

Deficiency in DNAH12 causes male infertility by impairing DNAH1 and DNALI1 recruitment in humans and mice

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eLife Assessment

This **fundamental** study further validates DNAH12 as a causative gene for asthenoteratozoospermia and male infertility in both humans and mice. **Compelling** evidence supports the notion that DNAH12 is essential for proper axonemal development. This work will be of interest to reproductive biologists studying spermatogenesis and sperm biology, as well as andrologists focusing on male fertility.

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Abstract

Asthenoteratozoospermia, a prevalent cause of male infertility, lacks a well-defined etiology. DNAH12 is a special dynein featured by the absence of a microtubule-binding domain, however, its functions in spermatogenesis remain largely unknown. Through comprehensive genetic analyses involving whole-exome sequencing and subsequent Sanger sequencing on infertile patients and fertile controls from six distinct families, we unveiled six biallelic mutations in *DNAH12* that co-segregate recessively with male infertility in the studied families. Transmission electron microscopy (TEM) revealed pronounced axonemal abnormalities, including inner dynein arms (IDAs) impairment and central pair (CP) loss in sperm flagella of the patients. Mouse models (*Dnah12*^{-/-} and *Dnah12*^{mut/mut}) were generated and recapitulated the reproductive defects in the patients. Noteworthy, DNAH12 deficiency did not show effects on cilium organization and function. Mechanistically, DNAH12 was confirmed to interact with two other IDA components DNALI1 and DNAH1, while disruption of DNAH12 leads to failed recruitment of DNALI1 and DNAH1 to IDAs and compromised sperm development. Furthermore, DNAH12 also interacts with radial spoke head proteins

RSPH1, RSPH9, and DNAJB13 to regulate CP stability. Moreover, the infertility of *Dnah12*^{-/-} mice could be overcome by intracytoplasmic sperm injection (ICSI) treatment. Collectively, DNAH12 plays a crucial role in the proper organization of axoneme in sperm flagella, but not cilia, by recruiting DNAH1 and DNALI1 in both humans and mice. These findings expand our comprehension of dynein component assembly in flagella and cilia and provide a valuable marker for genetic counseling and diagnosis of asthenoteratozoospermia in clinical practice.

Introduction

Infertility has emerged as a global health concern, impacting around 8-12% of couples during their reproductive years, approximately 50% of infertility cases are attributed to male factors (1). Among these, asthenoteratozoospermia, characterized by poor sperm motility and notable morphological abnormalities, stands out as a prevalent contributor to male infertility (2, 3). Recent investigations have unveiled a multitude of gene mutations associated with asthenoteratozoospermia and male infertility, including Dynein Axonemal Heavy Chain (DNAH) inner dynein arm (IDA) genes like *DNAH1*, *DNAH2*, *DNAH6*, *DNAH7*, and *DNAH10*, as well as outer dynein arm (ODA) genes such as *DNAH8* and *DNAH17*, all implicated in primary male infertility with asthenoteratozoospermia (4–11). Additionally, genes encoding cilia and flagella-associated proteins (CFAP), including *CFAP43*, *CFAP44*, *CFAP57*, and *CFAP61*, have been identified as linked to asthenoteratozoospermia (12–15). Despite these advances, the intricate and heterogeneous features of asthenoteratozoospermia leave a significant number of cases unexplained. It is anticipated that additional genetic factors play a role in sperm anomalies and dysfunction.

The axoneme, a fundamental structure of cilia and flagella, exhibits a well-organized pattern with nine peripheral doublet microtubules (DMTs) and a central pair of microtubules (CP), constituting the classical “9+2” microtubular arrangement (16). In motile cilia or flagella, the radial spoke (RS) head contacts the CP accessory structures, transmitting mechanical and chemical signals from the central microtubules to the axonemal dynein arms, thus coordinating the flagellar or ciliary movement (17). Within mammalian species, the DNAH family plays a crucial role in the axonemal structure, encompassing eight IDA members (*DNAH1-3*, *DNAH6*, *DNAH7*, *DNAH10*, *DNAH12*, *DNAH14*) and five ODA members (*DNAH5*, *DNAH8*, *DNAH9*, *DNAH11*, *DNAH17*). Despite the collaborative effort of IDAs and ODAs in generating the force necessary for flagellar and ciliary motion, they serve distinct functions. Chlamydomonas ODAs, composed of α , β , and γ chains, primarily regulate beat frequency, while IDAs, with seven major subspecies (a-g), contribute to bending formation and beating form (18). Mutations in genes encoding ODA or IDA components have been linked to asthenoteratozoospermia, with or without symptoms of primary ciliary dyskinesia (PCD), a genetic disorder characterized by chronic airway infections or situs inversus (19). Notably, deficiency in certain IDAs and ODAs coding genes, such as *DNAH7* or *DNAH11*, may lead to PCD (20–22). Patients carrying mutations in *DNAH1* and *DNAH2* typically lack PCD symptoms (4, 23), although a study suggested the potential contribution of the *DNAH1* p.Lys1154Gln mutation to PCD (24). The extensive genetic heterogeneity and clinical variabilities underscore the intricacy and uniqueness of dynein arms in flagella and cilia. However, the susceptibility of uncharacterized dynein factors that related to asthenoteratozoospermia and/or PCD remains unclear, given the absence of human genetic evidence or corresponding animal models.

Dynein axonemal heavy chain 12, *DNAH12* (MIM: 603340, GenBank: NM_001366028.2), also known as DHC3, is a unique dynein arm component (25). Notably, it is the shortest member and lacks a typical microtubule-binding domain (MTBD) in the DNAH family. Mutations of *DNAH12* have been predicted to be associated with male patients' infertility, however, limited pathogenic evidence was provided, and the underlying pathogenesis remained unclear (26, 27). Besides, although

DNAH12 has been identified as a possible axonemal dynein and its mRNA is predominantly detected in two tissues containing axonemal structures: trachea and testis (25 [↗](#), 28 [↗](#)), its physiological functions remain largely elusive.

In this study, we enrolled 796 infertile patients from unrelated Pakistani and Chinese families diagnosed with asthenozoospermia. Utilizing whole-exome sequencing (WES) followed by Sanger sequencing, we identified six novel bi-allelic mutations in *DNAH12* in infertile males from six unrelated families. All these mutations resulted in the loss of DNAH12 protein in the sperm of patients. Two mouse models, *Dnah12*^{-/-} and *Dnah12*^{mut/mut}, were generated and exhibited asthenoteratozoospermia and male infertility, mimicking our patients' phenotypes. Deficiency in DNAH12 led to axoneme defects characterized by impaired IDAs and CP loss in sperm flagella, while no such defects were observed in cilia. DNAH12 interacts with other IDA components like DNALI1 and DNAH1 in mouse testes, and disruption of DNAH12 impaired the recruitment of dynein components into sperm flagella. Our findings unveil, for the first time, that DNAH12 plays a pivotal role in facilitating the recruitment of IDA components specifically in flagella, but not in cilia. Furthermore, we demonstrate that bi-allelic mutations in *DNAH12* are identified as causative factors for asthenoteratozoospermia and male infertility in both humans and mice. These results enhance our understanding of the development of axonemal structures in flagella versus cilia and provide valuable clues for genetic counseling and future individualized treatment for infertile males carrying *DNAH12* mutations in clinical practice.

Materials and methods

Human subjects

Informed consent and peripheral blood were obtained from the participants and the patients' information was registered in the Human Reproductive Disease Resource Bank database. Sperm analyses were performed according to the WHO guidelines (29 [↗](#)). During their medical consultation, all patients answered a questionnaire regarding clinical presentations. This study was approved by the institutional ethics committee of University of Science and Technology of China (USTC) with the approval number 2019-KY-168.

Whole exome sequencing (WES) and bioinformatics analyses

Whole-exome capture and sequencing were performed by capturing exons from genomic DNAs of peripheral blood cells, using the SureSelect XT 50 Mb Exon Capture Kit (Agilent, California, USA), and then high-throughput sequencing was conducted on the HiSeq 2000 platform (Illumina, San Diego, CA, USA) following instructions. Variants were called using the Genome Analysis Toolkit HaplotypeCaller and annotated through ANNOVAR (30 [↗](#)). Variants within exons or exon-intron boundaries were retained and filtered through the criteria we previously described (31 [↗](#)). Bcftools was applied to detect runs of homozygosity (32 [↗](#)). Runs of homozygosity over 1.5 Mb were used to calculate the inbreeding coefficients. The genotypes were validated using Sanger sequencing validation. Additional primers are listed in Supplemental Table 1. For phylogenetic analysis, different DNAH12 protein sequences were extracted from the UniProt database, and the sequence alignment and phylogenetic tree construction were generated using MEGA11 (33 [↗](#)). Newick format tree files were extracted and processed in the Evolview v2 tool (34 [↗](#)). For gene family and conserved domain analyses, the batch sequence search tool in Pfam was used to analyze features of proteins and further visualized with the assistance of Evolview v2 tool and TBtools (35 [↗](#), 36 [↗](#)). For protein structure analysis, the structure models were predicted and constructed with the Phyre2 tool (37 [↗](#)), interactions were analyzed by the HDock (<http://hdock.phys.hust.edu.cn/> [↗](#)) and AlphaFold3 (<https://golgi.sandbox.google.com/> [↗](#)). The predicted protein structures and possible interactions were further explored and visualized by UCSF Chimera (Version: 1.15, California, USA) according to the manual instruction (38 [↗](#)).

Scanning electron microscopy (SEM) and Transmission electron microscopy (TEM)

Fresh semen samples were obtained and centrifuged at 900 g for 3 min. Spermatozoa were washed with phosphate buffer saline (PBS) three times. Then the sediment was fixed in 0.1 M phosphate buffer (pH 7.4) containing 4% paraformaldehyde, 8% glutaraldehyde, and 0.2% picric acid at 4°C for at least 8 hours. Scanning electron microscopy (SEM) and Transmission electron microscopy (TEM) were then conducted as we previously described (39 [40](#)).

Generation and verification of the polyclonal anti-DNAH12 antibody

The characteristics of the DNAH12 including its region immunogenicity, hydrophilicity, surface leakage groups, and sequence homology were analyzed and the 1-200 amino acids sequence of mouse DNAH12 protein (UniProt accession no. Q69Z23) was regarded as the ideal antigen for immunization. Subsequently, the DNAH12 polyclonal antibody was generated in rabbits and rats using the mentioned peptide by Dia-An Biotech, Inc., in Wuhan, China. The specificity of antibodies was further validated using *Dnah12*^{+/+} and *Dnah12*^{-/-} mice in this study.

Immunoblotting

Protein lysate was prepared as we previously described (40 [41](#)). Tissues or cells were lysed using lysis buffer (50 mM tris (pH 7.5), 150 mM NaCl, 5 mM EDTA, and 0.1% NP-40). For human sperm cells, proteins were extracted using the TRIzol reagent. The protein sediment was washed with 95% ethanol containing 0.3M guanidine hydrochloride, dissolved in 1% SDS. Then the protein lysate was boiled for 10 minutes, and diluted in SDS-sample buffer (62.5 mM TRIS, 2% (w/w) SDS, 10 % (w/v) glycerol, and 5% 2-mercaptoethanol). Whereafter the proteins were separated by SDS-PAGE and transferred onto nitrocellulose blotting membranes with a pore size of 0.45 μm (GE Healthcare, 10600002). The membranes were blocked with TBS-Tween buffer (50 mM Tris, 150 mM NaCl, and 0.5% Tween-20, pH 7.4) containing 5% skim milk for 30 minutes to reduce unspecific binding and then incubated with primary antibodies diluted in TBST buffer containing 5% skim milk at 4°C overnight. Incubation with secondary antibodies for 1.5 hours was followed by chemiluminescence development system (GE Healthcare, Pittsburgh, USA). Antibodies details are listed in Supplemental Table 2-3.

Immunofluorescence (IF) staining

Human semen smear slides were prepared following the guidelines of WHO (29 [30](#)). Mouse sperm smear slide preparation and immunofluorescence staining were conducted as we described (42 [43](#)). For tissue IF staining, tissues were collected and fixed in 4% paraformaldehyde (PFA), the 5μm tissue sections were deparaffinized in xylene, rehydrated in a graded series of ethanol (100%, 95%, 90%, 80%, 70%, and 50% ethanol) and sterile water, and permeabilized with 0.2% Triton X-100 in PBS for 30min. Antigen retrieval was then conducted by boiling slides in a microwave for 30 minutes in 10 mM citrate buffer (pH 6.0). The other steps are consistent with sperm IF staining. Images of spermatozoa were captured using a Nikon ECLIPSE 80i microscope equipped with a charge-coupled device (Hamamatsu Photonics, Hamamatsu, Japan). The fluorescent signal intensity profiles were depicted using Fiji software (<https://imagej.net/software/fiji/> [44](#)). Information of antibodies is available in Supplemental Table 2-3.

Animal models and fertility test

Two mouse models (*Dnah12*^{-/-} and *Dnah12*^{mut/mut}) were generated using CRISPR/Cas9 Technology respectively (43 [44](#)). Briefly, to generate the *Dnah12*^{-/-} mouse model, guide RNAs (gRNAs) targeting exon 5 of *Dnah12*, a key exon shared by all its transcripts, were transcribed in vitro (Thermo

Fisher, CA, USA). Then, the gRNAs and the Cas9 protein were electroporated into C57BL/6 mouse zygotes. The zygotes were introduced into Institute of Cancer Research (ICR) female mice, and the genotypes of the pups were checked using Polymerase Chain Reaction (PCR) with designed primers, the mice with the frameshift mutation c.386_389 del that may lead to paralyzed DNAH12 function were kept for further study. Similar conductions were performed targeting exon 18 of *Dnah12* to generate the *Dnah12*^{mut/mut} mice. The genotyping primers are listed in Supplemental Table 1. For male fertility test, each 8-10 weeks male mouse was caged with wild-type female mice; for female fertility test, each 8-10 weeks female mouse was caged with one 8-10 weeks *Dnah12*^{+/-} male mouse, the number of offspring was recorded.

Reverse-transcription PCR and real-time quantitative PCR

For reverse-transcription PCR (RT-PCR), total RNA was extracted from mouse tissues using TRIzol reagent (TAKARA, Japan). Whereafter 1 µg of RNA was reverse-transcribed using the PrimeScript RT Reagent Kit (TAKARA, Japan). The cDNA was diluted to a comparable concentration and then used for RT-PCR. All amplified DNA fragments were visualized by 2% agarose gel electrophoresis containing GelRed (APEX BIO, Houston, TX, USA). For real-time quantitative PCR (qPCR), the experiments were performed by using Hieff UNICON® Universal Blue qPCR SYBR Green Master Mix reagent (Yeasen Biotechnology, Shanghai, China) on CFX96® Real-Time System (Bio-Rad, CA, USA) according to the manufacturer's guidelines. Details of the primers were listed in Supplemental Table 1.

Histological analysis

As previously described (31), after sacrificing mice, the testes, epididymides, tracheas, and oviducts were harvested and fixed in Bouin's fixative solution (Sigma-Aldrich, Missouri, USA) for hematoxylin and eosin (H&E) staining or 4% PFA fixative solution for immunofluorescence or TUNEL assay. The pictures were captured using a microscope equipped with a charge-coupled device camera (Nikon, Tokyo, Japan).

