistical analyses**

Statistical analyses, Mann-Whitney rank sum test and Student’s t-test, were performed with SigmaStat® (Systat Software Inc.). Student’s t-test was used for the data of ciliary length, eye size and KV area measurement. In the case of the comparison of body gap angle, t-test could not be applied. Therefore, the differences in the variance between control- and morphant-embryo body gap angle were analyzed using the Mann-Whitney Rank Sum Test. The probability value of P<0.05 was considered statistically significant (***, P<0.001; **, P<0.01; *, P<0.05). X2 (chi-square) test was performed for the result of morphant rescue experiments. Degrees of freedom = 1, alpha level of significance = 0.05 (College of Saint Benedict & Saint John’s University) [http://www.physics.csbsju.edu/stats/contingency_NROW_NCOLUMN_form.html]

**Supporting Information**

**Figure S1** Alignment of the amino acid sequences of the 2 variants of maternally expressed znp7p7.2 transcripts. The alignment shows that 1 amino acid is substituted (S99R) and 18 amino acids are deleted (aa 101–118) in maternally expressed transcript variant 2 (Transcript 2) compared to transcript variant 1 (Transcript 1) of znp7p7.2. (TIF)

**Figure S2** Antisense morpholino oligonucleotides against zbb1. (A) MO (red line) targeting zbb1 translational start codon (AUG MO) and MO targeting exon 2 splice donor site (SP MO) of zbb1. (B) 1 μg of plasmids DNA containing wild-type full-length zbb1 in pCS2+ vector was mixed with or without MOs in vitro translation reactions. zbb1 AUG MO efficiently interfered with zbb1 protein expression. (TIF)

**Figure S3** Grading of dysmorphy. (A) Embryos at 48–55 hpf were assessed and scored as wild-type-like or with a degree of dysmorphy ranging from mild (D1) to severe (D3). (B) zbb1 or znp7p7.2 morphant embryos were always consistent of a mixed population of individuals with dorsally or ventrally curved body axis. (TIF)
Figure S4 Antisense morpholino oligonucleotides against znp7.2. (A) 2 independent MOs against exon 3 donor site (SP1 MO) and exon 6 acceptor site (SP2 MO) of znp7.2 were designed. (B and C) RT-PCR was performed with znp7.2 SP1 MO-injected embryos at 55 hpf. The following PCR with the primers (short blue lines) designed to produce 325 bp amplicon of wild-type znp7.2 cDNA was performed. znp7.2 SP1 MO efficiently interfered with normal splicing to cause the insertion of intron sequence between exon 3 and exon 4 resulting in 2 abnormally large amplicons. (Exon 3 is marked as green to visualize the mRNA splicing process.) (D) These abnormal splicing products have a stop codon in the intronic sequence leading to a truncation within the first ZF domain after translating 6 extra amino acids from intronic sequence. This eliminates all other ZF domains together with the C-terminus of zNphp7.2.

(TIF)

Figure S5 zBbs1- or zNphp7.2-deficient embryos showed defects in organ laterality. (A, B) In situ hybridisation of both zbbs1 and znp7.2 morphants at 24 hpf with lefty2 probe showed defective left-right asymmetry patterning. (C and D) Defective laterality of liver (foxA3) and pancreas (ins) was observed in zBbs1- and zNphp7.2-depleted embryos at 48 hpf.

(TIF)

Figure S6 Co-Injection of zbbs1 or znp7.2 mRNA partially rescued the corresponding morphant phenotype. Co-injection of 20 pg of zbbs1 mRNA (A and B) or 10 pg of znp7.2 mRNA (C and D) together with the corresponding MO partially rescued the dysmorphic changes caused by the MO-mediated knockdown, decreasing the dorsal body curvature and rescued pronephric cyst formation. (A: scale bar = 500 µm, B: scale bar = 100 µm) (D, $X^2 = 5.27$, $P = 0.022$; H, $X^2 = 14.5$, $P<0.001$).

(TIF)

Figure S7 Dextran injection revealed fluid excretion via the cloaca in zbbs1 and znp7.2 morphants. Zebrasfish control and morphant embryos with pronephric cysts at 96 hpf were injected with 5% FITC-conjugated dextran solution (70 kD) into the circulation. Fluorescent dye excretion with the urine at the cloaca (black and white arrows) was observed in control embryos (22/22), zbbs1 (16/16) and znp7.2 (21/23) morphants. The lower panel represents a zbbs1 morphant embryo with missing fluorescent dye excretion due to persistent closure of the cloaca (arrowheads). Images on the left column represent transmitted light images and images on the right column represent fluorescent images of the same embryo for each setting.

(TIF)

Figure S8 zbbs1 and znp7.2 morphants display reduced eye size and defective retinal layer formation. The eye size as area ($\mu m^2$) was measured for control embryos, zbbs1 and znp7.2 morphant embryos at 80 hpf. (A) Representative brightfield images showing reduced eye size for zbbs1 and znp7.2 morphants in comparison to the control (Scale bar = 200 µm). (B) Statistical quantification of the measurements proved that the reduction in eye size for zbbs1 and znp7.2 morphants was significant in comparison to the control. (C) Histological cross-sections of 96 hpf zbbs1 and znp7.2 morphants revealed defective layer formation in comparison to the control. Two representative images are shown for each setting. (Scale bar = 100 µm).

