

Dynein axonemal assembly factors (*dnaaf*) 5 and 9 are expressed in ciliated organs of zebrafish embryos

USHARANI NAYAK, KALYANI SAHOO, RAJEEB K. SWAIN*

*Institute of Life Sciences, NALCO Square, Chandrasekharpur, Bhubaneswar, India
and Regional Centre for Biotechnology, Faridabad, India*

ABSTRACT Dynein axonemal assembly factors (DNAAFs) play crucial roles in the formation and function of motile cilia, and their dysfunction often results in primary ciliary dyskinesia (PCD). We report the spatio-temporal expression patterns of *dnaaf5* and *dnaaf9* mRNA in zebrafish embryos, providing insight into their possible functions during development. We show that *dnaaf5* and *dnaaf9* mRNAs are expressed in motile ciliated tissues, such as the Kupffer's vesicle, pronephros, floor plate, brain and olfactory placode. The *dnaaf5* and *dnaaf9* crispants develop ciliopathic defects during zebrafish development. These data suggest that *dnaaf5* and *dnaaf9* may regulate motile cilia biogenesis and function in zebrafish. Our findings suggest functional redundancy and divergence among dynein arm assembly factors in vertebrates.

KEYWORDS: DNAAF, primary ciliary dyskinesia, ciliogenesis, CRISPR/Cas, zebrafish

Introduction

Primary ciliary dyskinesia (PCD) is a motile ciliopathy, clinically categorised with hydrocephalus, chronic respiratory infection, infertility, situs inversus totalis or heterotaxy and hearing loss. These rare and genetically heterogeneous disorders result from mutations in genes coding for proteins mediating motile ciliogenesis and function. Structurally motile cilia possess a basic feature as primary cilia (basal body, transition zone and axonemal microtubule) and several structural augmentations to generate motion. This additional element includes a central singlet microtubule (9+2 axoneme arrangement), radial spoke, nexin link connecting the outer nine microtubule doublet with central singlet and dynein arms (inner and outer dynein arm, IDA and ODA), that function as ATP-driven motor proteins responsible for microtubule sliding and bending motions (Raidt *et al.*, 2023). Several studies have identified that the mutations in genes associated with dynein arms assembly and trafficking cause PCD (Fassad *et al.*, 2023, Jat *et al.*, 2024). Axonemal dynein outer arm and inner arms are multimeric proteins that require a cytoplasmic assembly complex, which involves dynein axonemal assembly factors (DNAAF) and a transportation complex that includes the intraflagellar transport proteins.

DNAAF family proteins act as chaperone core or associated protein, facilitate assembly, stabilisation of dynein arms and sometimes their trafficking to the cilia axoneme. Like other ciliary proteins, DNAAFs are highly conserved across eukaryotes and several studies in multiple model systems, including zebrafish,

have elucidated their role in motile cilia formation and function (Mitchison *et al.*, 2012, Wan *et al.*, 2023). These studies supported the conserved function of DNAAFs and the usefulness of zebrafish for motile cilia studies. In addition, zebrafish develop different motile ciliated organs during embryogenesis, similar to higher vertebrates, such as the brain ventricle, ependymal cell lining of the central nervous system, pronephros, Kupffer's vesicle, otic vesicle and olfactory pit. The consensus nomenclature of DNAAF lists 19 proteins under the DNAAF category (Braschi *et al.*, 2022). Zebrafish has all orthologs except DNAAF8.

This study aims to characterise the spatiotemporal expression pattern and functional roles of *dnaaf5* and *dnaaf9* during early embryogenesis using zebrafish as a model organism. Zebrafish *dnaaf5* and *dnaaf9* show overlapping and distinct expression patterns during zebrafish embryogenesis. The expression of *dnaaf5* and *dnaaf9* in the motile ciliated organs during zebrafish embryogenesis is consistent with their role as dynein assembly factors.

Results

Zebrafish *Dnaaf5* and *Dnaaf9* show strong protein sequence conservation with human homologues

To assess the evolutionary relationship between zebrafish and human DNAAF5 and DNAAF9 homologues, their amino acid sequences were compared. DNAAF5 has a key functional domain called HEAT repeat (PF24573), which is conserved in zebrafish *Dnaaf5* (Fig. 1A). Human DNAAF9 has a shulin_C20orf194-like do-

*Address correspondence to: Rajeeb K. Swain. Institute of Life Sciences, NALCO Square, Chandrasekharpur, Bhubaneswar, India.
E-mail: rkswain@ils.res.in

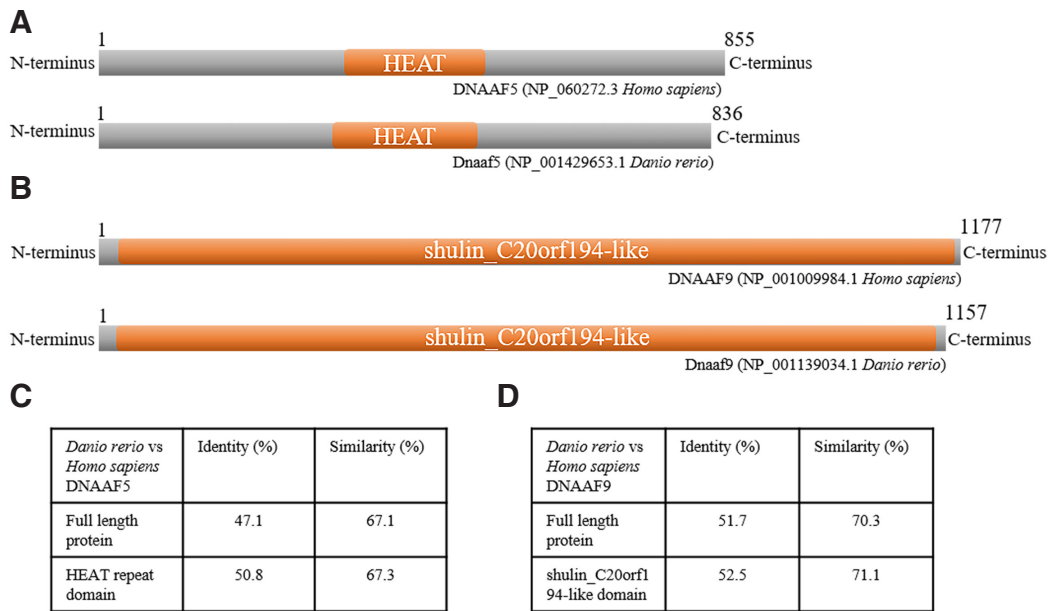


Fig. 1. Human and zebrafish DNAAF5 and DNAAF9 show conserved amino acid sequence and domain architecture. (A,B) Schematic representation of the protein marking the HEAT domain in human and zebrafish DNAAF5 and the shulin_C20orf194-like domain in human and zebrafish DNAAF9. The length of the schematic representation is proportional to the number of amino acids in the proteins. **(C,D)** Tables summarising the percentage sequence identity and similarity between human and zebrafish DNAAF5 and DNAAF9 full-length protein and active domain, respectively.

main (cd22936), which is also present in zebrafish Dnaaf9 (Fig. 1B).