Sperm counts, motility and morphology analysis

As previously described (44), H&E staining or Papanicolaou staining kit (Solabio, Beijing, China) was used for the evaluation of sperm morphology of patients. For mouse sperm parameters analysis, 8-10-week-old mice were sacrificed by cervical dislocation, the sperm number per epididymis was then assessed by cutting the epididymis into pieces in PBS and incubating them for 30 minutes at 37°C to release the sperm, then counting with a hemocytometer. Sperm motility were assessed by computer-aided sperm analysis (CASA) as we described (14). Sperm morphology was assessed on spermatozoa collected from epididymal cauda after H&E staining.

Testicular cell smear slides and seminiferous tubule squash slides preparation

To prepare testicular cell smears, mice were sacrificed and testes were then detached with one time of PBS washing. Then testes were peeled off the testicular capsule, torn into small pieces, and then cut into tiny pieces in PBS. Finally, the cell suspension was filtered through a 200-mesh cell strainer, the filtered cell suspensions were added 10% FBS before spread onto glass slides. For testicular seminiferous tubule squash slide preparation, the testes were dissected and carefully isolated into single testicular seminiferous tubules in PBS dishes. After 10 mins of prefixation in fix/lysis solution (2% PFA diluted in PBS), we then applied 100 µL fix/lysis solution onto poly-L-lysine coated glass slides and carefully teased apart approximately 20 seminiferous tubules with the assistance of 2 sterile forceps on the slides. After that, coverslips were applied using slide forceps and pressed with the heel of the palm for 10 seconds, then the glass slides were frozen in a small

container with liquid N₂ immediately till the liquid ceases to bubble. The slides were saved in the -80°C fridge keeping the coverslip on the slides or removing coverslips for immunofluorescence staining.

TUNEL assay

As previously described (44 [DOI](#)), In Situ Cell Death Detection Kit, Fluorescein (Sigma–Aldrich, Missouri, USA) was applied according to the manufacturer’s protocol. Images were captured using a Nikon Eclipse 80i microscope equipped with Hamamatsu C4742-80 digital camera and analyzed using the NIS-Element Microscope imaging software (Nikon, Tokyo, Japan).

Cell culture and transfection

As previously described (40 [DOI](#)), HEK293T (ATCC, VA, USA) cells were cultured to 70%-80% density in 12-well plates, followed by transfected with plasmids expressing mCherry-tagged mouse DNAH12 and GFP-tagged mouse DNALI1 using Lipofectamine 3000 (Thermo Fisher, CA, USA) according to the manufacturer’s instructions. Cells were harvested for in-vitro IP and immunoblotting the next day after transfection. The sequences of the primers for plasmid construction are listed in Supplemental Table 4.

Co-immunoprecipitation (Co-IP)

Tissue extracts were prepared using lysis buffer supplemented with PMSF, Protease Inhibitor Cocktail (Solarbio, Beijing, China) on ice using a glass homogenizer. Lysate was then centrifuged at 12,000 g at 4°C for 10 min, and the supernatant was divided into equal aliquots. Each aliquot was incubated with 1.5 µg of target antibody or rabbit IgG nonspecific antibody. After incubation at 4°C overnight with rotation, beads were washed five times with lysis buffer. Finally, the beads were resuspended in SDS-sample buffer and boiled at 100°C for 10 min. The samples were either analyzed by immunoblotting immediately or stored at -80°C. For Co-IP in cultured cells, HEK293T cells were plated in six-well plates, followed by transfection using Lipofectamine 3000 reagents. MG132 (5 µM) was added into the cultured cells 6 hours before cell harvesting. Cells were washed with PBS for 10 min, cells were then collected and lysed in the same lysis buffer used for Co-IP from testicular samples. The cell lysate was then incubated with prewashed Protein A/G Magnetic Beads B23202 (Bimake, Texas, USA) and primary antibodies at 4°C overnight, followed by washing and boiling as described above for the Co-IP protocols for testicular samples.

Mass spectrometry proteomics

To identify potential partners that interact with DNAH12, we performed IP of mouse testes lysate. Tissue extracts were prepared using lysis buffer supplemented with PMSF, Protease Inhibitor Cocktail (Solarbio, Beijing, China) with the same operation of the Co-IP protocol. Beads were washed 3 times with lysis buffer and 3 times with 50 mM NH₄HCO₃, the pull-down proteins were digested with 0.2 µg Trypsin overnight and stopped by 5% Formic acid, (Thermo Fisher, CA, USA). The peptide mixture was then desalted, dried with a SpeedVac, and resuspended in 0.1% Formic acid, followed by the MS analysis using timsTOF Pro 2 mass spectrometer (Bruker, Massachusetts, USA). The proteomics data are available on the ProteomeXchange Consortium via the iProX partner repository (45 [DOI](#)), with the dataset identifier PXD051681.

Intracytoplasmic sperm injection (ICSI)

3.5-4-week-old female mice were given an intraperitoneal injection of 5 IU Pregnant Mare Serum Gonadotropin (PMSG, Ningbo Sansheng, China). After 48 hours, a second intraperitoneal injection of 5 IU human Chorionic Gonadotropin (hCG, Ningbo Sansheng, China) was administered. The mice were euthanized by cervical dislocation after 15 hours post-hCG injection. The oviducts were excised and the ampullary regions were opened to release the cumulus-oocyte complexes (COCs). The cumulus cells surrounding the oocytes were digested using hyaluronidase, and the

morphologically round MII stage oocytes were selected and washed three times in M2 medium (Sigma–Aldrich, Missouri, USA) before being set aside for use. Male mice are euthanized by cervical dislocation, the epididymides are excised and gently punctured with a syringe, allowing the spermatozoa to flow out. The spermatozoa were then placed in the human tubal fluid (HTF) medium (EasyCheck, Nanjing Aibei Biotechnology, China) and then frozen and thawed repeatedly to remove sperm tails, the single sperm head was microinjected into an MII oocyte and then cultured in potassium simplex optimized medium (KSOM, EasyCheck, Nanjing Aibei Biotechnology, China) at 37 °C under 5% CO₂.

Statistics analysis

Statistical analyses were performed using GraphPad Prism 8.0 (GraphPad Software, San Diego, CA, USA). Values in tables and graphs are expressed as mean ± SEM. Two groups of mice were compared with Student's 2-tailed t-test for independent data and $P > 0.05$ was considered not significant (ns); * $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$, and **** $P < 0.0001$ were considered significant.

Results

Identification of biallelic *DNAH12* variants in men with asthenoteratozoospermia

To identify the genetic factors contributing to male infertility, we recruited patients with primary infertility and their family members from Pakistan and China. WES was performed on these individuals, followed by comprehensive analyses of the WES data (Supplemental Figure 1, A–D). Seven mutations in *DNAH12* were identified as potential pathogenic factors for male infertility in patients P1–3 of family 1, P4–5 of family 2, P8 of family 3, and P9, P10, and P11, all of whom suffered from asthenoteratozoospermia (Fig. 1A). Initially, a homozygous frameshift mutation c.944_945 del (p. Phe315*) in family 1, and two homozygous missense mutations, c. A164G (p. Gln55Arg) in family 2 and c. A2560G (p. Ile854Val) in family 3, were identified in *DNAH12* as potential causes of infertility. Subsequently, four other mutations in *DNAH12* were discovered in three sporadic Chinese patients with asthenoteratozoospermia: a homozygous stop-gain mutation c.2680 C>T (p. Gln2077*) in subject P9, a homozygous missense mutation c.2813T>C (p. Val938Ala) in P10, and compound heterozygous mutations c.2680C>T (p. Arg894Cys) and c.5964-3A>G in P11 (Fig. 1A and B).

All seven variants were rare in public human genome databases, including the 1000 Genome Project and gnomAD (v3). Additionally, these variants were predicted to be deleterious by in-silico software tools such as SIFT, Polyphen2, fathmm-MKL, and CADD2 (Table 1). Sanger Sequencing further validated these variants and revealed the recessive pathogenic pattern of these mutations (Fig. 1A). Collectively, these results suggested that the identified bi-allelic *DNAH12* variants could potentially be the cause of asthenoteratozoospermia in our patients.

DNAH12 protein is conserved in mammals and highly expressed in the testes

DNAH12 encodes a functionally uncharacterized protein and is predicted to be abundantly expressed in testes of humans and mice based on our bioinformatics tool FertilityOnline (46). Experimentally, our RT-PCR, qPCR, and immunoblotting detection of *DNAH12* expression in different tissues indicated that both *DNAH12* mRNA and protein showed high expression in testes and relatively lower expression in lungs and tracheas of male mice and oviducts of female mice (Supplemental Figure 2, A–C). It's worth noting that in epididymides, *DNAH12* protein, but not mRNA, was detected, indicating the *DNAH12* protein detected in epididymides is from testes, most

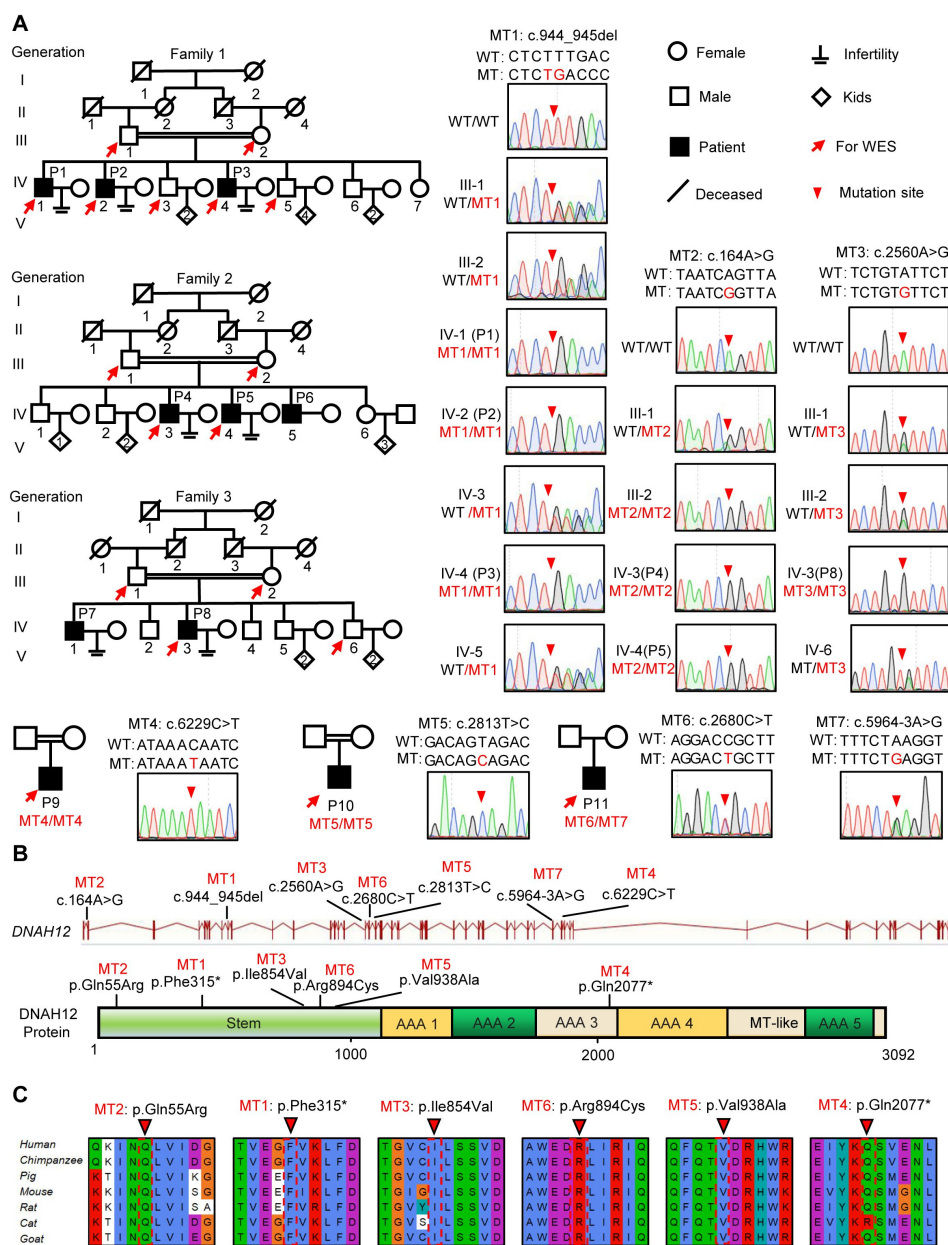


Figure 1.

Identification of bi-allelic *DNAH12* variants in infertile men with asthenoteratozoospermia

(A) Pedigrees of family 1 with 3 infertile males, P1 (IV:1), P2 (IV:2), and P3 (IV:4), family 2 with 3 infertile males, P4 (IV:3), P5 (IV:4) and P6 (IV:5), family 3 with 2 infertile males, P7 (IV:1) and P8 (IV:3), and three sporadic infertile cases P9, P10 and P11. Red arrows point to the individuals whom whole-exome sequencing was performed. Candidate mutations were validated by Sanger sequencing and the chromatograms were shown on the right. Red arrowheads point to mutant sites. Double horizontal lines represent consanguineous marriages. WT, wild-type allele; MT, the mutant allele. (B) The positions of the *DNAH12* variants at the transcript (ENST00000351747.2) and protein levels (Q6ZR08-1). MT-like, Microtubule-binding-like domain. (C) Conservation analyses of affected amino acids across different species, the arrowheads indicate the mutation sites.

	Genomic position on chr3	cDNA Change ^a	Mutation type	Genetic pattern	SIFT	Polyphen2	fathmm-MKL	CADD	1000 Genomes	gnomAD
Family 1	57,489,883	MT1: c.944_945del	frameshift substitution	AR	NA	NA	NA	NA	0	0.00002844
Family 2	57,528,434	MT2: c.164A>G	nonsynonymous SNV ^b	Male-limited AR	T	D	D	D	0.000599042	0.0004
Family 3	57,447,323	MT3: c.2560A>G	nonsynonymous SNV	AR	T	D	D	D	0.000599042	0.0002
P9	57,391,670	MT4: c.6229C>T	stop-gain	AR	NA	NA	D	NA	0	0.00002714
P10	57,445,368	MT5: c.2813T>C	nonsynonymous SNV	AR	D	D	D	D	0.000199681	0.00003383
P11	57,394,265	MT6: c.5964-3A>G	splicing	AR	NA	NA	NA	NA	0.000199681	0.0002
	57,445,501	MT7: c.2680C>T	nonsynonymous SNV		D	D	D	D	0	0.000007178

a. The accession number of human *DNAH12* is ENST00000351747.2. **b.** Abbreviations are listed as follows: SNV, single nucleotide variants; NA, not assessed; T, tolerated; D, deleterious. SIFT, Sorting Intolerant From Tolerant; PolyPhen-2, Polymorphism Phenotype v2; FATHMM-MKL, Functional Analysis Through Hidden Markov Models-Multiple Kernel Learning; CADD, Combined Annotation-Dependent Depletion; and a mutation is predicted deleterious if the CADD Phred score > 20 in CADD. gnomAD, Genome Aggregation Database.

Table 1

Detailed information of bi-allelic *DNAH12* variants identified in infertile men

probably from sperm. Furthermore, DNAH12 protein was initially detected in testes around 21 days postpartum (dpp) and thereafter maintained a high level (Supplemental Figure 2D). The expression pattern of DNAH12 in testes suggests that it is possibly involved in spermiogenesis, and its abnormal expression could lead to asthenozoospermia or asthenoteratozoospermia, further indicating that the identified *DNAH12* mutations are candidate pathogenic mutations for patients.