(TIF)

Figure S9 The morphants of zbbs1 and znp7.2 exhibited impaired motility of cilia in the pronephric tubule. (A) The areas where the movies were recorded are shown by the dashed box. (B) Acetylated tubulin staining demonstrated normal development of cilia in the nasal pit (Scale bar = 10 µm). 3-dimensional (3D) images (upper panel) and z-plane images (lower panel) show that the cilia formation in the morphants of zbbs1 and znp7.2 is normal compared to control embryo.

Movie S1 The anterior part of pronephric tubule in a control embryo. The pronephric cilia of control embryo show normal motility.

(AVI)

Movie S2 The posterior part of pronephric tubule in a control embryo. The pronephric cilia of control embryo show normal motility.

(AVI)

Movie S3 The cloaca area of a control embryo. The pronephric cilia of control embryo show normal motility.

(AVI)

Movie S4 The anterior part of pronephric tubule in a zbbs1 morphant embryo. The morphant embryo (zbbs1 0.5 mM) exhibited greatly reduced or abnormal motility. The arrows point to the individual cilia showing impaired movement. The movies were recorded from morphant embryos with clear pronephric cysts.

(AVI)

Movie S5 The middle part of pronephric tubule in a zbbs1 morphant embryo. Most of the cilia in a morphant embryo (zbbs1 0.5 mM) exhibited greatly reduced or abnormal motility. The movies were recorded from morphant embryos with clear pronephric cysts.

(AVI)

Movie S6 The cloaca area of a zbbs1 morphant embryo. Most of the cilia in a morphant embryo (zbbs1 0.5 mM) exhibited greatly reduced or abnormal motility. The movies were recorded from morphant embryos with clear pronephric cysts.

(AVI)

Movie S7 The anterior part of pronephric tubule in a znp7.2 morphant embryo. Most of the cilia in a morphant embryo (znp7.2 0.2 mM) exhibited greatly reduced or abnormal motility. The movies were recorded from morphant embryos with clear pronephric cysts.

(AVI)

Movie S8 Another anterior part of pronephric tubule in a znp7.2 morphant embryo. Most of the cilia in a morphant embryo (znp7.2 0.2 mM) exhibited greatly reduced or abnormal motility. The movies were recorded from morphant embryos with clear pronephric cysts.

(AVI)

Movie S9 The cloaca area of a znp7.2 morphant embryo. Most of the cilia in a morphant embryo (znp7.2 0.2 mM) exhibited greatly reduced or abnormal motility. The movies were recorded from morphant embryos with clear pronephric cysts.

(AVI)

Movie S10 The cilia in nasal pit of a control embryo. The cilia in nasal pit of control embryo showed normal uniformed beating along the whole epithelial layer. Arrows indicate the border of the nasal pit.

(AVI)
Movie S11  The cilia in nasal pit of a zbsl morphant embryo. The morphant embryo (zbsl 0.4 mM) exhibited remarkably impaired motility with preserved motility only in focal areas. Arrows indicate the border of the nasal pit. (AVI)

Movie S12  The cilia in nasal pit of a znphp7.2 morphant embryo. The morphant embryo (znphp7.2 0.2 mM) exhibited remarkably impaired motility with preserved motility only in focal areas. Arrows indicate the border of the nasal pit. (AVI)

Movie S13  The middle part of pronephric tubule in a control morphant embryo. The pronephric cilia of control morphant embryo (Cont MO 0.3 mM) show normal motility. (AVI)

Movie S14  The posterior part of pronephric tubule in a control morphant embryo. The pronephric cilia of control morphant embryo (Cont MO 0.3 mM) show normal motility. (AVI)

Movie S15  The cloaca area of a control morphant embryo. The pronephric cilia of control morphant embryo (Cont MO 0.3 mM) show normal motility. (AVI)

Movie S16  The middle part of pronephric tubule in a zbsl morphant embryo. The pronephric cilia of zbsl morphant embryo (zbsl 0.2 mM+Cont MO 0.1 mM) show active motility. The movies were recorded from the single knockdown morphant embryos without pronephric cysts. (AVI)

Movie S17  The cloaca area of a zbsl morphant embryo. The pronephric cilia of zbsl morphant embryo (zbsl 0.2 mM+Cont MO 0.1 mM) show active motility. The movies were recorded from the single knockdown morphant embryos without pronephric cysts. (AVI)

Movie S18  The middle part of pronephric tubule in a znphp7.2 morphant embryo. The pronephric cilia of znphp7.2 morphant embryo (znphp7.2 0.1 mM+Cont MO 0.2 mM) show active motility. The movies were recorded from the single knockdown morphant embryos without pronephric cysts. (AVI)

Movie S19  The posterior part of pronephric tubule in a znphp7.2 morphant embryo. The pronephric cilia of znphp7.2 morphant embryo (znphp7.2 0.1 mM+Cont MO 0.2 mM) show active motility. The movies were recorded from the single knockdown morphant embryos without pronephric cysts. (AVI)

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