We performed pairwise protein sequence alignments using EMBOSS Needle to measure the and found that zebrafish Dnaaf5 exhibited a high degree of conservation with human DNAAF5, with 47.1% identity and 67.1% amino acid similarity across the full-length protein (Fig. 1C). The human HEAT domain shares 50.8% identity and 67.3% similarity with the zebrafish HEAT domain (Fig. 1C). The zebrafish ortholog Dnaaf9 also showed substantial conservation, sharing 51.7% identity and 70.3% similarity with human DNAAF9, while the functional domains shulin_C20orf194-like are highly conserved, showing 52.5% identity and 71.1% similarity with the respective domain of human protein (Fig. 1D). These findings indicate that both zebrafish proteins retain essential structural features of human DNAAF5 and DNAAF9, supporting their functional relevance for modelling PCD-associated gene disruption.

***dnaaf5* and *dnaaf9* are expressed in motile cilia-bearing organs during zebrafish development**

To investigate the developmental expression dynamics of *dnaaf5* and *dnaaf9*, we carried out semi-quantitative RT-PCR and whole-mount *in situ* hybridisation (WISH) across multiple developmental stages of zebrafish. A 756 bp fragment of *dnaaf5* was used as a WISH probe, and the primers used to amplify this fragment were used in semiquantitative RT-PCR (Fig. 2A). RT-PCR results revealed that *dnaaf5* is expressed throughout zebrafish embryogenesis (Fig. 2B). WISH showed that *dnaaf5* is maternally deposited and ubiquitously expressed during early embryogenesis (1-cell to 6 hpf) (Fig. 2 C-E). Thereafter, organ-specific expression of *dnaaf5* can be seen (Fig. 2 F-N). It is detected in the dorsal forerunner cells (DFCs) at 9 hpf and, in a ciliated, vesicle-like organ, Kupffer's vesicle (KV) at 12 hpf (Fig. 2 F-G). Its expression can be seen in the eye, hind brain (HB) rhombomere, otic vesicle (OV), floorplate (FP), pronephros (P) and tail bud (TB) in 18 hpf embryos (Fig. 2 H-J). By 24 hpf, expression was evident in the telencephalon (TEL), diencephalon (DI), and midbrain (MB) (Fig. 2 I-J). At 48, 72 and 96 hpf, the *dnaaf5* is highly expressed in the brain compared to other organs (Fig. 2 K-N). It is also found to be expressed in the liver (L)

(Fig. 2 L,N). In the dorsal view of 72 hpf larvae, *dnaaf5* expression can be seen in the whole brain (B), olfactory pits (OP), and retina of the eye (R) (Fig. 2M). Sense riboprobe was used as a negative control for WISH (Fig. 2 O-P).

To examine the expression of *dnaaf9*, WISH and RT-PCR were performed using primers that span 868 bp of the transcript (Fig. 3A). Temporal expression of *dnaaf9*, examined by RT-PCR, shows low expression levels during developmental stages up to 48 hpf, followed by an increase thereafter (Fig. 3B). WISH revealed that the maternal deposit and early zygotic expression of *dnaaf9* is minimal (Fig. 3 C-D), compared to the early gastrulation stage (Fig. 3E). Its first organ-specific expression appears at dorsal forerunner cells (DFCs), then at KV (Fig. 3 F-G). 18 hpf onwards, *dnaaf9* is detected in the telencephalon (TEL), diencephalon (DI), hindbrain (HB), pronephros (P) and spinal cord (SC) (Fig. 3 H-J). After 24 hpf, expression extends to the midbrain (MB) (Fig. 3 I-N). At 48, 72 and 96 hpf, *dnaaf9* is strongly expressed in the brain (B) (Fig. 3 K-N). Expression of *dnaaf9* is also seen in the liver (Fig. 3N). Dorsal view of 72 hpf larvae marked *dnaaf9* expression in whole brain (B), olfactory pits (OP), and retina of eye (R) (Fig. 3M). Sense riboprobe was used as a negative control for WISH (Fig. 3 O-P). These results mark the dynamic spatiotemporal expression of *dnaaf5* and *dnaaf9* in different ciliated organs, during zebrafish development, suggesting roles in early embryogenesis and organ-specific functions.

***dnaaf5* crispants exhibit a primary ciliary dyskinesia (PCD)-like phenotype**

The zebrafish Dnaaf5 contains the conserved HEAT domain, spanning from 316th to 514th amino acid residue, corresponding to the coding sequences of exon 4 to exon 8 of the *dnaaf5* gene. To elucidate the role of *dnaaf5* in early development, we generated *dnaaf5* crispants using CRISPR/Cas9 with a guide RNA (gRNA) targeting the exon 5 of *dnaaf5* (Fig. 4A). Microinjection of this gRNA (12.5 pg or 25 pg per embryo) together with Cas9 mRNA into 1-cell embryos resulted in 80-90% embryos with genetic lesions (Fig. 4 B-C). The *dnaaf5* crispants exhibited body curvature, glomerular cyst and pericardial edema, at different stages. The

body axis curvature was observable at 2-day post fertilisation (dpf), while pericardial edema and glomerular cyst were prominent at 3 dpf. The penetrance of the phenotypes increased in a gRNA dose-dependent manner. At 12.5 pg/embryo gRNA injection, 22 out of 115 (19%) crispants showed combined glomerular cyst and body axis curvature phenotype, whereas 51 out of 130 (39%) in 25 pg/embryo gRNA injected *dnaaf5* crispants exhibited the same phenotypic combination (Fig. 4D). To confirm that the phenotypes result from a mutation in the *dnaaf5* gene, crispants with the phenotype were genotyped. Genotyping result confirmed the presence of mutations in the crispant showing developmental defects (Fig. 4E). Together, these results demonstrate that loss of *dnaaf5* might lead to PCD-like phenotypes in zebrafish.