Further phylogenetic analysis revealed the conservative nature of DNAH12 from *Chlamydomonas* to humans (Supplemental Figure 3). All the amino acids affected by mutations were conserved in mammals and were located in functional regions, including the ATPase domains or the Stem domain of DNAH12 (Fig. 1C), indicating that these mutations are highly likely to hinder the structure, function, or stability of the DNAH12 protein. Altogether, our results suggested that the identified mutations in *DNAH12* are probable pathogenic factors contributing to asthenoteratozoospermia in our patients.

Impaired sperm motility and morphology in patients with bi-allelic *DNAH12* variants

To assess the pathological effects of the identified bi-allelic *DNAH12* variants, semen analyses were conducted following WHO guidelines (29). All affected individuals exhibited significantly reduced sperm motility and progressive motility (Table 2). Detailed sperm morphology analyses were performed on patients P3 from family 1, P5 from family 2, P8 from family 3, and three Chinese sporadic patients (P9, P10, and P11). Spermatozoa from fertile controls displayed normal heads and tails, while spermatozoa from the patients exhibited a range of abnormal morphologies, including short, coiled, absent, bent, or irregular caliber flagella, accompanied by aberrant heads (Fig. 2A, Supplemental Figure 4, A and B). Scanning electron microscopy (SEM) analyses were further carried out on a fertile control and P10, confirming the presence of abnormal morphologies of the sperm head and flagellum in the patients (Fig. 2B). Thus, patients with the bi-allelic *DNAH12* variants demonstrated significantly decreased sperm motility and increased abnormal sperm morphology.

The bi-allelic *DNAH12* variants result in absence of DNAH12 protein in patients' sperm

To investigate the effects of the identified bi-allelic *DNAH12* variants on the DNAH12 protein, we obtained sperm samples from three affected Chinese patients, and the expression of DNAH12 was detected using a custom-made DNAH12 rabbit antibody that recognized the N-terminal 1-200 amino acids of the DNAH12 protein. The validated antibody could theoretically detect the truncated protein if it were present in P9 with a homozygous stop-gain mutation M4 (p. Gln2077*). The predicted DNAH12 protein size was detected in the sperm lysate from a fertile control but not in P9 with the homozygous stop-gain mutation M4, P10 with a homozygous missense mutation M5, and P11 with compound heterozygous mutations M6 and M7, indicating these mutations possibly lead to the disruption of the DNAH12 protein (Fig. 2C).

To further explore the effects of the identified *DNAH12* variants on the DNAH12 protein, we performed immunofluorescence assays on sperm from patients. The results showed that DNAH12 signals were localized along the flagella of the fertile control, however, no DNAH12 signals were observed in sperm flagella of patients P3 (family 1), P5 (family 2), P8 (family 3), P9, P10, and P11 (Fig. 2D). Collectively, these findings indicated that all the detected *DNAH12* variants cause the absence of DNAH12 protein in the sperm of our patients, reinforcing that these *DNAH12* mutations are pathogenic mutations of asthenoteratozoospermia in the patients.

	Reference values ^a	Fertile controls ^b	IV-1(P1)	Family1 IV-2(P2)	IV-4(P3)	Family2 IV-5(P4)	IV-7(P5)	Family3 IV-4(P8)	P9	P10	P11
cDNA mutation	–	–	MT1/MT1	MT1/MT1	MT1/MT1	MT2/MT2	MT2/MT2	MT3/MT3	MT4/MT4	MT5/MT5	MT6/MT7
Age (y) ^c	–	–	58	49	42	40	38	32	33	40	31
Semen parameters											
Semen volume (ml)	>1.5	–	ND	1.5±0.5	2.8±0.3	4.0	3.0	3.0	3.2	3.1	2.4
Sperm concentration (10 ⁶ /ml)	>15	–	ND	58±28.5	50.0±15.0	6.0	8.0	2.8	24.7	83.1	57.1
Semen pH	Alkaline	–	ND	Alkaline	Alkaline	Alkaline	Alkaline	Alkaline	Alkaline	Alkaline	Alkaline
Motile sperm (%)	>40	–	ND	29.3±9.0	7.5±7.5	14.0	20.0	20	15.1	17.4	7.4
Progressively motile sperm (%)	>32	–	ND	13.3±10.3	16.7±17	5.0	8.0	5	13.4	16.8	24
Sperm morphology											
Normal (%)	–	76.5±2.0	ND	ND	19.9 ^d	ND	21.7	17.0	14.0	17.4	7.4
Abnormal head (%)	–	7.0±1.9	ND	ND	16.4	ND	5.4	6.0	3.5	16.4	20.0
Abnormal tail (%)	–	12.4±3.4	ND	ND	43.8	ND	41.4	44.0	42.4	37.4	16.3
Abnormal head & tail (%)	–	4.1±1.3	ND	ND	19.9	ND	31.5	33.0	40.2	28.8	56.3
Sperm flagella morphology											
Normal (%)	–	83.1±2.3	ND	ND	26.4	ND	26.6	22.7	17.9	31.7	27.4
Absent (%)	–	2.7±1.1	ND	ND	16.4	ND	7.9	15.8	9.6	16.6	11.6
Short (%)	–	3.6±0.6	ND	ND	16.4	ND	11.3	14.9	20.1	11.7	8.4
Coiled (%)	–	2.8±0.7	ND	ND	17.3	ND	24.6	12.9	11.8	14.6	7.0
Bent (%)	–	6.3±0.6	ND	ND	12.7	ND	15.8	10.0	13.5	7.8	33.5
Irregular caliber (%)	–	1.5±0.2	ND	ND	10.9	ND	13.8	23.8	27.1	17.6	12.1

a. The reference limits are shown according to WHO 6th edition standards. **b.** Reference values are shown by observation of sperm morphology in 3 fertile individuals. **c.** "Age" represents age in 2024. ND, not determined. **d.** Values in red colors show obvious declined proportions of normal sperm morphology compared with fertile controls.

Table 2

Clinical characteristics of the infertile patients with bi-allelic *DNAH12* mutations

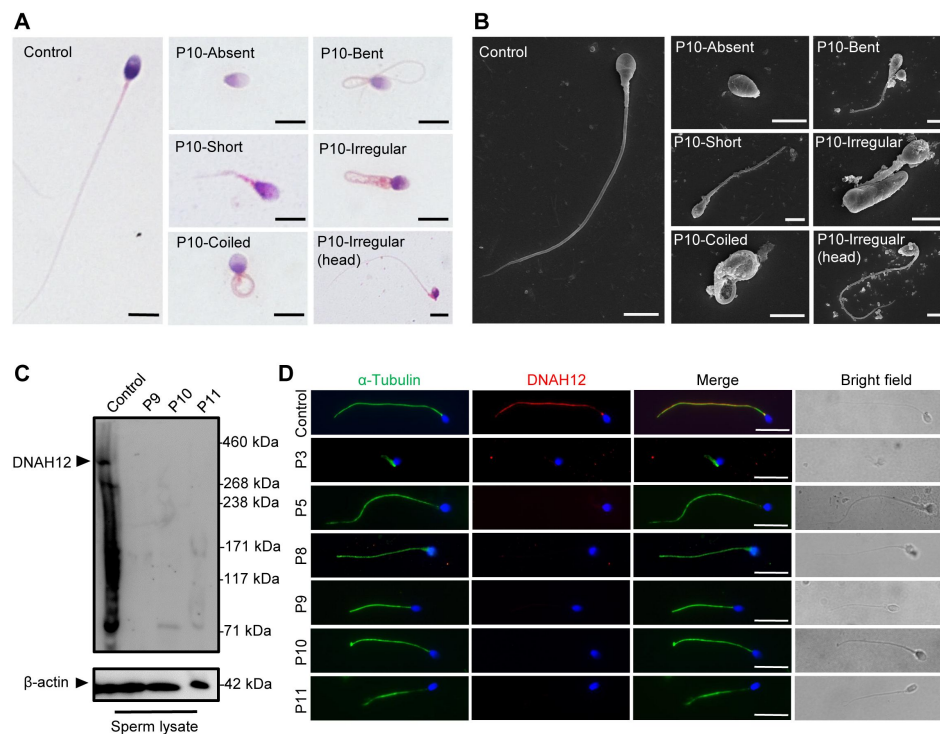


Figure 2.

Abnormal sperm morphology and undetectable DNAH12 expression in men with bi-allelic *DNAH12* variants

(A) Representative micrographs show normal sperm morphology from a fertile control and abnormal spermatozoa from P10 with absent, short, coiled, bent, irregular flagella or abnormal head under light microscopy. Scale bars, 5μm. (B) Representative SEM micrographs show details about the sperm morphology of a fertile control and P10. The normal sperm morphology can be observed in the fertile control (left) and abnormal types like absent, short, coiled, bent, irregular tails, and head abnormality in P10 (right). Scale bars, 5μm. (C) Immunoblotting assay revealed that DNAH12 was absent in the spermatozoa from P9, P10, and P11 harboring *DNAH12* variants. β-actin was used as a loading control. (D) Sperm cells were co-stained with α-Tubulin (green) and DNAH12 (red) antibodies while no DNAH12 signals were observed in sperm flagella of patients. DNA was counterstained with Hoechst 33342. Scale bars, 10μm.

Patients with bi-allelic *DNAH12* variants exhibit flagellar axoneme defects

Since patients with the bi-allelic *DNAH12* variants exhibited compromised sperm motility and abnormal sperm morphology, we then investigated whether sperm flagellar ultrastructure was also affected in these patients. TEM examination was performed on sperm flagella from both patients and normal fertile controls. The flagella of fertile controls displayed the typical “9+2” microtubule structure with well-organized DMTs and CP. In contrast, various defects were found in our patients, including loss of CP (54.5%), disorganization of microtubules (26%), and absence of DMTs (16.5%) in the examined cross-sections (Fig. 3A, 3C and Supplemental Figure 5A). To examine the IDAs impairment in axonemes, the sections with intact ODAs were analyzed. Compared with axonemes in controls that almost all the sections presented intact IDAs, approximately 70% of sections examined showed impaired IDA structures in the analyzed three patients (Fig. 3B, 3D and Supplemental Figure 5A).

We then performed immunofluorescence assays on sperm smears to assess the expression and localization of specific proteins of the axonemes. These included SPAG6 (a CP component), DNAH1 and DNALI1 (IDA components), DNAH17 (an ODA component), and ODF2 (an outer dense fiber, ODF component). The sperm flagella of patients showed remarkably reduced SPAG6 signals when compared with those from controls (Fig. 3E). Similarly, DNAH1 and DNALI1 signals were both obviously decreased in the patients’ sperm flagella, indicating impaired IDAs in the patients (Fig. 3F and G). Besides, DNAH17 and ODF2 signals in patients’ flagella were comparable to those in a normal control (Fig. 3H and Supplemental Figure 5B). In summary, these findings indicated that the bi-allelic *DNAH12* variants cause pronounced flagellar axoneme defects, including impaired IDAs and loss of CP, but do not affect ODAs and ODFs.

Deletion of *Dnah12* impairs spermiogenesis and causes male infertility in mice

Considering the bi-allelic *DNAH12* variants resulting in absence of DNAH12 protein in patients’ sperm, to functionally confirm the deletion of *DNAH12*, we generated *Dnah12* knockout mice using CRISPR-Cas9 approach (Supplemental Figure 6A). Fortunately, we obtained a mouse model with a 4-bp deletion (c.386_389del) in exon 5 of *Dnah12* (Supplemental Figure 6A). In contrast to the observations in control mice, immunoblotting of testis lysate using two antibodies targeting the N-terminal of DNAH12 failed to detect DNAH12 proteins, and immunofluorescence staining of sperm flagella did not show DNAH12 signals in *Dnah12*^{-/-} mice (Supplemental Figure 6, B and C). *Dnah12*^{-/-} males exhibited normal development but were infertile (Fig. 4A), while *Dnah12*^{-/-} female mice showed normal development and fertility, and no obvious defective oviductal and tracheal cilia (Supplemental Figure 7, A-C). The body weight of *Dnah12*^{-/-} and *Dnah12*^{+/+} males was comparable, while smaller testes were observed in *Dnah12*^{-/-} males (Fig. 4B). Besides, the average number of sperm per epididymis of *Dnah12*^{-/-} mice dramatically decreased to approximately 2% of that of *Dnah12*^{+/+} mice (Fig. 4C). The epididymal caudal spermatozoa of *Dnah12*^{-/-} male mice were 100% immotile, compared to those of *Dnah12*^{+/+} mice (Fig. 4D, Supplemental Videos 1 and 2). H&E staining of testis and epididymis sections also showed a striking reduction in the number of sperm in *Dnah12*^{-/-} mice (Fig. 4E). Importantly, morphological analysis of sperm from the caudal epididymis of *Dnah12*^{-/-} mice showed similar types and frequencies of flagellum and head abnormalities to those observed in our patients (Fig. 4F and G).

To understand how *DNAH12* depletion causes abnormal sperm morphology, we conducted immunofluorescence assays on testicular cells from *Dnah12*^{+/+} and *Dnah12*^{-/-} mice, in *Dnah12*^{-/-} mice, while DNAH12 signals are missing in round, elongating, and elongated spermatids; the abnormal manchette structure and flagellum were observed (Fig. 4H). We then performed

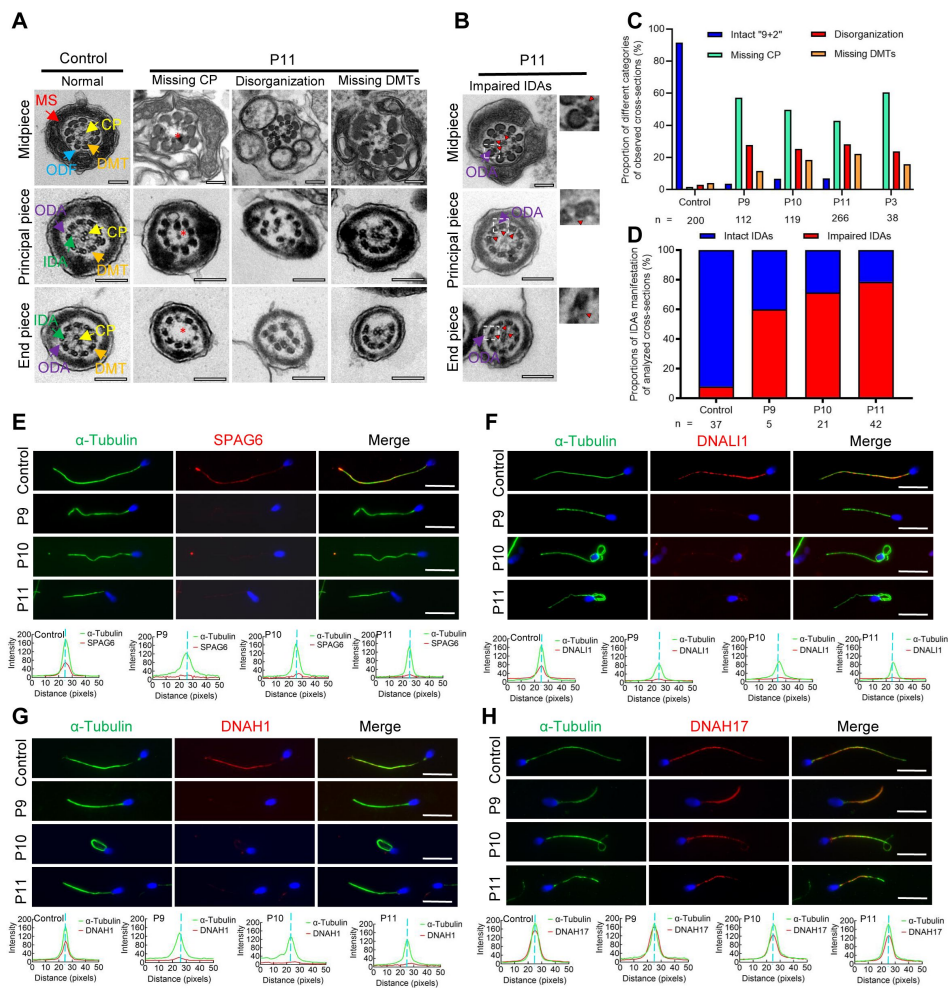


Figure 3.