dnaaf9 crispants exhibit scoliosis-like phenotype

The zebrafish *dnaaf9* comprises 37 exons. Exons 6 and 30 were targeted to create *dnaaf9* crispant with maximal disruption of the shulin_C20orf194-like domain (Fig. 5A). Both gRNAs exhibited above 70% efficiency in inducing mutations when injected at a 25 ng/ μ L concentration (Fig. 5 B-C). To generate *dnaaf9* crispant for phenotypic analysis, both gRNAs were co-injected into 1-cell embryos along with Cas9 mRNA. The crispants did not show morphological defects up to 3 dpf. The notochord deformities mimicking a scoliosis-like phenotype were seen in 18 out of 72 (20%) crispants by 4 to 6 dpf (Fig. 5D). To confirm that the observed phenotype resulted from the mutation of the *dnaaf9* gene, crispants exhibiting the phenotype were genotyped. The PCR was performed

using a forward primer from exon 6 and a reverse primer from exon 30. The expected amplicon size for wild-type DNA using these primers was 24.7 kb. In contrast, these primers will amplify a 156 bp fragment in the crispants where double-strand breaks are induced in exon 6 and exon 30, thus removing the DNA sequence between these two exons and joining them together. The PCR conditions were designed to selectively amplify the shorter fragment, while the DNA above 500 bp were not amplified. The anticipated amplicon was detected in crispants showing phenotype, while wild-type DNA failed to amplify, confirming the large deletion (Fig. 5E). To confirm the mutation, the amplified products were cloned and subjected to Sanger sequencing. Sanger sequencing confirmed deletion of a 24 kb genomic region with a small number of nucleotide insertions at the junction (Fig. 5F). This deletion corresponds to 2166 bp of the protein coding sequence, encod-

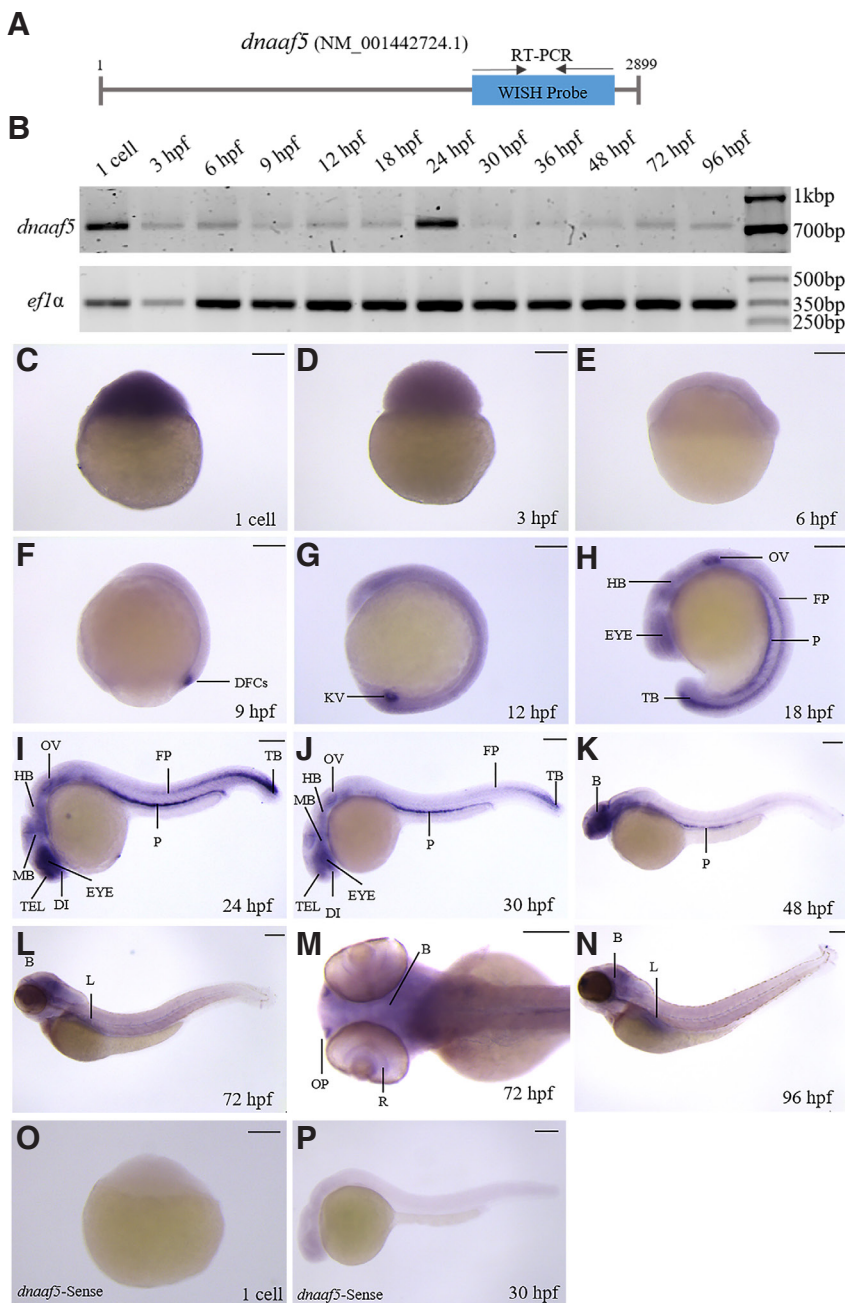


Fig. 2. Spatiotemporal expression of *dnaaf5* during zebrafish embryogenesis. (A) Schematic representation of *dnaaf5* mRNA showing the position of primers used for whole-mount *in situ* hybridisation (WISH) probe synthesis and RT-PCR analysis. (B) Temporal expression of *dnaaf5* mRNA during zebrafish development. Zebrafish *ef1a* is used as a loading control. (C-N) Whole-mount *in situ* hybridisation showing *dnaaf5* transcript localisation from the 1-cell to 96 hours post-fertilisation (hpf). (C-E) During early development (1-cell to 6 hpf), *dnaaf5* is ubiquitously expressed, with strong maternal deposition of the transcript and early zygotic expression. (F-N) Later, it is expressed in specific organs. (F,G) *dnaaf5* is expressed in dorsal forerunner cells (DFC) and Kupffer's vesicle (KV) at 9 hpf and 12 hpf respectively. (H-J) 18 hpf onwards it is expressed in eye, hindbrain (HB), pronephros (P) and floorplate (FP), tail bud (TB) and otic vesicle (OV). (I-N) *dnaaf5* is expressed in the telencephalon (TEL), diencephalon (DI) and midbrain (MB) after 24 hpf. (K-N) Expression is more intense in the brain than in other organs by 48 hpf. (L, N) After 72 hpf, it is expressed in the liver (L). (M) Dorsal view of 72 hpf embryo head, clarifying *dnaaf5* expression throughout the brain (B), retina (R) of the eye and olfactory pit (OP). (O,P) WISH with the sense *dnaaf5* probe in 1 cell and 30 hpf zebrafish embryos shows no detectable signal. Scale bars, 150 μ m.

ing 722 amino acids (residue 185 to 907), which encompasses the majority of the shulin_C20orf194-like domain, thereby confirming a functional mutation. Collectively, these results demonstrate that *dnaaf9* crispants exhibit scoliosis-like phenotype in zebrafish.

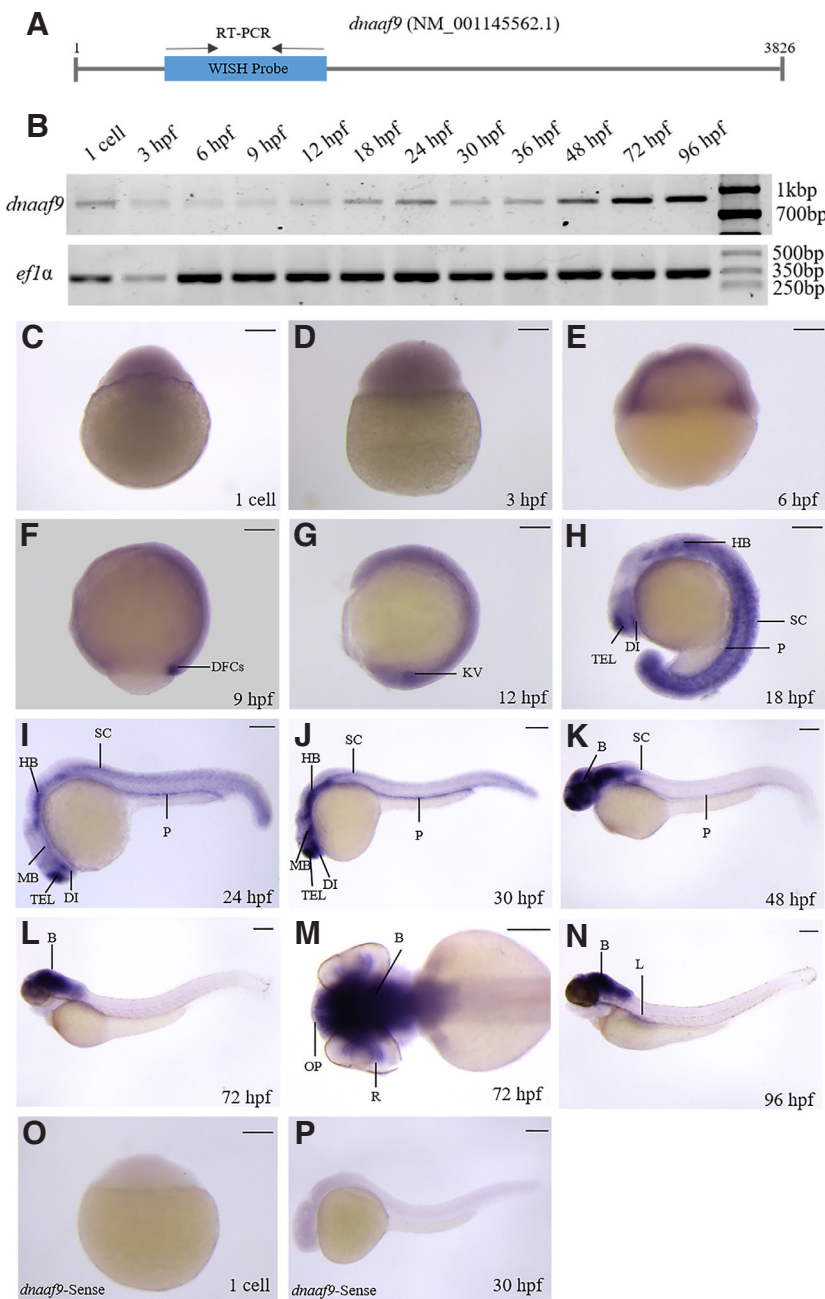
Discussion

Here, we report the spatio-temporal expression pattern of *dnaaf5* and *dnaaf9* during zebrafish development (Figs. 2 and 3). The expression patterns of *dnaaf5* and *dnaaf9* in motile ciliated organs, including the KV, brain, floor plate, spinal cord, olfactory pit, and pronephros, in zebrafish embryos are consistent with the known functions of the dynein assembly factors in motile cilia. Many *dnaafs* exhibit a broad expression domain, while some are

restricted to motile ciliated organs. Several dynein assembly factors showing maternal expression are reported (Gao et al., 2010; Panizzi et al., 2012; Falkenberg et al., 2021). *wdr69* (*dnaaf18*) is a cilia-associated gene, expressed only in motile ciliated organs like Kupffer's vesicle, floor plate, pronephros and otic vesicle, and is functionally involved in ODA assembly. *wdr69* deficient zebrafish embryos developed pronephric cyst, body curvature, pericardial edema, otolith defect, and laterality defect in different organs (Gao et al., 2010). Whereas *dnaaf19* (*ccdc103*), along with ciliary expression, is expressed in hemato-vascular progenitors, and found to be functionally involved in myeloid cell proliferation, which is independent of cilia. Classical dynein assembly function is also exhibited by *dnaaf19* (Panizzi et al., 2012; Falkenberg et al., 2021). In zebrafish, four PIH family protein genes *pih1d1* (*dnaaf14*), *pih1d2* (*dnaaf15*), *ktu* (*dnaaf2*), and *twister* (*dnaaf6*) are expressed in Kupffer's vesicle, floor plate, pronephros and otic vesicle, along with some parts of brains. Their functional characterisation demonstrated their importance in sperm motility and Kupffer's vesicle cilia motility, and also uncovered their role in dynein protein assembly (Yamaguchi et al., 2018). Disruption of the dyslexia candidate gene *dyx1c1* (*dnaaf4*), expressed in KV, olfactory placode, brain, otic vesicle and pronephros, resulted in loss of cilia dynein arm and pronephric brush border defect (Chandrasekar et al., 2013). The maternal deposition of *dnaaf5* and *dnaaf9* expression during gastrula stages suggests that these genes may have roles during early embryogenesis.

CRISPR/Cas9-mediated F0 crispants have become valuable for preliminary characterisation of gene functions in zebrafish (Wu et al., 2018; Kroll et al., 2021). Crispants have been found to faithfully recapitulate the phenotypes seen in stable knock-out lines (Lin et al., 2025). Based on gene architecture and protein functional domain organisation, distinct CRISPR knockout strategies were employed in this study. Zebrafish *dnaaf5* has a conserved HEAT domain encoded by exons 4-8.