Flagellar axoneme defects were detected in the patients with *DNAH12* variants

(A-B) Representative TEM micrographs showing cross-sections of the midpiece, principal piece, and end piece of sperm flagella from a fertile control and P11. The axoneme structure in control presents a "9+2" microtubules arrangement, including mitochondrial sheath (MS, indicated with red arrows), central pair of microtubules (CP, indicated with yellow arrows), outer dense fibers (ODFs, indicated with cerulean arrows), peripheral doublet microtubules (DMTs, indicated with orange arrows), inner dynein arms (IDAs, indicated with green arrows) and outer dynein arms (ODAs, indicated with purple arrows), while in P11, axonemal defects like missing CP, disorganized axonemal structures or missing DMTs were observed, the red asterisks mark CP loss (A). Impaired IDAs in the sperm axoneme of patient P11 (B). The red triangle marks impaired IDAs while adjacent ODAs are identifiable. Scale bars, 200 nm. (C-D) The proportion of different categories of observed cross-sections in the control and patients. Cross-sections were classified into four categories: Intact "9+2", missing CP, disorganization, and missing DMTs (C), and the proportion of intact IDAs or impaired IDAs in the cross-sections of flagellar axoneme of which ODAs were identifiable (D). n, the total number of cross-sections for quantification. (E-G) Representative images of spermatozoa from fertile controls and patients carrying bi-allelic *DNAH12* variants co-stained α -Tubulin with SPAG6 (E), DNAH1 (F), DNALI1 (G), or DNAH17 (H), and Hoechst 33342 for DNA (blue). The fluorescent signal intensity profiles were shown on the bottom. Scale bars, 10 μ m.

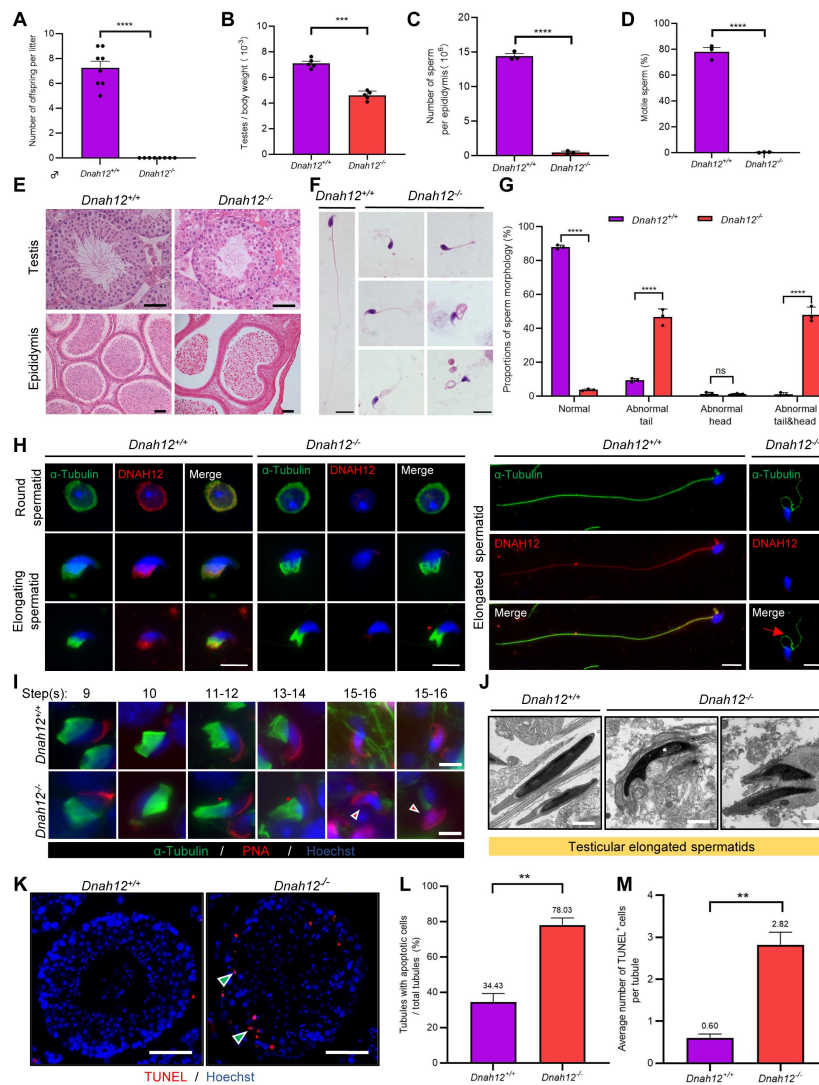


Figure 4.

Disruption of *Dnah12* results in impaired spermatogenesis and abnormal spermatozoa morphology

(A) Fertility test of adult *Dnah12*^{+/+} and *Dnah12*^{-/-} male mice. (B) The testes to body weight ratios of *Dnah12*^{+/+} and *Dnah12*^{-/-} mice. (C) The number of sperm per epididymis of *Dnah12*^{+/+} and *Dnah12*^{-/-} male mice. (D) Percentages of motile spermatozoa in *Dnah12*^{+/+} and *Dnah12*^{-/-} male mice. (E) Histological sections of testis and epididymis from *Dnah12*^{+/+} and *Dnah12*^{-/-} mice after H&E staining. Scale bars, 50 μ m. (F) Morphology of the sperm from caudal epididymis of *Dnah12*^{+/+} and *Dnah12*^{-/-} mice. Scale bars, 10 μ m. (G) Quantitative analysis of sperm morphology of *Dnah12*^{+/+} and *Dnah12*^{-/-} male mice. The experiments were repeated three times with at least 200 spermatozoa counted every time. The above data were obtained from 3 adult mice for each genotype. (H) Representative images of testicular cells from *Dnah12*^{+/+} and *Dnah12*^{-/-} mice co-stained DNAH12 and α -Tubulin antibodies. Scale bars, 10 μ m. The red asterisk indicates abnormal manchette structure and the red arrow indicates abnormal sperm flagellum at the elongated spermatid stage of *Dnah12*^{-/-} mice. (I) Representative immunofluorescence images of seminiferous tubule squash from 8-week-old *Dnah12*^{+/+} and *Dnah12*^{-/-} mice, co-stained by PNA (red) and α -Tubulin (green). DNA was stained with Hoechst 33342. Scale bars, 10 μ m. The asterisks indicate the manchette defects and the red triangles mark the abnormal sperm heads. Scale bars, 10 μ m. (J) Representative TEM micrographs of testicular elongated spermatids from *Dnah12*^{+/+} and *Dnah12*^{-/-} mice. Scale bars, 1 μ m. (K) TUNEL assay on the testicular sections from *Dnah12*^{+/+} and *Dnah12*^{-/-} mice. The green triangles mark the apoptotic elongating or elongated spermatids. Scale bars, 50 μ m. (L) The proportion of tubules with apoptotic cells in *Dnah12*^{+/+} and *Dnah12*^{-/-} mice. (M) Average numbers of TUNEL-positive cells per tubule in *Dnah12*^{+/+} and *Dnah12*^{-/-} mice. Data are obtained from three mice with at least 50 tubules scored in each repeated experiment. ***p* < 0.01; ****p* < 0.001; *****p* < 0.0001, Student's *t* test.

immunofluorescence assays on testicular sections using α -Tubulin and lectin peanut agglutinin (PNA, a marker of the acrosome) to examine the process of spermiogenesis. The manchette structure in *Dnah12*^{-/-} was roughly normal until steps 9-10 (Stage IX-X) but was aberrantly stretched from steps 11-12 (Stage XI-XII) when compared with those of *Dnah12*^{+/+} mice (Supplemental Figure 8). To meticulously characterize these defects, we stained α -Tubulin and PNA on squashed seminiferous tubules and found obvious manchette defects and abnormal sperm heads in *Dnah12*^{-/-} mice (Fig. 4I). TEM analysis showed severely abnormal head morphology in elongated spermatids of *Dnah12*^{-/-} males when compared with those of *Dnah12*^{+/+} males (Fig. 4J). Additionally, TUNEL assays of testis sections showed dramatically increased proportions of seminiferous tubules containing TUNEL-positive cells in *Dnah12*^{-/-} mice, compared with those in *Dnah12*^{+/+} mice, suggesting the aberrant spermatids may undergo apoptosis (Figure 4K-M). Altogether, these results indicated that deletion of *Dnah12* impairs spermiogenesis, results in decreased sperm counts, and increased sperm morphological abnormalities in mice.

Deletion of *Dnah12* causes axoneme defects in sperm flagella but not cilia in mice

TEM analysis was conducted on sperm from cauda epididymis of *Dnah12*^{+/+}, *Dnah12*^{+/-} and *Dnah12*^{-/-} mice. Flagellar axonemes of *Dnah12*^{+/+} and *Dnah12*^{+/-} mice displayed a typical “9+2” microtubular arrangement with clearly visible CP, IDAs, and ODAs. In contrast, sperm from *Dnah12*^{-/-} mice exhibited severe abnormalities in all the analyzed sections, characterized by CP loss and disorganized axoneme structure (Fig. 5A). Immunoblotting was performed to assess the expression levels of DNAH1, DNALI1, and SPAG6 in sperm lysates, *Dnah12*^{-/-} sperm showed a significant reduction in the abundance of DNAH1 and DNALI1, while the amount of DNAI2 (a component of ODAs) remained comparable to *Dnah12*^{+/+} and *Dnah12*^{+/-} sperm (Fig. 5B). Additionally, immunofluorescence assays revealed the near absence of signals for SPAG6, SPEF2 (a component of CP complex), DNAH1, and DNALI1 in *Dnah12*^{-/-} sperm flagella, whereas DNAH17 and DNAI2 (components of ODAs) remained unaffected (Fig. 5C-H). These results indicate that DNAH12 deficiency specifically impairs CP and IDAs, but not ODAs, in sperm flagella. Furthermore, immunoblotting of SPAG6 and DNALI1 in testis lysate exhibited a sharp decrease in *Dnah12*^{-/-} mice (Supplemental Figure 9A). Additionally, TEM examination of testicular sperm axonemes showed defects like loss of CP and disarranged axonemes in *Dnah12*^{-/-} mice (Supplemental Figure 9B). These results indicate that impaired IDAs and loss of CP possibly occur during testicular sperm flagellar development.

Another *Dnah12* mutant mouse model (*Dnah12*^{mut/mut}) targeting exon18 was also generated (Supplemental Figure 10A), which may produce a truncated DNAH12 protein (p.Cys793Glu, fs*10). The mutation effect was checked but no truncated DNAH12 protein was detected, indicating that DNAH12 expression was disrupted in *Dnah12*^{mut/mut} mice (Supplemental Figure 10B). Moreover, *Dnah12*^{mut/mut} mice developed normally, while decreased testicular size was observed (Supplemental Figure 10, C and D). Consistent with *Dnah12*^{-/-} mice and patients carrying bi-allelic *DNAH12* mutations, *Dnah12*^{mut/mut} presented obviously decreased sperm count, increased sperm morphological abnormalities, and sperm axoneme defects similar to *Dnah12*^{-/-} mice (Supplemental Figure 10, E-H). Therefore, these results reconfirmed that DNAH12 is essential for sperm flagellar development.

Given the detection of DNAH12 protein in the trachea and lung (Supplemental Figure 2, A-C), we also examined whether the deficiency of DNAH12 would cause PCD symptoms in males. Intriguingly, both *Dnah12*^{mut/mut} and *Dnah12*^{-/-} males developed normally and showed no tracheal motility or structure defects, despite the expression of DNAH12 being detected in tracheal cilia cells (Supplemental Figure 11, A-E; Supplemental Videos 3 and 4). Besides, the ultra-structures of cilia axonemes were comparable in wild-type and *Dnah12*^{-/-} groups (Supplemental Figure 11F). These findings collectively suggested that DNAH12 deficiency is crucial for sperm axonemal integrity in humans and mice without manifesting PCD symptoms.

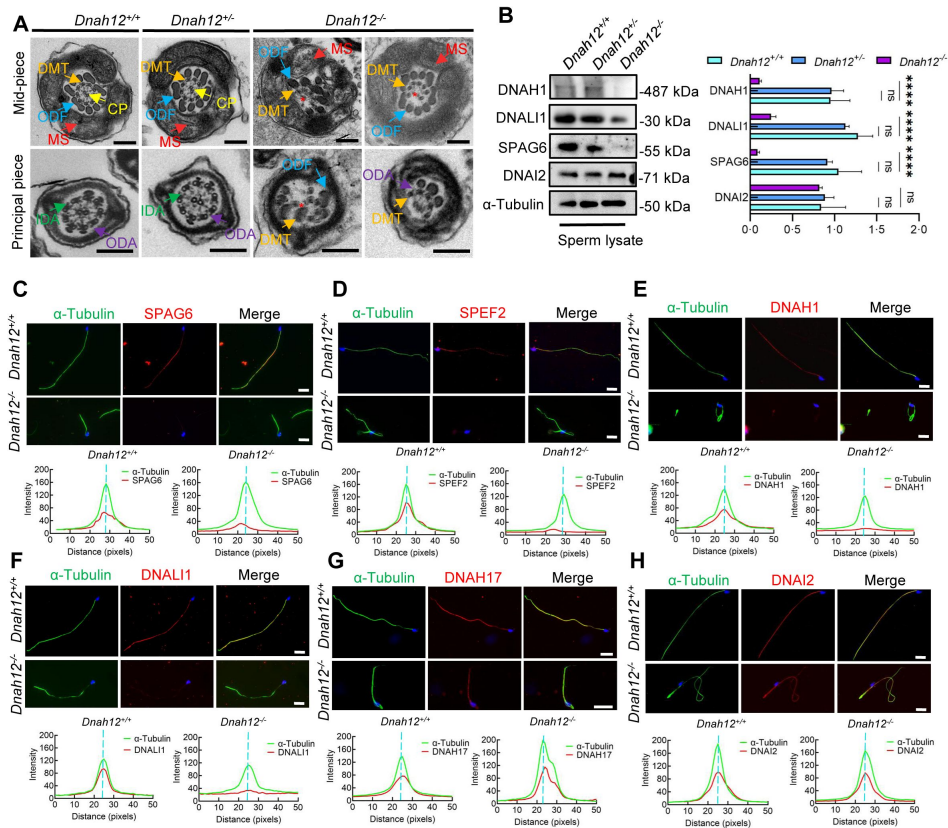


Figure 5.