Fig. 3. *dnaaf9* exhibits a broad expression domain across the early developmental stages of zebrafish. (A) Schematic representation of *dnaaf9* mRNA showing the position of primers used for WISH probe synthesis and RT-PCR analysis. (B) Temporal expression analysis of *dnaaf9* mRNA during zebrafish development, using *ef1a* as a loading control. (C–N) Whole-mount *in situ* hybridisation marking *dnaaf9* transcript localisation from the 1-cell to 96 hpf. (C–E) *dnaaf9* is ubiquitously expressed during early development (1-cell to 6 hpf), with low maternal deposition and early zygotic expression. (F,G) Expression becomes restricted to dorsal forerunner cells (DFC) and Kupffer's vesicle (KV) at 9 and 12 hpf, respectively. (H–J) From 18 hpf onward, *dnaaf9* is detected in the telencephalon (TEL), diencephalon (DI), hindbrain (HB), pronephros (P) and spinal cord (SC). (I–N) At 30 hpf and later, expression extends to the midbrain (MB). (K–N) After 48 hpf, expression intensifies in the brain relative to other tissues. (N) By 96 hpf, expression is also detected in the liver (L). (M) Dorsal view of a 72 hpf embryo head showing *dnaaf9* expression in the brain (B), retina (R), and olfactory pit (OP). (O,P) WISH using the *dnaaf9* sense probe in 1 cell and 30 hpf zebrafish embryos shows no detectable signal. Scale bars, 150 μ m.



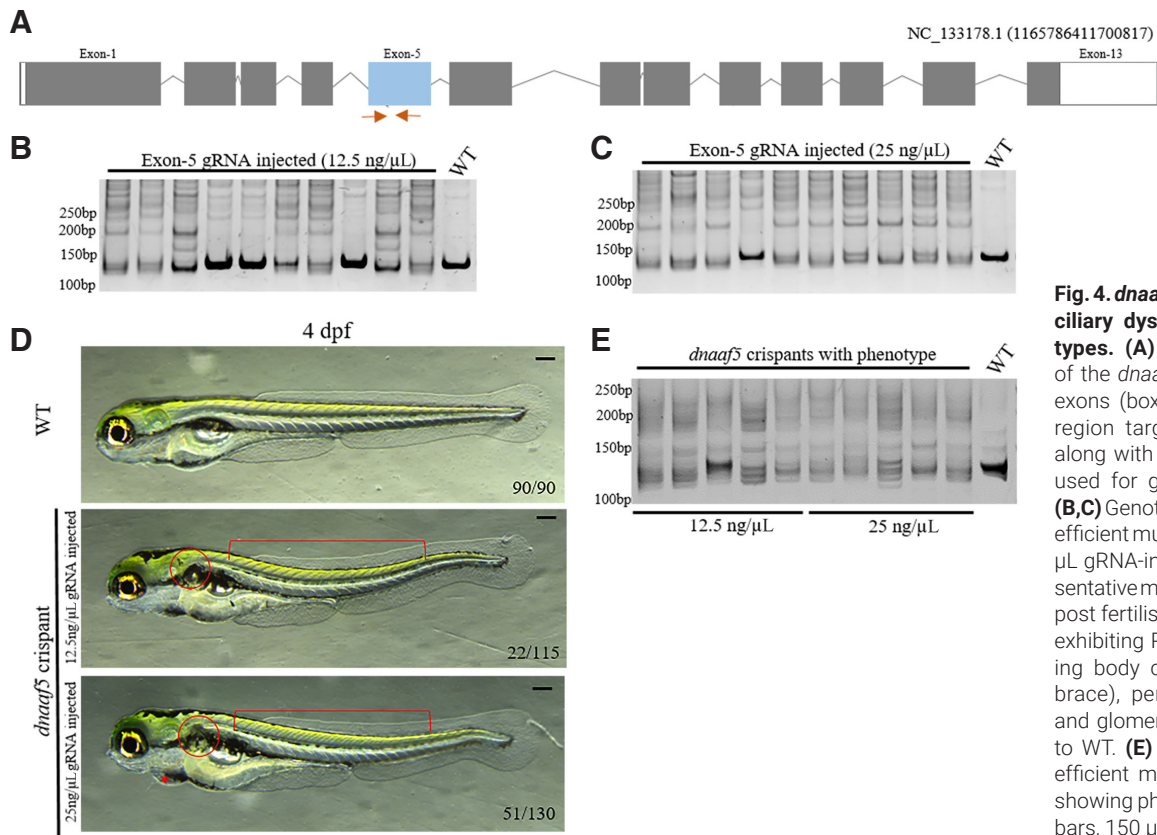


Fig. 4. *dnaaf5* crisprants exhibit primary ciliary dyskinesia (PCD)-like phenotypes. (A) Schematic representation of the *dnaaf5* gene structure, showing exons (boxes), introns (lines) and the region targeted by gRNA (blue box) along with the position of the primers used for genotyping (Orange arrow). (B,C) Genotyping results demonstrating efficient mutagenesis in 12.5 and 25 ng/ μ L gRNA-injected crisprants. (D) Representative morphological images of 4-day post fertilisation (dpf) *dnaaf5* crisprants exhibiting PCD-like phenotypes, including body curvature (square grouping brace), pericardial edema (asterisk), and glomerular cyst (circle) compared to WT. (E) Genotyping result showing efficient mutation in *dnaaf5* crisprants showing phenotype combination. Scale bars, 150 μ m.

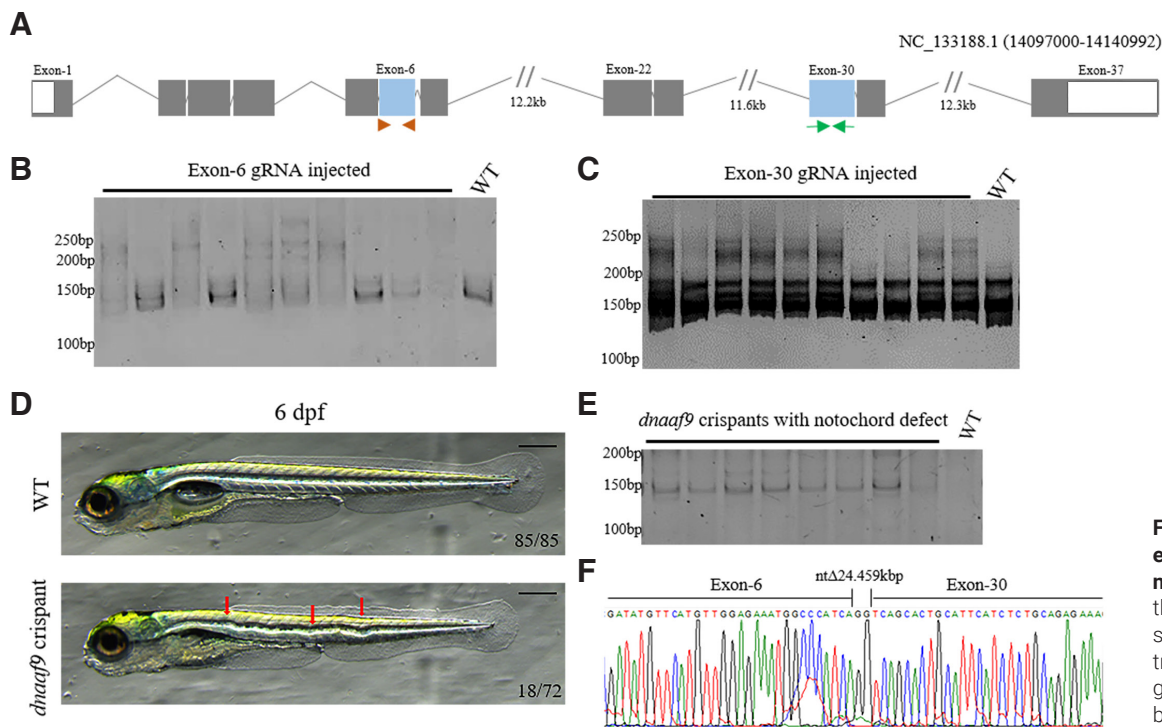


Fig. 5. *dnaaf9* crisprants exhibit scoliotic-like phenotype. (A) Illustration of the *dnaaf9* gene structure, showing exons (boxes), introns (lines) and exons targeted by guide RNAs (blue boxes) along with the primer positions used for genotyping (Orange arrowhead-Exon 6, Green arrow-Exon 30). (B,C) Heteroduplex assay result confirming high efficiency of guide RNA-mediated mutagenesis in F0 crisprants. (D) Representative morphological images of 6 dpf *dnaaf9* crisprants exhibiting notochord malformation (red arrows). (E) Genotyping result showing efficient mutation in *dnaaf9* crisprants exhibiting notochord defect. (F) Representative sequencing results showing the position of the deletion of 24.7 kb between exons 6 and 30. Scale bars, 150 μ m.

(Orange arrowhead-Exon 6, Green arrow-Exon 30). (B,C) Heteroduplex assay result confirming high efficiency of guide RNA-mediated mutagenesis in F0 crisprants. (D) Representative morphological images of 6 dpf *dnaaf9* crisprants exhibiting notochord malformation (red arrows). (E) Genotyping result showing efficient mutation in *dnaaf9* crisprants exhibiting notochord defect. (F) Representative sequencing results showing the position of the deletion of 24.7 kb between exons 6 and 30. Scale bars, 150 μ m.

Hence, the exon 5 of this gene was targeted to disrupt the HEAT domain, thus creating a zebrafish crispant with a potentially non-functional Dnaaf5. In contrast, *dnaaf9* with 37 protein-coding exons required a large fragment deletion using two gRNA targeting exon 6 and 30 to ensure major loss to the protein (Wu et al., 2018; Kim and Zhang, 2020). We observed PCD-like phenotypes in *dnaaf5* crispants including pericardial edema, glomerular cyst and body curvature (Fig. 4D). These phenotypes align with previously reported phenotypes associated with mutations in human *DNAAF5* (Horani et al., 2023). The *dnaaf9* crispants showed scoliosis-like phenotype during late embryogenesis, with a wavy notochord (Fig. 5D). Defects in motile cilia leading to scoliosis are reported both in zebrafish and higher vertebrates (Wang et al., 2020, Marie-Hardy et al., 2021), but their underlying mechanisms are not well understood. The functional data reported here are from the F0 crispants that do not represent a homozygous mutant line. Despite this lacuna, the preliminary data presented here suggest that the zebrafish *dnaaf5* and *dnaaf9* knockout lines may serve as a valuable resource for investigating how defects in dynein assembly factors contribute to PCD and scoliosis.

Materials and Methods

Zebrafish maintenance

Zebrafish were maintained in a circulating system at 28.5°C. Embryos were raised in E3 medium at 28.5°C and staged according to Kimmel et al., (Kimmel et al., 1995). The Zebrafish Tübingen (Tü) strain was used in all experiments. All the experiments were conducted as per the approval of the Institutional Animal Ethics Committee (ILS/IAEC-250-AH/FEB-22).

Bioinformatic analysis

Zebrafish orthologue for all human DNAAFs retrieved from the ZFIN and the NCBI database. The NCBI Conserved Domain Database and NCBI ensemble were used to identify the domains present in zebrafish Dnaaf9 (NP_001139034.1), human DNAAF5 (NP_060272.3) and DNAAF9 (NP_001009984.1). The functional domain for zebrafish Dnaaf5 (NP_001429653.1) was predicted using InterProScan software. Protein sequence alignments and protein sequence identity calculation were done using EMBOSS Needle.

Gene cloning and WISH probe synthesis

Total RNA was isolated from 24 hpf zebrafish embryos using the Direct-zol RNA Miniprep kit (Zymo Research, USA). cDNA was synthesised from isolated RNA using SuperScript IV First-

Strand Synthesis System (Thermo Fisher Scientific). The *dnaaf5* (NM_001442724.1) and *dnaaf9* (NM_001145562.1) genes were amplified by Phusion polymerase (Thermo Fisher Scientific), using gene-specific primers (Table 1), using cDNA as template. The PCR products were cloned into the PCR-Blunt-II-TOPO vector (Thermo Fisher Scientific) and were verified by Sanger sequencing. The plasmids containing the above genes were linearised with XhoI and BamHI, for antisense and sense probe synthesis, respectively. Digoxigenin-labelled antisense and sense riboprobe were synthesised by SP6 and T7 RNA polymerase.