Deletion of *Dnah12* causes axoneme defects in mice

(A) Representative TEM micrographs showing cross-sections of sperm flagella from *Dnah12*^{+/+}, *Dnah12*^{+/-}, and *Dnah12*^{-/-} mice. In *Dnah12*^{+/+}, *Dnah12*^{+/-} groups, the normal axoneme “9+2” microtubule arrangement mainly consisted of the MS, (indicated with red arrows), CP (yellow arrows), ODFs (cerulean arrows), DMTs (orange arrows), IDAs (green arrows) and ODAs (purple arrows). However, the CP and DMTs structures were missing or disarranged in the *Dnah12*^{-/-} group. Red asterisks mark the loss of CP. Scale bars, 200 nm. (B) Immunoblotting of sperm lysate from *Dnah12*^{+/+}, *Dnah12*^{+/-} and *Dnah12*^{-/-} mice using DNAH1, DNALI1 or SPAG6 antibodies. DNAI2 and α -Tubulin were used as the loading controls. 4 replicates of western blotting were performed. The relative band grayscale of proteins to α -Tubulin in sperm lysate from *Dnah12*^{+/+}, *Dnah12*^{+/-} and *Dnah12*^{-/-} mice were shown on the right. Data are mean $\pm$ SEM. ns: Not significant; (C-H) Representative images of caudal epididymal sperm from *Dnah12*^{+/+} and *Dnah12*^{-/-} mice co-stained α -Tubulin and SPAG6 (C), SPEF2 (a component of CP complex) (D), DNAH1 (E), DNALI1 (F), DNAH17 (G) or DNAI2 (H) antibodies. The fluorescent signal intensity profiles were shown on the bottom. Scale bars, 10 μ m.

DNAH12 interacts with IDA components DNAH1 and DNALI1

Through phylogenetic and domain analyses of DNAH family members, we found that DNAH12 is a distinctive IDA lacking the microtubule-binding domain (MTBD), which is crucial for ATP-sensitive microtubule binding and motor direction control (Fig. 6A, Supplemental Figure 12). Therefore, DNAH12 likely collaborates with other members to carry out its functions. To delve into the molecular mechanism of DNAH12, we conducted co-immunoprecipitation (Co-IP) followed by mass spectrometry analysis using a DNAH12 rabbit antibody in testicular lysate from adult *Dnah12*^{+/+} mice (Fig. 6B). Gene ontology analysis showed that the potential interactors were mainly enriched in motor protein, actin binding, myosin and RNA binding proteins (Fig. 6C). The motor proteins were predominantly categorized into three groups: dynein, kinesin, and myosin protein members, interestingly, among these, dynein proteins such as DNAH1, DNALI1, DNAH1, DNAH3, and DNAH2 were annotated to be involved in IDA assembly, which is closely related to the defects found in *DNAH12* mutants (Fig. 6D). This piqued our interest to explore these proteins in detail. Besides the predominant enrichment of DNAH12 (top1), potential IDA interactors such as DNALI1 (top2), DNAH1 (top3), DNAH3 (top4), and DNAH2 (top5) were co-enriched in the candidate interacting dynein protein list (Supplemental Table 5). Immunoblotting confirmed the co-enrichment of DNAH12, DNALI1, and DNAH1 in the immunoprecipitated complex by the DNAH12 rabbit antibody in *Dnah12*^{+/+} mice but not in *Dnah12*^{-/-} mice, indicating specific interactions between DNAH12 and DNALI1, DNAH1 (Fig. 6E). Reciprocal in vivo Co-IP using an anti-DNALI1 antibody further validated strong interactions among these three IDA components (Fig. 6F).

Given the potential role of DNALI1 as an IDA assembly factor (47), we delved into the interaction details between DNAH12 and DNALI1. Employing Phyre2, we constructed the structural models of both proteins and further elucidated their interaction dynamics via DNAH12-DNALI1 docking using the HDock server. The results showed that the N-terminal Stem domain of DNAH12 possibly interacts with DNALI1 in the top three best-fit conformations (Fig. 6G). Considering DNAH12 is encoded by 3092 amino acids and their full-length protein interactions have been proved in vivo, we constructed the Stem domain (1-1212 amino acids) of DNAH12 and full-length DNALI1 plasmid, further in vitro Co-IP assays validated that the Stem domain of DNAH12 indeed mediates interactions with DNALI1 (Fig. 6H-J). Together, these findings proved the specific interactions among DNAH12, DNALI1, and DNAH1. Given the significantly decreased protein levels of DNALI1 and DNAH1 observed in the sperm of *Dnah12*^{-/-} mice (Fig. 5B), we proposed that the interactions involving DNAH12 (possibly restricted to the Stem domain of DNAH12), DNALI1, and DNAH1 are crucial for flagella development. As dynein proteins typically share similar domains, our findings also suggested that the Stem domain of DNAH family members may facilitate the interaction with DNALI1.

Deletion of DNAH12 impairs the recruitment of IDA components during flagella development

To evaluate the effect of DNAH12 deletion on DNALI1 and DNAH1 during spermiogenesis, we examined the abundance of the DNAH12 interaction partners, DNALI1 and DNAH1, in *Dnah12*^{+/+} and *Dnah12*^{-/-} mouse testis lysate. The protein levels of DNALI1 and DNAH1 were decreased in *Dnah12*^{-/-} testes, while the level of ODA member DNAI1 was comparable (Fig. 7A). We then stained DNALI1 or DNAH1 on testicular cell smear of *Dnah12*^{+/+} and *Dnah12*^{-/-} mice. Although the signals of DNALI1 or DNAH1 in round spermatids of *Dnah12*^{-/-} mice were comparable to those of *Dnah12*^{+/+} mice (Fig. 7B and 7C), the recruitment of DNALI1 and DNAH1 to sperm flagella was severely impaired in elongated spermatids of *Dnah12*^{-/-} mice (Fig. 7D and 7E). Considering that the deficiency of DNAH12 also caused scarcely visible immunofluorescence signals of DNALI1 and DNAH1 in sperm flagella of humans and mice (Fig. 3F, 3G, 5E, and 5F), we proposed that the interaction among DNAH12, DNALI1, and DNAH1 is conserved in mammals, and is required for flagellar development during spermiogenesis. Since the deficiency of DNAH12 does

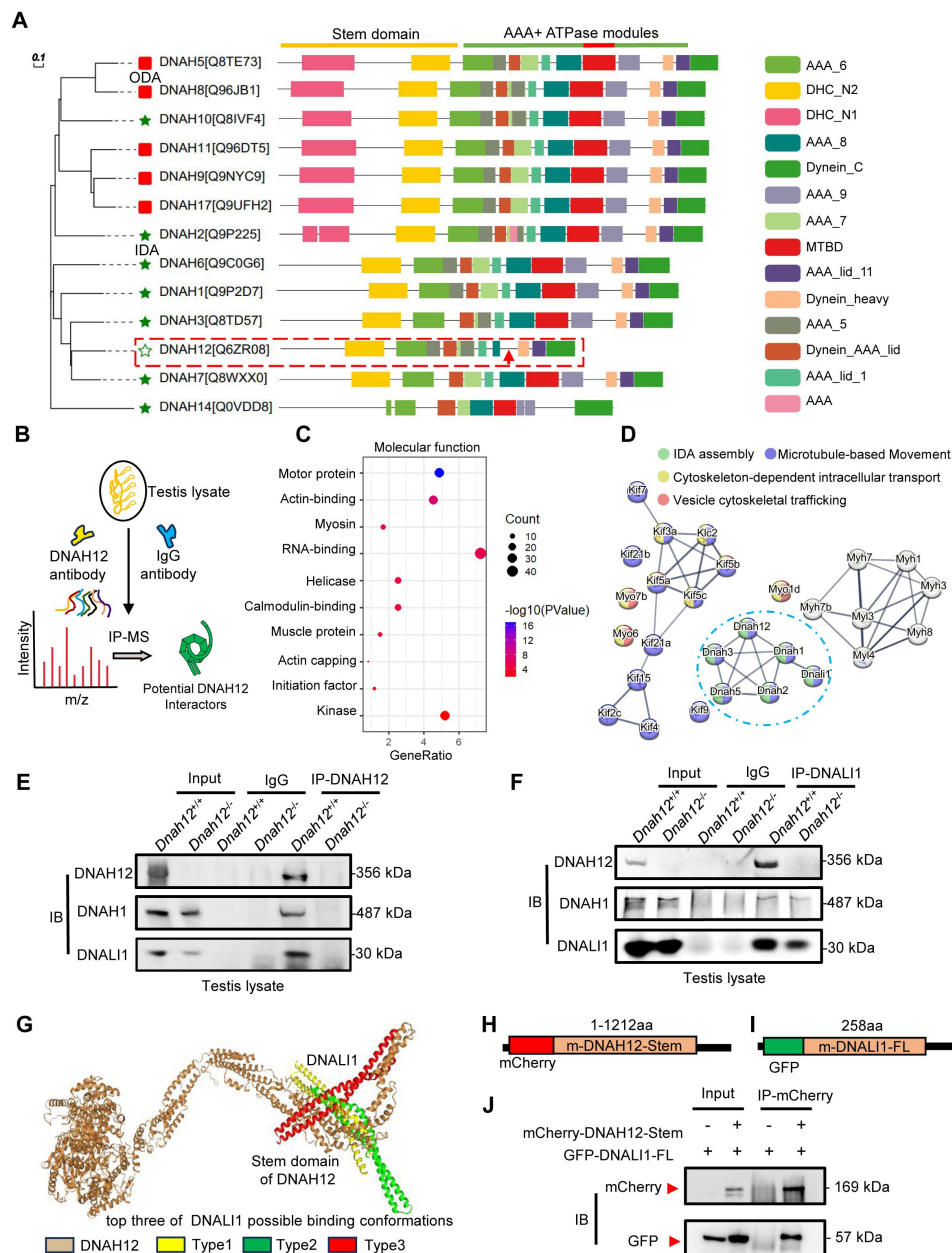


Figure 6.

DNAH12 interacts with IDA components DNALI1 and DNAH1

(A) Phylogenetic and protein domain analyses of the DNAH family. The dashed box indicates the shortest-length member, DNAH12, while the arrow points to the missing microtubule-binding domain (MTBD) region. (B) Schematic diagram showing the procedure of the immunoprecipitation/mass spectrometry (IP/MS) proteomics experiment. (C) GO term enrichment analysis of DNAH12 potential interactors. (D) Protein-protein interaction network of DNAH12 among the motor protein term. The blue circle marks the proteins involved in IDA assembly. (E-F) Co-IP assays using DNAH12 antibody (E) or DNALI1 antibody (F) showed strong and specific interactions among DNAH12, DNALI1, and DNAH1. (G) Predicted DNAH12-DNALI1 interaction details by the HDock server. The top three of DNALI1 possible binding conformations were shown in types 1-3. (H-J) Construction of mouse-mCherry-DNAH12-Stem (H) and mouse-GFP-DNALI1-full length (FL) (I) plasmids, and the validation of the interaction between the Stem domain of DNAH12 and full-length of DNALI1 (J).

not cause PCD symptoms in humans and mice, we hypothesized that flagella and cilia may exhibit distinct features in the process of dynein assembly. We thus checked the abundance of DNALI1 and DNAH1 in the oviduct and trachea, the results showed that DNALI1 and DNAH1 were comparable in the *Dnah12*^{+/+} and *Dnah12*^{-/-} groups (Fig. 7F and 7G). Furthermore, no discernible disparities were observed in the intensity and localization of DNALI1 and DNAH1 signals within the oviduct and trachea (Fig. 7H-K). In summary, these results underscore the specific involvement of DNAH12 in flagellar development, rather than in ciliary development. Moreover, we supposed that the assembly of IDA components differs between flagella and cilia.

DNAH12 is prerequisite for the recruitment of DNALI1 and DNAH1 to sperm flagella

In patients and mice with DNALI1 deficiency, flagellar DNAH1 but not DNAH12 signals were absent (47). To unravel the recruitment relationships among DNAH12, DNALI1, and DNAH1 during sperm flagellar formation, we conducted staining of DNALI1 and DNAH12 on sperm flagella obtained from the infertile asthenoteratozoospermic male P13 with a homozygous *DNAH1* mutation (c.11275C>T) of a family (Fig. 7L, and Supplemental Figure 13 A and B), wherein DNAH1 signals were absent along the flagella (Fig. 7M). In comparison to those in the control, the DNALI1 signals were absent, while no discernible differences in DNAH12 signals were observed in the flagella of the patient with the *DNAH1* mutation (Fig. 7N and 7O). These findings suggested that the recruitment of DNAH12 to flagella is independent of DNAH1, and DNAH12 is the first protein to be recruited to flagella among the three proteins.

DNAH12 deficiency causes the decrease of RS head associated proteins

In addition to the enrichment of dynein proteins, we noticed that a clusters of RS head-related proteins such as RSPH1, RSPH9, or DNAJB13 were enriched in the testicular interactome of DNAH12. Considering that deficiency in RSPH1 (48), RSPH9 (49), or DNAJB13 (50) were associated with CP loss phenotype. We then explored their interaction with DNAH12 in the testis lysate, the results proved the interactions of these proteins with DNAH12 in the testis (Fig. 8A). Further analyses by AlphaFold3 showed that these proteins may interacts with DNAH12 through Stem domain (Fig. 8B). The expression of RSPH1, RSPH9, and DNAJB13 showed obvious decrease in testis lysate of *Dnah12*^{-/-} mice (Fig. 8C). Additionally, immunofluorescence assays revealed the loss of signals for RSPH1, a marker of RS head, in *Dnah12*^{-/-} sperm flagella (Fig. 8D). Together, these results indicate that deficiency of DNAH12 lead to the impairment of radial spoke head, which could contribute to CP loss in flagella of *Dnah12*^{-/-} mice.

ICSI may overcome the infertility caused by DNAH12 deficiency

Since the DNAH12 mutated patients and the *Dnah12*^{-/-} mice have similar deficiency, to evaluate the prospects of whether the asthenoteratozoospermia patients caused by *DNAH12* variants could be treated with assisted reproductive technology, spermatozoa from *Dnah12*^{+/+} and *Dnah12*^{-/-} mice were used for ICSI. Promisingly, the percentages of 2-cell and blastocyst-stage embryos in *Dnah12*^{-/-} group showed no difference compared to those of *Dnah12*^{+/+} group (Fig. 8E-G), indicating that ICSI could also be an effective treatment for infertile men carrying *DNAH12* variants.