Whole mount *in situ* hybridisation (WISH)

Embryos were treated with PTU to stop the pigmentation. For whole mount *in situ* hybridisation, embryos at different developmental stages were fixed in 4% paraformaldehyde (PFA), dehydrated through a graded methanol:PBST series, and stored in 100% methanol at -20 °C until use. The fixed and dehydrated embryos were rehydrated in PBST, permeabilised with proteinase-K and re-fixed in 4% PFA. Then the embryos were incubated in hybridisation buffer for 2 hours (without the probe), then the probe was added and kept at 65°C overnight. The next day, the embryos were washed with formamide/SSC buffer at 65°C to remove unbound probes and transferred to MABT (0.1% Tween-20 in 1X MAB) at RT. Then the embryos were transferred to blocking buffer (10% fetal bovine serum (FBS) and 2% blocking reagent (Roche 1109617600) in MABT, 2 hours at RT). Then it was incubated overnight with alkaline phosphatase (AP) conjugated anti-digoxigenin (DIG) at 4°C. On the third day, gene transcripts were detected by developing with BM purple (Roche 11442074001). The reaction was stopped by adding PBS and fixed with 4% PFA kept at 4°C for overnight.

RT-PCR

cDNA was prepared by taking an equal amount of total RNA extracted from different developmental stages. The above-mentioned WISH primer pairs were used to amplify 756 bp and 868 bp of *dnaaf5* and *dnaaf9*, respectively (Table 1). Zebrafish ef1 alpha (now called eukaryotic translation elongation factor 1 alpha 1, like 1 (eef1a1l1), NM_131263.1) was used as a loading control.

gRNA design and synthesis

gRNA target for *dnaaf5* and *dnaaf9* was designed using CRISPR RGEN tools (<http://www.rgenome.net/cas-designer/>). gRNAs were designed against exon 5 (5'-GATTCCTTACCACCAGCTCC-3') of *dnaaf5*. For *dnaaf9* crispants, gRNAs were designed to target

TABLE 1

Gene name	Forward primer	Reverse primer	Purpose	Product length
<i>dnaaf5</i>	CTTCCAGCAGAGAGTGGT	CATCTCAAATCTGGGAGACAAC	WISH/RT-PCR	756
<i>dnaaf9</i>	ATTGTCTGACTGAAACAGAGTACGA	CCCTTACTGAATCAACCAAAGC	WISH/RT-PCR	868
<i>dnaaf5</i>	TAAACCCAGTGCTTTTT	CAACCAGTCCCAATATC	Genotyping (<i>dnaaf5</i> exon 5)	132
<i>dnaaf5</i>	GTGTTTCTCTCTGTAGGCA	AGAGCAGCGTTCTGACA	Genotyping (<i>dnaaf5</i> exon 10)	119
<i>dnaaf9</i>	TTGATATTTAATAGGAC	CTTATATTCATGGTGAA	Genotyping (<i>dnaaf5</i> exon 6)	122
<i>dnaaf9</i>	GGTTTGACTATGGAGCAA	GTATGTATGTATGTAAGAAG	Genotyping (<i>dnaaf5</i> exon 30)	148
<i>ef1a</i>	CTTCTCAGGCTGACTGTGC	CCGCTAGCATTACCCTCC	RT-PCR	358

exon 6 and exon 30, having sequence 5'-GGAGAAATGGCCCAT-CATTC-3' and 5'-GATGAATGCAGTGCTGAGAT-3' respectively. DNA templates for the designed gRNA target and the universal oligo were obtained from IDT and annealed to prepare the template for gRNA transcription. gRNA synthesised from annealed template using MEGAscript SP6 or T7 kit (Thermo Fisher Scientific) (Gagnon *et al.*, 2014).

Microinjection

Crispantns were generated by injecting 1nl of gRNA and Cas9 mRNA mixture, containing each gRNA (for one gene) at a concentration of 12.5 ng/μL or 25 ng/μL and Cas9 mRNA at a concentration of 100 ng/μL. Microinjection was performed at the single cell stage of zebrafish embryos using a Femtojet microinjector (Eppendorf).

High resolution melt curve analysis (HRM) and Heteroduplex assay (HD)

Genomic DNA was isolated from the whole embryos 48 hours post gRNA/Cas9 injection in PCR tubes by incubating in TE buffer at 95°C and digested by proteinase K at 55°C (Zhu *et al.*, 2014). For DNA amplification, primers were designed flanking the CRISPR target site (Table 1). The reaction for HRM was prepared by adding 2X SYBR® Green JumpStart™ Taq ReadyMix, genomic DNA, forward and reverse primers (10 μM) and nuclease-free water. PCR reaction method (95°C/15 sec., 50-60°C /20sec and 72°C /20 sec) was set for 40 cycles. During melt curve reaction temperature was increased from 60-95°C with an increment of 0.2°C and fluorescence was recorded at every 0.2 sec.

In the HD assay, the amplified DNA obtained from HRM was denatured at 95°C and reannealed at RT. The products were then loaded on 12% polyacrylamide gel and electrophoresis was carried out at 120V for 90 minutes, to analyse the heteroduplex and homoduplex bands formed in the crispant compared to the WT homoduplex.

Microscopy

The live zebrafish embryos were anaesthetised in Tricaine methanesulfonate solution, and then dropped on 3% methylcellulose bed. Thereafter, the bright-field images were obtained, while fixed samples were mounted in 75% glycerol and images were taken. All the imaging was taken using a Leica M205FA stereo microscope.

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Conflicts of interest

The authors declare no conflict of interest.

Author contributions

RKS conceived, supervised and provided resources for the study. UN and KS carried out the experiments. UN wrote the manuscript. RKS and KS revised the manuscript. All the authors approved the final version of the manuscript.