Taken together, we suggest that DNAH12 interacts with DNALI1 and DNAH1, and this interaction is a prerequisite for the proper localization of DNALI1 and DNAH1 to sperm flagella, which is essential for the intact IDAs and CP structures in both humans and mice. Additionally, DNALI1 and DNAH1 exhibit mutual dependence for their localization to sperm flagella. Deficiency of DNAH12 leads to the failure of DNALI1 and DNAH1 recruitment to sperm flagella, resulting in abnormal

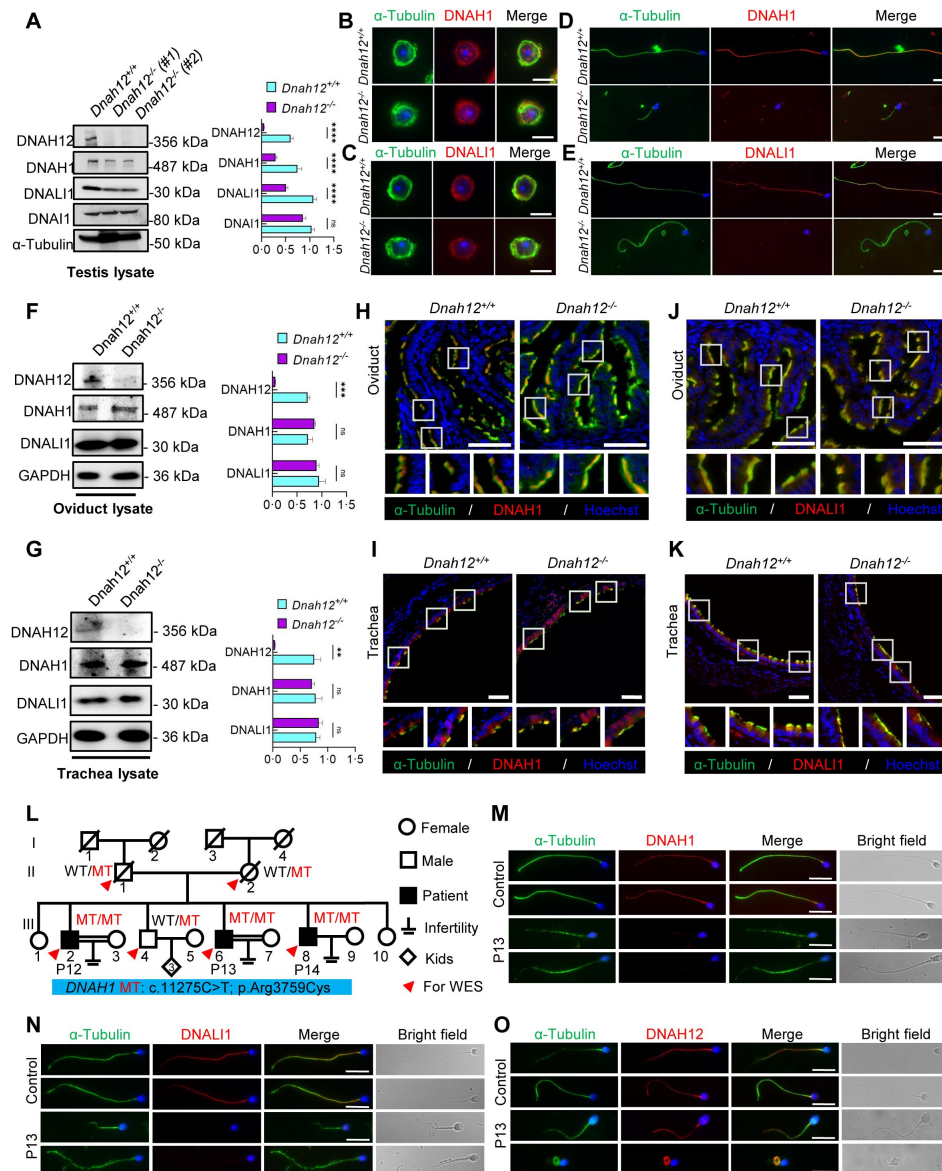


Figure 7.

DNAH12 facilitates the recruitment of DNALI1 and DNAH1 to flagella, but not cilia

(A) Immunoblotting of testis lysate from *Dnah12*^{+/+}, *DNAH12*^{+/-} and *Dnah12*^{-/-} mice using DNAH12, DNAH1, and DNALI1 antibodies. DNAI1 (a component of ODAs) and α -Tubulin were used as the loading controls. 5 replicates of western blotting were performed. The relative band grayscale values of proteins to α -Tubulin in testis lysate from *Dnah12*^{+/+}, and *Dnah12*^{-/-} mice were shown on the right. Data are mean $\pm$ SEM. ns: Not significant; (B-C) Immunofluorescence assays of α -Tubulin and DNAH1 (B), or DNALI1 (C) antibodies on round spermatids from adult *Dnah12*^{+/+} and *Dnah12*^{-/-} mice. Scale bars, 10 μ m. (D-E) Immunofluorescence assays of α -Tubulin and DNAH1 (D), or DNALI1 (E) antibodies on elongated spermatids from adult *Dnah12*^{+/+} and *Dnah12*^{-/-} mice. Scale bars, 10 μ m. (F-G) Immunoblotting of oviduct (F) or trachea (G) lysate from *Dnah12*^{+/+}, *Dnah12*^{+/-} and *Dnah12*^{-/-} mice using DNAH12, DNAH1, and DNALI1 antibodies. GAPDH was used as the loading control. At least 4 replicates of western blotting were performed. The relative band grayscale values of proteins to GAPDH in oviduct lysate (F) or trachea lysate (G) from *Dnah12*^{+/+}, and *Dnah12*^{-/-} mice were shown on the right. Data are mean $\pm$ SEM. ns: not significant; (H-I) Immunofluorescence assays of the oviduct (H) or trachea (I) sections with α -Tubulin and DNAH1 antibodies. Scale bars, 50 μ m. (J-K) Immunofluorescence assays of the oviduct (J) or trachea (K) sections with α -Tubulin and DNALI1 antibodies. Scale bars, 50 μ m. (L) Pedigree of a family with 3 infertile males, P12 (III-2), P13 (III:6), and P14 (III:8); MT: the *DNAH1* mutation c.11275C>T; p.Arg3759Cys. (M-O) Representative images of spermatozoa from a fertile control and P13 co-stained by α -Tubulin antibody and DNAH1 (M), DNALI1 (N), or DNAH12 (O) antibodies, respectively. Scale bars, 10 μ m.

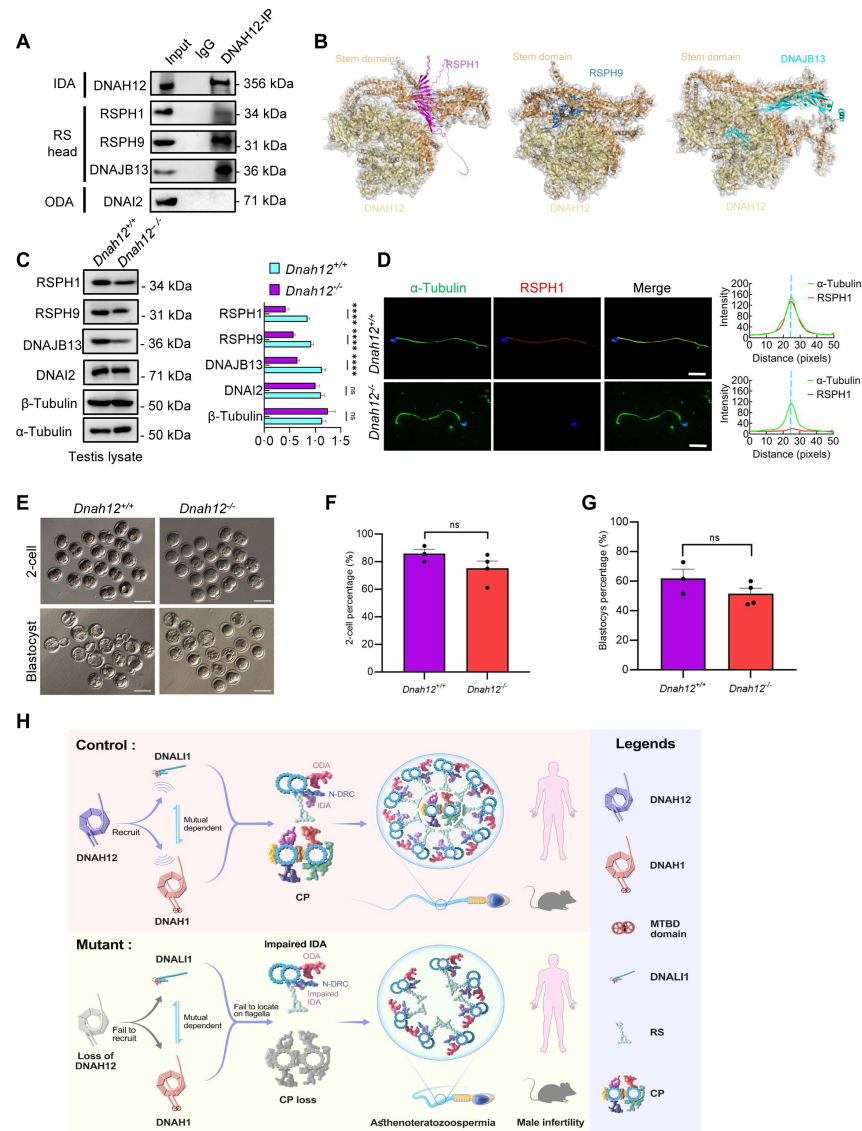


Figure 8.

DNAH12 may interact with RS head associated proteins to regulate CP stability

(A) Co-IP assays of the potential interactions of DNAH12 and RS head proteins RSPH1, RPSH9, or DNAJB13. DNAI2 antibody was used as a negative control. (B) Interaction analyses of DNAH12 and RS head proteins RSPH1, RPSH9, or DNAJB13 by AlphaFold3. (C) Immunoblotting of testis lysate from *Dnah12*^{+/+} and *Dnah12*^{-/-} mice using RSPH1, RPSH9, DNAJB13 antibodies. DNAI2 (a component of ODAs), α -Tubulin and β -Tubulin were used as the loading controls. 4 replicates of western blotting were performed. The relative band grayscale of proteins to α -Tubulin in testis lysate from *Dnah12*^{+/+} and *Dnah12*^{-/-} mice were shown on the right. Data are mean $\pm$ SEM. ns: not significant; (D) Immunofluorescence assays of α -Tubulin and RSPH1 antibodies on sperm collected from *Dnah12*^{+/+} and *Dnah12*^{-/-} mice caudal epididymis. The fluorescent signal intensity profiles were shown on the right. Scale bars, 10 μ m. (E) Representative two-cell embryos and blastocysts of *Dnah12*^{+/+} and *Dnah12*^{-/-} male mice after intracytoplasmic sperm injection. Scale bar, 100 μ m. (F) Percentages of 2-cell-stage embryos of *Dnah12*^{+/+} and *Dnah12*^{-/-} groups. (G) Percentages of blastocyst-stage embryos of *Dnah12*^{+/+} and *Dnah12*^{-/-} groups. (H) Schematic diagram showing the proposed function of DNAH12 in sperm flagellar development in humans and mice. DNAH12 is essential for recruiting DNALI1 and DNAH1 to sperm flagella and maintaining the proper axonemal arrangement, especially IDAs and CP structures. Loss of DNAH12 causes the failure of DNALI1 and DNAH1 to be recruited to sperm flagella and results in abnormal sperm morphology with compromised axoneme organization, causing male infertility. ODA, outer dynein arm; IDA, inner dynein arm; N-DRC, nexin-dynein regulatory complex; CP, central pair of microtubules; MTBD, microtubule-binding domain; RS, radial spoke.

sperm morphology characterized by compromised axoneme organization, causing male infertility (Fig. 8H). Moreover, our results indicate that the assembly of IDA components occurs intricately, possibly not synchronously in time and space.

Discussion

In this study, we identified six novel bi-allelic mutations in *DNAH12* that cause the loss of DNAH12 proteins in asthenoteratozoospermia patients. Spermatozoa from these patients exhibited abnormal morphology and compromised axonemal defects, including a high percentage of impaired IDAs and loss of CP. Two mouse models (*Dnah12*^{-/-} and *Dnah12*^{mut/mut}) were generated, recapitulating the abnormal flagellum morphology and structure observed in patients, thereby confirming the pathogenic role of *DNAH12* mutations in asthenoteratozoospermia. Through testicular Co-IP and mass spectrometry proteomics, we screened and validated that DNAH12 interacts with IDA components like DNALI1 and DNAH1 and is indispensable for recruiting DNALI1 and DNAH1 to sperm flagella, highlighting the crucial role of DNAH12 in sperm flagellar development. Notably, deficiency of DNAH12 does not manifest PCD symptoms in humans and mice, underscoring the unique role of DNAH12 in flagellar development. Collectively, our study not only unveils the functional significance of the previously unexplored gene *DNAH12* but also elucidates the pathogenic link between *DNAH12* mutations and asthenoteratozoospermia.

DNAH12 and other dynein arm proteins share the N-terminal Stem domain, which is crucial for structure and accessory subunit binding (51). The motor domain which encompasses ~6 AAA+ ATPase modules forming the ring-shaped oligomeric complexes responsible for force production (52). Five of the identified *DNAH12* mutations, primarily in the Stem domain (critical for DNALI1 interaction), result in loss-of-function effects, indicating a weak tolerance property of the Stem domain. Similar weak tolerance features were observed in mutations of other dynein arm components like *DNAH1* and *DNAH2* (53, 54). The profound impact of these variants, particularly missense mutations, underscores delicate features of the Stem domain and DNAH family members. Notably, DNAH12 stands out as the shortest in sequence length and the only one lacking the essential MTBD region within the DNAH family (55). MTBD, a 15-nanometer coiled-coil stalk structure, crucially supports ATP-sensitive microtubule binding properties and determines dynein motility direction (56). Our investigation revealed that DNAH12 interacts with other IDA components, specifically DNALI1 and DNAH1. Despite the mutual dependence between DNAH1 and DNALI1 for their flagellar localization, DNAH12 is prerequisite for the proper assembly of DNALI1 and DNAH1 onto sperm flagella in both humans and mice. Notably, *Dnah12*^{-/-} mice exhibit more severe phenotypes than *Dnali1* and *Dnah1* knockout mice (4, 47, 57), including a more severe reduced sperm count (~ 2% of normal values), this observation further supports our perspective.

Deficiency in DNAH12 presented sperm ultrastructural defects like CP loss and impaired IDAs indicating essential roles of DNAH12 in the maintenance of axonemal integrity and arrangement. These results indicate essential roles of DNAH12 in the maintenance of axonemal integrity and arrangement. Noteworthy, deficiency of DNAH12 potential partner DNAH1, a well-known asthenoteratozoospermia-related gene, presents quite similar defects with DNAH12, indicating DNAH12 and DNAH1 possibly exhibit synergic functions in axonemal proper arrangement (58–60). In addition to DNAH12 and DNAH1, deficiency in DNAH2 or DNAH6 also leads to high proportions of CP loss and disorganized axonemal structures (5, 54, 61). However, DNALI1, DNAH7, or DNAH10 deficiency results in no obvious CP loss but impaired IDA integrity (7, 8, 22, 47). These studies indicate that specific IDA components, like DNAH12 and DNAH1, are required for the maintenance of CP integrity. Moreover, discrepancies observed in the phenotypes resulting from deficiencies in specific IDAs suggest that multiple dynein proteins are produced to support the intricate organization of sperm flagella. Additionally, these proteins may exhibit distinct characteristics and perform unique functions to some extent. Noteworthy, deficiency in RS

head proteins RSPH1, RSPH9, or DNAJB13 were related to axonemal CP loss phenotype (48 [↗](#), 50 [↗](#), 62 [↗](#)), indicating that DNAH12, and other IDAs, possibly regulate the stability of CP through the cooperation with RS head proteins. Notably, while the adoption of AlphaFold3 and HDock contributes to a clearer model of protein structures and their interactions, further techniques like cryo-EM may facilitate a deeper understanding of these proteins at higher resolution. Elucidating the dependencies among the assembly of sperm components could enhance our understanding of the structural basis of sperm motility and human fertility. Additionally, while an enrichment of IDA and RS proteins, but not nexin-dynein regulatory complex (N-DRC) proteins, was observed in the DNAH12 interactome, further exploration is needed to determine whether DNAH12 may indirectly affect the stability and functionality of the N-DRC structure.

The localization of DNAH12 presents dynamic changes during spermiogenesis. DNAH12 is initially expressed in the cytoplasm of the round spermatid, while it's enriched in the manchette structure of the elongating spermatid and the tail of the elongated spermatid. The manchette and the sperm tail axoneme are two microtubular platforms that transport proteins to the developing head and tail (63 [↗](#)). Manchette is a transient skirt-like structure and functions as a scaffold for morphological remodeling, contributing to the formation of the sperm head, neck, and tail (64 [↗](#)). Here, we found that DNAH12 deficiency results in abnormal manchette structure and compromised sperm head and tail morphology. Additionally, in sperm of *Dnah12*^{-/-} males, several CP complex proteins like SPAG6, and SPEF2 are not properly assembled and the axoneme structure of the tail was severely disorganized, indicating dynein components are responsible for proper axonemal components organization. Noteworthy, DNALI1, a potential DNAH12 interacting partner showed similar localization patterns to DNAH12, a study presented that DNALI1 recruits and stabilizes manchette component PACRG, and DNALI1 deficiency causes impaired sperm flagellar assembly (57 [↗](#)). These data hint that dynein proteins like DNAH12 and DNALI1, possibly work together in the manchette organization as well as sperm flagellar components assembly during sperm development.

DNAH12 was also detected in the lungs, tracheas, and female oviducts while no cilia morphology defects in these tissues or PCD symptoms were observed in both two mice models. Besides, none of the patients carrying bi-allelic *DNAH12* mutations claimed symptoms of PCD via visits except for the reproductive problems. *Dnah12*^{-/-} female mice showed normal development and fertility and the mother (III:2) of patients from family 2, carrying homozygous *DNAH12* mutation M2 had 6 offspring. Intriguingly, we found that the localization and abundance of DNALI1 and DNAH1 in oviductal or tracheal cilia were not affected after *Dnah12* deletion, indicating that DNAH12 works differently in flagella and cilia. Deficiency of specific DNAH members like DNAH6 (65 [↗](#)), or DNAH7 (22 [↗](#)), but not DNAH2 (54 [↗](#)), results in PCD symptoms. Hence, DNAH family members possibly present disparate processing of axonemal biogenesis and organization through evolution in flagella and cilia, further explorations could contribute to a more comprehensive understanding of the differences between flagella and cilia, and personalized clinical diagnosis of related diseases like male/female infertility and/or PCD (66 [↗](#)).

Previously, two genetic screening studies mentioned unverified compound heterozygous mutations and a homozygous missense mutation in *DNAH12*, predicting their association with patients' infertility. However, limited genetic evidence was provided, and the pathogenesis remained unclear (26 [↗](#), 27 [↗](#)). Through investigations involving patients with *DNAH12* variants and mutant mouse models, we provide direct genetic and molecular evidence that variants in *DNAH12* are genetic causes of asthenoteratozoospermia, and DNAH12 along with DNAH1, DNALI1 is indispensable for sperm flagellar biogenesis. These data strongly support DNAH12 as a valuable marker in genetic counseling and diagnosis of male infertility.

ICSI is a widely used assisted reproductive technology in treating infertile men with asthenoteratozoospermia. Deficiency in most dynein proteins leads to asthenoteratozoospermia with severely compromised sperm motility and morphology. Previous studies have shown that

patients carrying mutations in *DNAH17* (10 [DOI](#)) have poor ICSI outcomes while patients with *DNAH1* (59 [DOI](#)), *DNAH2* (5 [DOI](#), 61 [DOI](#)), *DNAH3* (67 [DOI](#)), *DNAH7* (22 [DOI](#)), or *DNAH8* (68 [DOI](#)) mutations have achieved favorable clinical pregnancy. Thus, the comparative study of asthenoteratozoospermia-related genes on their ICSI outcomes will be beneficial to developing individualized treatment recommendations in clinical practice. In our study, the successful ICSI outcomes of *Dnah12*^{-/-} mice suggest that ICSI could be used as a potential treatment for infertile men carrying *Dnah12* mutations. It's meaningful to evaluate the prognosis for ICSI treatments in patients carrying *DNAH12* variants in a large population.

In conclusion, our study functionally and mechanistically sheds light on a special and poorly characterized protein DNAH12, during the intricate process of sperm flagellar biogenesis. These results pave the way for an enhanced understanding of the roles of dynein proteins in sperm flagellar development and their functional difference in flagella and cilia, offering a diagnostic target for genetic counseling and future individualized treatment of male infertility in clinical practice.

Data availability

All data that support the findings in this study are available from the corresponding author upon reasonable request. Proteomics data supporting this study have been deposited to the ProteomeXchange Consortium via the iProX partner repository with the dataset identifier PXD051681.

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Additional information

Author contributions

B.S. and Q.S. conceived and supervised the study; M.Y., X.H., J.Y., M.C., A.Z., T.L., A.Z., and N.A. performed the experiments and analyzed the data; H.M.J., M.K., Z.M., R.K., A.A., and W.S. collected the patients' samples; J.Z. and H.Z. performed the WES sequencing and data analyses; M.Y. wrote the manuscript and B.S., Q.S., H.M., and M.Y. modified the manuscript; B.S., Q.S., H.M., B.X., H.Z., and A.M. were involved in discussions of the manuscript.

Ethics statement

This work was approved by the institutional ethics committee of University of Science and Technology of China (USTC) with the approval number 2019-KY-168, and informed consent was obtained from patients and controls. All animal experiments were approved by the Institutional Animal Care and Use Committee of the USTC with the approval number USTCACUC1301021.

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Additional files

Supplemental Fig 1-13 [🔗](#)

Supplemental video 1-2.zip [🔗](#)

Supplemental video 3-4.zip [🔗](#)

Supplemental Tables 1-5 [🔗](#)

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Reviewer #1 (Public review):

Summary:

Even though this is not the first report that the mutation in the DNAH12 gene causes asthenoteratozoospermia, the current study explores the sperm phenotype in-depth. The authors show experimentally that the said mutation disrupts the proper axonemal arrangement and recruitment of DNALI1 and DNAH1 - proteins of inner dynein arms. Based on these results, the authors propose a functional model of DNAH12 in proper axonemal development. Lastly, the authors demonstrate that the male infertility caused by the studied mutation can be rescued by ICSI treatment at least in the mouse. This study furthers our understanding of male infertility caused by a mutation of axonemal protein DNAH12, and how this type of infertility can be overcome using assisted reproductive therapy.

Strengths:

This is an in-depth functional study, employing multiple, complementary methodologies to support the proposed working model.

Weaknesses:

The structure and interaction model between DNAH12, DNALI1, and DNAH1 relies on *in silico* methodologies, and further studies are required to validate these predictions.

<https://doi.org/10.7554/eLife.100350.2.sa2>

Reviewer #2 (Public review):

Summary:

The authors first conducted whole exome sequencing for infertile male patients and families where they co-segregated the biallelic mutations in the Dynein Axonemal Heavy Chain 12 (DNAH12) gene. Sperm from patients with biallelic DNAH12 mutations exhibited a wide range of morphological abnormalities in both tails and heads, reminiscing a prevalent cause of male infertility, asthenoteratozoospermia. To deepen the mechanistic understanding of DNAH12 in axonemal assembly, the authors generated two distinct DNAH12 knockout mouse lines via CRISPR/Cas9, both of which showed more severe phenotypes than observed in patients. Ultrastructural observations and biochemical studies revealed the requirement of DNAH12 in recruiting other axonemal proteins and that the lack of DNAH12 leads to the

aberrant stretching in the manchette structure as early as stage XI-XII. At last, the authors proposed intracytoplasmic sperm injection as a potential measure to rescue patients with DNAH12 mutations, where the knockout sperm culminated in the blastocyst formation with a comparable ratio to that in WT.

Strengths:

The authors convincingly showed the importance of DNAH12 in assembling cilia and flagella in both human and mouse sperm. This study is not a mere enumeration of the phenotypes, but a strong substantiation of DNAH12's essentiality in spermiogenesis, especially in axonemal assembly.

The analyses conducted include basic sperm characterizations (concentration, motility), detailed morphological observations in both testes and sperm (electron microscopy, immunostaining, histology), and biochemical studies (co-immunoprecipitation, mass-spec, computational prediction). Molecular characterizations employing knockout animals and recombinant proteins beautifully proved the interactions with other axonemal proteins.

Many proteins participate in properly organizing flagella, but the exact understanding of the coordination is still far from conclusive. The present study gives the starting point to untangle the direct relationships and order of manifestation of those players underpinning spermatogenesis. Furthermore, comparing flagella and trachea provides a unique perspective that attracts evolutionary perspectives.

Weaknesses:

Seemingly minor, but the discrepancies found in patients and genetically modified animals were not fully explained. For example, both knockout mice vastly reduced the count of sperm in the epididymis and the motility, while phenotypes in patients were rather milder. Addressing the differences in the roles that the orthologs play in spermatogenesis would deepen the comprehensive understanding of axonemal assembly.

Comments on revisions:

The reviewer is satisfied with the authors' response.

<https://doi.org/10.7554/eLife.100350.2.sa1>

Author response:

The following is the authors' response to the original reviews.

eLife Assessment

This important study further validates DNAH12 as a causative gene for asthenoteratozoospermia and male infertility in humans and mice. The data supporting the notion that DNAH12 is required for proper axonemal development are generally convincing, although more experiments would solidify the conclusions. This work will interest reproductive biologists working on spermatogenesis and sperm biology, as well as andrologists working on male fertility.

We thank the editor and the two reviewers for their time and careful evaluation of our manuscript. We sincerely appreciate their encouraging feedback and insightful guidance on improving our study. In the revised manuscript, we have performed additional experiments and provided quantitative data regarding the reviewers' comments.

Public Reviews:

Reviewer #1 (Public Review):

Summary:

Even though this is not the first report that the mutation in the DNAH12 gene causes asthenoteratozoospermia, the current study explores the sperm phenotype in-depth. The authors show experimentally that the said mutation disrupts the proper axonemal arrangement and recruitment of DNALI1 and DNAH1 - proteins of inner dynein arms. Based on these results, the authors propose a functional model of DNAH12 in proper axonemal development. Lastly, the authors demonstrate that the male infertility caused by the studies mutation can be rescued by ICSI treatment at least in the mouse. This study furthers our understanding of male infertility caused by a mutation of axonemal protein DNAH12, and how this type of infertility can be overcome using assisted reproductive therapy.

Strengths:

This is an in-depth functional study, employing multiple, complementary methodologies to support the proposed working model.

Thank you for your recognition of the strength of this study. Your positive feedback motivates us to continue refining our research and methodological rigor in future studies.

Weaknesses:

The study strength could be increased by including more controls such as peptide blocking of the inhouse raised mouse and rat DNAH12 antibodies, and mass spectrometry of control IP with beads/IgG only to exclude non-specific binding. Objective quantifications of immunofluorescence images and WB seem to be missing. At least three technical replicates of western blotting of sperm and testis extracts could have been performed to demonstrate that the decrease of the signal intensity between WT and mutant was not caused by a methodological artifact.

Thank you for your comments. In order to study in-depth, we have analyzed the protein sequence features of DNAH12 protein, 1-200 amino acids of DNAH12 were selected as the ideal antigen considering its good performance (1. high immunogenicity; 2. High hydrophilicity; 3. Good Surface Leakage Groups; 4. Sequence homology analysis to avoid unspecific recognition to other proteins;). The two different anti-DNAH12 antibodies were developed with the help Dia-An Biotech company in 2022, we have tried to acquire the polypeptide fragments of target proteins to do peptide blocking but the material were discard after the service. Luckily, we have got the target band of DNAH12 protein in western blotting experiment while the band was not detected in knockout mice group; the immunofluorescence signals of DNAH12 were strong but not present in knockout mice group. Besides, we have tested that the inhouse raised rabbit antibody were suitable for IP experiment. The IP experiment also showed the raised rabbit antibody were able to immunoprecipitated the DNAH12 band in the *Dnah12*^{+/+} mice but not in *Dnah12*^{-/-} mice. Collectively, these data could support the specificity of the raised DNAH12 antibodies. In IP assay, we have added the IgG group in the IP-mass spectrometry to exclude non-specific binding. And the experimental design was described in Figure 6B. The raw data were deposited in iProX partner repository (accession number: PXD051681), and we have coordinated with the repository manager to make the data publicly accessible (<https://www.iprox.cn/page/subproject.html?id=IPX0008674001>).

Besides, we have conducted replicates of western blotting of sperm and testis extracts at least 3 times and added the objective quantifications of immunofluorescence signals and WB images. The quantifications of the blot were shown in figures to help readers understand these results easily.

Reviewer #2 (Public Review):

Summary:

The authors first conducted whole exome sequencing for infertile male patients and families where they co-segregated the biallelic mutations in the Dynein Axonemal Heavy Chain 12 (DNAH12) gene.

Sperm from patients with biallelic DNAH12 mutations exhibited a wide range of morphological abnormalities in both tails and heads, reminiscing a prevalent cause of male infertility, asthenoteratozoospermia. To deepen the mechanistic understanding of DNAH12 in axonemal assembly, the authors generated two distinct DNAH12 knockout mouse lines via CRISPR/Cas9, both of which showed more severe phenotypes than observed in patients. Ultrastructural observations and biochemical studies revealed the requirement of DNAH12 in recruiting other axonemal proteins and that the lack of DNAH12 leads to the aberrant stretching in the manchette structure as early as stage XI-XII. At last, the authors proposed intracytoplasmic sperm injection as a potential measure to rescue patients with DNAH12 mutations, where the knockout sperm culminated in the blastocyst formation with a comparable ratio to that in WT.

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The authors convincingly showed the importance of DNAH12 in assembling cilia and flagella in both human and mouse sperm. This study is not a mere enumeration of the phenotypes, but a strong substantiation of DNAH12's essentiality in spermiogenesis, especially in axonemal assembly.

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Many proteins participate in properly organizing flagella, but the exact understanding of the coordination is still far from conclusive. The present study gives the starting point to untangle the direct relationships and order of manifestation of those players underpinning spermatogenesis. Furthermore, comparing flagella and trachea provides a unique perspective that attracts evolutionary perspectives.

Thank you for your thoughtful and positive feedback. We are delighted that you found our study to be a strong substantiation of DNAH12's essential role in spermiogenesis, particularly in axonemal assembly. We believe that this study represents a meaningful step toward unraveling the intricate coordination of axonemal proteins during spermatogenesis, and your comments further inspire us to continue exploring these complex mechanisms in future work. Thank you once again for your valuable insights and summary of this work.

Weaknesses:

Seemingly minor, but the discrepancies found in patients and genetically modified animals were not fully explained. For example, both knockout mice vastly reduced the

count of sperm in the epididymis and the motility, while phenotypes in patients were rather milder. Addressing the differences in the roles that the orthologs play in spermatogenesis would deepen the comprehensive understanding of axonemal assembly.

This is an interesting question. Actually, it seems that although humans and mice share the male infertility phenotypes with deficiency in dynein proteins essential for sperm flagellar development, they are different in some ways. For instance, it has been reported that deficiency in DNAH17 (Clin Genet. 2021. PMID: 33070343) or DNAH8 (Am J Hum Genet. 2020. PMID: 32619401; PMCID: PMC7413861), two other members of Dynein Axonemal Heavy Chain family, also cause more severe phenotype in mice, comparing with that of human patients carrying bi-allelic DNAH17 or DNAH8 loss-of-function mutations. In knockout mice, sperm counts are lower, and the proportion of abnormal sperm morphology is higher, whereas the phenotypes in human patients tend to be milder. These observations suggest that orthologs may influence spermatogenesis to slightly different extents in humans and mice. We plan to investigate the mechanisms underlying these discrepancies in future studies, which will provide deeper insights into axonemal assembly and the evolutionary aspects of spermatogenesis. Thank you again for bringing up this important issue.