References

- BRASCHI B., OMRAN H., WITMAN G. B., PAZOUR G. J., PFISTER K. K., BRUFORD E. A., KING S. M. (2022). Consensus nomenclature for dyneins and associated assembly factors. *Journal of Cell Biology* 221: e202109014. <https://doi.org/10.1083/jcb.202109014>
- CHANDRASEKAR G., VESTERLUND L., HULTENBY K., TAPIA-PÁEZ I., KERE J. (2013). The Zebrafish Orthologue of the Dyslexia Candidate Gene DYX1C1 Is Essential for Cilia Growth and Function. *PLoS ONE* 8: e63123. <https://doi.org/10.1371/journal.pone.0063123>
- FALKENBERG L. G., BECKMAN S. A., RAVISANKAR P., DOHN T. E., WAXMAN J. S. (2021). Ccdc103 promotes myeloid cell proliferation and migration independent of motile cilia. *Disease Models & Mechanisms* 14: dmm048439. <https://doi.org/10.1242/dmm.048439>
- FASSAD M. R., RUMMAN N., JUNGER K., PATEL M. P., THOMPSON J., GOGGIN P., UEFFING M., BEYER T., BOLDT K., LUCAS J. S., MITCHISON H. M. (2023). Defective airway intraflagellar transport underlies a combined motile and primary ciliopathy syndrome caused by IFT74 mutations. *Human Molecular Genetics* 32: 3090-3104. <https://doi.org/10.1093/hmg/ddad132>
- GAGNON J. A., VALEN E., THYME S. B., HUANG P., AHKMETOVA L., PAULI A., MONTAGUE T. G., ZIMMERMAN S., RICHTER C., SCHIER A. F. (2014). Efficient Mutagenesis by Cas9 Protein-Mediated Oligonucleotide Insertion and Large-Scale Assessment of Single-Guide RNAs. *PLoS ONE* 9: e98186. <https://doi.org/10.1371/journal.pone.0098186>
- GAO C., WANG G., AMACK J. D., MITCHELL D. R. (2010). Oda16/Wdr69 is essential for axonemal dynein assembly and ciliary motility during zebrafish embryogenesis. *Developmental Dynamics* 239: 2190-2197. <https://doi.org/10.1002/dvdy.22355>
- HORANI A., GUPTA D. K., XU J., XU H., CARMEN PUGA-MOLINA L., SANTI C. M., RAMAGIRI S., BRENNAN S. K., PAN J., KOENITZER J. R., HUANG T., HYLANDER M., *et al.* (2023). The effect of Dnaaf5 gene dosage on primary ciliary dyskinesia phenotypes. *JCI Insight* 8: e168836. <https://doi.org/10.1172/jci.insight.168836>
- JAT K. R., FARUQ M., JINDAL S., BARI S., SONI A., SHARMA P., MATHEWS S., SHAMIM U., AHUJA V., UPPILLI B., YADAV S. C., LODHA R., *et al.* (2024). Genetics of 67 patients of suspected primary ciliary dyskinesia from India. *Clinical Genetics* 106: 650-658. <https://doi.org/10.1111/cge.14590>
- KIM B. H., ZHANG G. J. (2020). Generating Stable Knockout Zebrafish Lines by Deleting Large Chromosomal Fragments Using Multiple gRNAs. *G3 Genes/Genomes/Genetics* 10: 1029-1037. <https://doi.org/10.1534/g3.119.401035>
- KIMMEL C. B., BALLARD W. W., KIMMEL S. R., ULLMANN B., SCHILLING T. F. (1995). Stages of embryonic development of the zebrafish. *Developmental Dynamics* 203: 253-310. <https://doi.org/10.1002/aja.1002030302>
- KROLL F., POWELL G. T., GHOSH M., GESTRI G., ANTINUCCI P., HEARN T. J., TUNBAKH., LIM S., DENNIS H. W., FERNANDEZ J. M., WHITMORE D., DREOSTI E., *et al.* (2021). A simple and effective F0 knockout method for rapid screening of behaviour and other complex phenotypes. *eLife* 10: e59683. <https://doi.org/10.7554/eLife.59683>
- LIN S. J., HUANG K., PETREE C., QIN W., VARSHNEY P., VARSHNEY G. K. (2025). Optimizing gRNA selection for high-penetrance F0 CRISPR screening for interlocking disease gene function. *Nucleic Acids Research* 53: gkaf180. <https://doi.org/10.1093/nar/gkaf180>
- MARIE-HARDY L., CANTAUT-BELARIF Y., PIETTON R., SLIMANI L., PASCAL-MOUSSELLARD H. (2021). The orthopedic characterization of cfap298tm304 mutants validate zebrafish to faithfully model human AIS. *Scientific Reports* 11: 7392. <https://doi.org/10.1038/s41598-021-86856-1>
- MITCHISON H. M., SCHMIDTS M., LOGES N. T., FRESHOUR J., DRITSOULA A., HIRST R. A., O'CALLAGHAN C., BLAU H., AL DABBAGH M., OLBRICH H., BEALES P. L., YAGI T., *et al.* (2012). Mutations in axonemal dynein assembly factor DNAAF3 cause primary ciliary dyskinesia. *Nature Genetics* 44: 381-389. <https://doi.org/10.1038/ng.1106>
- PANIZZI J. R., BECKER-HECK A., CASTLEMAN V. H., AL-MUTAIRI D. A., LIUY., LOGES N. T., PATHAK N., AUSTIN-TSE C., SHERIDAN E., SCHMIDTS M., OLBRICH H., WERNER C., *et al.* (2012). CCDC103 mutations cause primary ciliary dyskinesia by disrupting assembly of ciliary dynein arms. *Nature Genetics* 44: 714-719. <https://doi.org/10.1038/ng.2277>
- RAIDT J., LOGES N. T., OLBRICH H., WALLMEIER J., PENNEKAMP P., OMRAN H. (2023). Primary ciliary dyskinesia. *La Presse Médicale* 52: 104171. <https://doi.org/10.1016/j.lpm.2023.104171>

- WAN F., YU L., QU X., XIA Y., FENG K., ZHANG L., ZHANG N., ZHAO G., ZHANG C., GUO H. (2023). A novel mutation in PCD-associated gene DNAAF3 causes male infertility due to asthenozoospermia. *Journal of Cellular and Molecular Medicine* 27: 3107-3116. <https://doi.org/10.1111/jcmm.17881>
- WANG Y., LIU Z., YANG G., GAO Q., XIAO L., LI J., GUO C., TROUTWINE B. R., GRAY R. S., XIE L., ZHANG H. (2020). Coding Variants Coupled With Rapid Modeling in Zebrafish Implicate Dynein Genes, dnaaf1 and zmynd10, as Adolescent Idiopathic Scoliosis Candidate Genes. *Frontiers in Cell and Developmental Biology* 8: 582255. <https://doi.org/10.3389/fcell.2020.582255>
- WU R. S., LAM I. I., CLAY H., DUONG D. N., DEOR C., COUGHLIN S. R. (2018). A Rapid Method for Directed Gene Knockout for Screening in G0 Zebrafish. *Developmental Cell* 46: 112-125.e4. <https://doi.org/10.1016/j.devcel.2018.06.003>
- YAMAGUCHI H., ODA T., KIKKAWA M., TAKEDA H. (2018). Systematic studies of all PIH proteins in zebrafish reveal their distinct roles in axonemal dynein assembly. *eLife* 7: e36979. <https://doi.org/10.7554/eLife.36979>
- ZHU X., XU Y., YU S., LU L., DING M., CHENG J., SONG G., GAO X., YAO L., FAN D., MENG S., ZHANG X., et al. (2014). An Efficient Genotyping Method for Genome-modified Animals and Human Cells Generated with CRISPR/Cas9 System. *Scientific Reports* 4: 6420. <https://doi.org/10.1038/srep06420>