Recommendations for the authors:

Reviewer #1 (Recommendations For The Authors):

This reviewer is impressed by the study's depth and the extent of the methodology used in the study. The study is well-designed, and the results are very interesting. The reviewer's enthusiasm was reduced by the lack of some controls (provided that the reviewer did not miss them). Further are point-to-point suggestions that this reviewer believes will increase the merit of the present study.

Title:

(1) Why a "special" dynein? What makes it special when compared to other dyneins? I suggest removing the word special.

Through phylogenetic and protein domain analyses of the DNAH family, we found that DNAH12 is the shortest member and the only one that lacks a typical microtubule-binding domain (MTBD) in the DNAH family, thus we want to describe it as a "special" dynein. We have fully considered your valuable suggestion and decided to remove it from the title.

Abstract:

(2) L23: same as above, why special?

We identified DNAH12 as the shortest member of the DNAH family and uniquely lacking the typical microtubule-binding domain (MTBD). This distinct characteristic prompted us to describe it as a 'special' dynein in the abstract part.

(3) L37: the reviewer did not find a figure (neither main nor supplementary) that would demonstrate the proper organization of microtubules in cilia. Figure S11 only shows the presence of cilia in DNAH12^{-/-} mouse. A TEM image of cilia is required to confirm or reject the claim that DNAH12 does not play a crucial role in proper microtubule organization in cilia.

We have now added TEM images of cilia in wild-type and *Dnah12^{-/-}* mice. The ultra-structures of cilia axonemes were comparable in wild-type and *Dnah12^{-/-}* groups, suggesting that

DNAH12 may not play crucial role in proper microtubule organization. The results have now been added to Supplemental Figure 11F.

(4) L122-6: Did the authors also confirm these structures by cryo-EM? If not, this needs to be pointed out as a shortcoming in the discussion, that the structures and interactions are predicted in silico only.

Thank you for your comment. Due to resource limit, we do not perform cryo-EM to confirm these structures. We will pursue the structures details at an atomic resolution structure in further study. We understand this point and now we have addressed this as a shortcoming in the discussion part.

(5) L134: Be more specific about what characteristics of DNAH12 were analyzed.

Thank you for your comment. We have now updated these in the method part. The characteristics of the DNAH12 including its region immunogenicity, hydrophilicity, surface leakage groups, and sequence homology were analyzed.

(6) L137: Be more specific about how the antibodies validated were. Were the antibodies validated for both immunofluorescence and western blotting? I suggest doing peptide blocking of the antibody, for instance for ICC, preincubation of ab with immunizing peptide followed by primary ab incubation with studied cells/tissues.

Thank you for your comments and suggestions. We validated the antibodies for both immunofluorescence and western blotting to ensure their effectiveness in our experiments. The two different anti-DNAH12 antibodies were developed with the help Dia-An Biotech company in 2022, we have attempted to acquire the polypeptide fragments of target proteins to do peptide blocking but the material were disposed after the service. Luckily, we have got the target band of DNAH12 protein which showed strong signal in western blotting experiment and the band was not detected in knockout mice group; the immunofluorescence signals of DNAH12 were strong but not present in knockout mice group. Besides, the IP experiment also showed the raised rabbit antibody were able to immunoprecipitated the DNAH12 band in the *Dnah12*^{+/+} mice but not in *Dnah12*^{-/-} mice. Collectively, these data could support the specificity of the raised DNAH12 antibodies. We sincerely admire your suggestion and will require for the peptide material if we develop new antibodies.

(7) L142: This reviewer is unfamiliar with using TRIzol for sperm protein extraction. Is there a specific reason for not using PAGE loading buffer for human sperm protein extraction?

Thanks for your suggestions. TRIzol reagent can be used for small amounts of samples (5×10^6 cells) as well as large amounts of samples ($>10^7$ cells). It is suitable for extraction of RNA and proteins at the same time. Our lab has adopted these methods in our previous work (Hum Reprod Open. 2023; PMID: 37325547; PMCID: PMC10266965.). This method is very useful to process valuable small amounts of samples for scientific work. The human sperm protein extraction was added with SDS-sample buffer [PAGE loading buffer] before SDS-PAGE separation. We have added this detail in the method part. We are sorry for making this misunderstanding.

(8) L144: Were these the final concentrations of the SDS loading buffer? 1 × Laemmli buffer contains 62.5 mM TRIS, 2% (w/w) SDS, 10 % (w/v) glycerol, and 5% 2-mercaptoethanol. Please, amend accordingly.

Thanks for your suggestions. We apologized for incorrect labelling of concentrations (The previous one is 3× SDS loading buffer). We have now amended the SDS loading buffer to 1 × Laemmli buffer as suggested.

(9) L151: Table S2 contains other homemade antibodies than DNAH12. Please, include references to the studies where the generation and validation of these antibodies is described.

Thank you for your suggestions. We have developed a DNAH1 antibody for use in Western blot assays, with its generation and validation detailed in *Frontiers in Endocrinology* (Lausanne), 2021 (PMID: 34867808; PMCID: PMC8635859). Additionally, we have produced a DNAH17 antibody for both immunofluorescence (IF) and Western blot, as described in *Journal of Experimental Medicine*, 2020 (PMID: 31658987; PMCID: PMC7041708). These references have now been included.

(10) L167: Please, spell out ICR at its first appearance.

Done as suggested, Thank you. The full name of ICR is Institute of Cancer Research.

(11)L169: This reviewer is confused. It seems that the mouse encodes DNAH12 on exons 5 and 18 simultaneously. Each mouse model has only one exon targeted for a knockout. Would not this mean that the expression of DNAH12 in both models is not completely knocked down? Please, give more background in this paragraph for those less familiar with CRISPR/Cas9.

Thank you for your insightful comment. We appreciate your attention to detail. To clarify, while the mouse model does indeed encode DNAH12 on exons 5 and 18 simultaneously, we specifically targeted the key exon 5 or exon 18 in each model to achieve different knockout strategies. This approach allows us to assess the functional implications of the remaining DNAH12 expression in both models. We have checked the DNAH12 expression in both models, and the result showed both models present with undetected DNAH12 proteins, indicating both models were completely knocked out of DNAH12 proteins. Additionally, we will revise the manuscript to include further details on the CRISPR/Cas9 methodology, ensuring accessibility for readers less familiar with this technique. Thank you again for your valuable feedback, which we believe will greatly enhance our manuscript.

(12) L201: 50 % PBS? As in 0.5 x concentrated PBS? Please, rewrite for clarity.

The term "50% PBS" refers to a 1:1 dilution of phosphate-buffered saline (PBS) with an appropriate diluent, resulting in a final concentration of 0.5x PBS. We will revise the text to explicitly clarify this, ensuring it is clear to all readers. Thank you for highlighting this point.

(13) L224: Please, state what beads those were (magnetic/agarose, conjugated to protein A/G...) Include catalog # and manufacturer.

Thank you for your suggestion. We have updated the manuscript to include this information. The beads used were Protein A/G Magnetic Beads (Catalog #B23202, Bimake, Texas, USA).

(14) L227: What was the reason for adding a proteasomal inhibitor? What concentration was used? Please, add this information to the text.

We adding MG132 in cell immunoprecipitation (IP) experiments is to inhibit proteasomal activity, thereby preventing the degradation of the target protein. This helps maintain the

stability of the target protein during the experiment (Sci Adv. 2022. PMID: 35020426; PMCID: PMC8754306.), enhancing its detectability in subsequent analyses. MG132 (5 μ M) was added. We have added this information in the revised the manuscript

(15) L233: *in vivo IP of mouse testis lysate? This does not make sense. I suggest removing "in vivo".*

Thank you for your careful review and comments on our manuscript. We have modified as suggested.

(16) L317: *Supplemental Figure 6 precedes Supplemental Figure 5 in the text, which is neither logical nor orderly.*

Thank you for your suggestion. Since the N-terminal DNAH12 antibody is already described in the Methods section (L317), we propose removing Supplemental Figure 6 from the content to improve the logical flow and maintain an orderly presentation.

(17) L345 and elsewhere: *how did the authors quantify the decrement of the signal? This needs to be measured objectively.*

Thank you for your valuable suggestion. We quantified the signal intensity using Fiji (Nat Methods. 2012. PMID: 22743772; PMCID: PMC3855844), which allows for precise analysis of pixel intensity. The results are presented in the figures to effectively illustrate the decrement in signal intensity. We appreciate your suggestion, and we have provided a description of the method in our methodology section.

(18) L371: *I recommend: ...and elongated spermatids; the abnormal...*

Done as suggested. Thank you.

(19) L412-4: *Cilia in both $Dnah12^{mut/mut}$ and $Dnah12^{-/-}$ are developed, but are they motile or immotile? This needs to be investigated. Is the DNAH12 in cilia truncated while still fulfilling its function?*

Thanks for your comment. We have checked the ciliary motility using an inverted microscope, and no significant difference of ciliary motility were observed between the knockout group and the control group. These results indicated that the ciliary motility was not affected by DNAH12 deficiency. The N-terminal DNAH12 antibody was developed to detect whether a truncated protein in mice tissues while we do not detect DNAH12 signals through immunofluorescence assay on trachea sections of the $Dnah12^{-/-}$ mice. These results indicate that DNAH12 may exert little influence on cilia, comparing to its important function in flagella.

(20) L414-6: *The results do not support this claim as the authors do not show that cilia are motile.*

Thanks for your comment. The supplemental videos 3-4 of trachea live of $Dnah12^{+/+}$ and $Dnah12^{-/-}$ mice have been uploaded to support this conclusion.

(21) L421-3: *Did the authors perform a negative test, where they let the testis lysate interact with beads/IgG only and performed the MS to identify non-specific binding? This is a crucial specificity test for this approach.*

We have performed negative test. In IP assay, we have added the IgG group in the IP-mass spectrometry to exclude non-specific binding. And the experimental design was described in Figure 6B. The raw data were deposited in iProX partner repository (PXD051681), which we have required the manager soon to update the status to public, so it will be visible to readers.

(22) L462: same as #18 the authors need to show that cilia are also motile. The mere presence of cilia in *DNAH12*^{-/-} as shown in Fig S11C&D is not sufficient to conclude that the mice do not manifest PCD symptoms.

Thanks for your comment. We do not observe obvious differences between the cilia of *Dnah12*^{+/+} and *Dnah12*^{-/-} mice. The supplemental videos 3-4 of trachea live of *Dnah12*^{+/+} and *Dnah12*^{-/-} mice have been uploaded to show the motility of the trachea.

(23) L529: MTBD region instead of domain, as "domain" is already part of the abbreviation.

Done as suggested

(24) L875: Sperm is both the singular and plural form. Spermatozoon vs spermatozoa can be used where the distinction between singular and plural needs to be made.

Thanks for your suggestion. We have checked and changed this usage.

(25) Figure 3H: Is there a specific reason why P11 is not shown?

Because limited smear slides of P11 were available, the P11 were not stained for DNAH17 antibody previously. We have now updated the experiment, which showed that DNAH17 expression were not affected in patient P11. We have now added this result to Figure 3H.

(26) Figure 8H: The authors in their MS do not describe what is happening to N-DRC proteins, yet they suggest in their model that it's unaffected in the mutant mouse/human. Please, address this in the MS and clearly state in the model that N-DRC needs further exploration in future studies.

Thanks for your suggestion, we have checked the MS data but do not observe the enrichment of nexin-dynein regulatory complex (N-DRC) protein, just one known N-DRC protein DRC1 present with only 1 unique peptide. Instead, enrichment of inner dynein arm proteins and radial spoke proteins were observed. However, we cannot determine the N-DRC structures maybe affected or not. We have stated this in the discussion part and will pursue this with high resolution technology like cryo-EM in the future.

(27) Figure 5F: Is it possible to choose a different *Dnah12*^{-/-} spermatozoon to see a reduced level of DNALI1 so that it corresponds with the WB detection in Fig 5B?

Thanks for your suggestion, we have chosen a *Dnah12*^{-/-} spermatozoon with faint remnants of the DNALI1 signal as the representative picture.

(28) Figure S2 and elsewhere: How were the authors able to resolve and calibrate 356 kDa protein using SDS PAGE? Agarose electrophoresis protein electrophoresis is more suitable for resolution of high molecular proteins. Most of the protein standards have as high molecular standard as 250 kDa.

We have found that high molecular proteins (like 356kDa) were able to resolve in concentration 4-12% gradient gel of polyacrylamide gels and employ appropriate voltages and more time during electrophoresis to improve resolution of high molecular weight proteins. The DNAH12 proteins were calibrated by the using of a HiMark™ Pre-Stained High Molecular Weight Protein Standard (30-460 kDa). We have now updated the blot images to show the size of the DNAH12 protein (Fig S6B,). The target band is obvious between 268 kDa and 460 kDa, which make it easy to calculate the target band of DNAH12 antibody elsewhere. Thanks for your suggestion.

(29) Figure S5: similar to #24: Why P10 and P11 are not shown?

Because limited smear slides of P10 or P11 were available, we did not stain ODF2 antibody previously. We have now updated the experiments, which showed that ODF2 expression were not affected in patient P10 or P11. We have now added this result to Figure S5.

(30) Figure S6B: The specificity of the anti-DNAH12 antibody against mouse DNAH12 seems to be questionable since the authors detect multiple bands on WB. I recommend doing peptide blocking to show that these are non-specific binding as opposed to off-target binding.

Thank you for your comments. In order to study in-depth, we have analyzed the protein sequence features of DNAH12 protein, 1-200 amino acids of DNAH12 were selected as the ideal antigen considering its good performance (1. high immunogenicity; 2. High hydrophilicity; 3. Good Surface Leakage Groups; 4. Sequence homology analysis to avoid unspecific recognition to other proteins;). The two different anti-DNAH12 antibodies were developed with the help Dia-An Biotech company in 2022, we have attempted to acquire the polypeptide fragments of target proteins to do peptide blocking but the material were disposed after the service. Luckily, we have got the target band of DNAH12 protein which showed strong signal in western blotting experiment and the band was not detected in knockout mice group; the immunofluorescence signals of DNAH12 were strong but not present in knockout mice group. Besides, we have tested that the inhouse raised rabbit antibody was suitable for IP experiment. The IP experiment also showed the raised rabbit antibody were able to immunoprecipitated the DNAH12 band in the *Dnah12*^{+/+} mice but not in *Dnah12*^{-/-} mice. Collectively, these data could support the specificity of the raised DNAH12 antibodies. We admire your suggestion and will require for the peptide material if we develop new antibodies.

Reviewer #2 (Recommendations For The Authors):

Recruitment of DNAH1 and DNALI1 to the flagella is dependent on DNAH12 expression, according to the data. What would be the mechanism that locates DNAH12 which lacks MTBD to the flagella?

Thank you for your insightful question. We are currently investigating the mechanisms that facilitate the loading of DNAH12 to the flagella. Based on existing data, we hypothesize that CCDC39 and/or CCDC40 may play a critical role in the recruitment of DNAH12 to sperm flagella during spermiogenesis (Nat Genet. 2011, PMID: 21131972; PMCID: PMC3509786; Nat Genet. 2011, PMID: 21131974; PMCID: PMC3132183). Furthermore, a structural study by Walton et al. showed that DNAH12 associates with CCDC39/CCDC40 proteins (Nature. 2023, PMID: 37258679; PMCID: PMC10266980). These findings suggest that CCDC39 and/or CCDC40 may play a role in facilitating the localization of DNAH12 to the flagella. Additional studies are needed to identify other potential factors involved in this process and to further elucidate the mechanisms underlying this complex biological phenomenon.

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