# Picking up speed: advances in the genetics of primary ciliary dyskinesia

Amjad Horani<sup>1</sup>, Steven L. Brody<sup>2</sup> and Thomas W. Ferkol<sup>1,3</sup>

Abnormal ciliary axonemal structure and function are linked to the growing class of genetic disorders collectively known as ciliopathies, and our understanding of the complex genetics and functional phenotypes of these conditions has rapidly expanded. While progress in genetics and biology has uncovered numerous cilia-related syndromes, primary ciliary dyskinesia (PCD) remains the sole genetic disorder of motile cilia dysfunction. The first disease-causing mutation was described just 13 y ago, and since that time, the pace of gene discovery has quickened. These mutations separate into genes that encode axonemal motor proteins, structural and regulatory elements, and cytoplasmic proteins that are involved in assembly and preassembly of ciliary elements. These findings have yielded novel insights into the processes involved in ciliary assembly, structure, and function, which will allow us to better understand the clinical manifestations of PCD. Moreover, advances in techniques for genetic screening and sequencing are improving diagnostic approaches. In this article, we will describe the structure, function, and emerging genetics of respiratory cilia, review the genotype-phenotype relationships of motor ciliopathies, and explore the implications of recent discoveries for diagnostic testing for PCD.

Primary ciliary dyskinesia (PCD) is a rare disease of children and was the first human disease linked to cilia dysfunction. Nearly four decades have passed since the disease was linked to ultrastructural defects of the axoneme (1,2). Following that time, and particularly in the last decade, there has been an explosion in knowledge related to cilia. Progress in cilia biology has been accompanied by the recognition that a large number of previously uncharacterized syndromes in children are the result of ciliary defects. This review will summarize major areas of progress in cilia-related disease, focusing on the genetics of PCD and areas for future investigation.

#### CILIA STRUCTURE AND FUNCTION

Cilia are present on the surface of most cells, and the basic axonemal structure is evolutionarily conserved. Cilia have historically been segregated into motile or primary (or sensory) types, and based on patterns of microtubule structure, fall into three general classes: motile "9+2", motile "9+0", and

nonmotile "9+0" (**Figure 1**). Several lines of evidence suggest that this classification system may be overly simplistic as their molecular features and functions overlap (3,4).

Cilia are anchored to the cytoplasm by a basal body, a specialized centriole located at the base that localizes the cilium in the extracellular space and organizes cilia assembly. Each cilium consists of hundreds of proteins arranged around a scaffold of  $\alpha$ - and  $\beta$ -monomers of tubulin arranged into helical protofilaments in the recognized pattern of paired microtubules (Figure 2). The central fibrillar structure, or axoneme, is covered by a membrane continuous with the plasma membrane. Major strides have been made in understanding how the structural and functional elements of cilia are synthesized and assembled within the cytoplasm and moved to the basal body prior to transport into the cilium. A welldescribed process called intraflagellar transport mediated by microtubule-associated motor proteins, kinesin and dynein, moves proteins cargos to and from the ciliary axoneme for the construction and maintenance of cilia (5). The mechanism by which proteins move from the cytoplasm to the apical domain of the ciliated epithelial cell for the proper construction of the cilium and how this process is disrupted in disease still needs to be defined.

Motile cilia are highly conserved to provide cell locomotion, fluid movement, and sexual reproduction. In humans, they line the upper and lower surface of the respiratory tract (including the Eustachian tubes), ependymal cells in the brain ventricles and spinal canal, as well as the Fallopian tubes. The flagellum of male spermatazoa has a similar axonemal structure. Cilium beating is generated by motor proteins that bind the nine outer microtubule doublets and interact with the central pair (central apparatus). Information regarding the motor proteins was first deduced from electron microscopic analysis of these dynein motors in the biflagellate green algae, Chlamydomonas reinhartii (6). Ultrastructural appearance and the localization of the dynein proteins relative to the microtubules have led to the term "inner" and "outer" arms to describe the different sets of motors. Subsequent biochemical analysis in alga and other organisms have revealed that the arms are complex structures, consisting of several heavy, intermediate, and light dynein chains that attach to the A microtubule, and contain ATPases.

<sup>1</sup>Department of Pediatrics, Washington University School of Medicine, St. Louis, Missouri; <sup>2</sup>Department of Medicine, Washington University School of Medicine, St. Louis, Missouri; <sup>3</sup>Department of Cell Biology and Physiology, Washington University School of Medicine, St. Louis, Missouri, Correspondence: Thomas W. Ferkol (ferkol\_t@kids.wustl.edu) Received 16 April 2013; accepted 20 August 2013; advance online publication 8 January 2014. doi:10.1038/pr.2013.200

Nexin links, which are components of the dynein regulatory complex, connect adjacent outer microtubular doublets, and with the radial spokes, control the sliding of cilia (7). The result is a ciliary stroke and rhythmic beating at ~8–14 Hz under normal conditions that contributes to a coordinated muco-ciliary wave moving with intra- and intercellular synchrony.

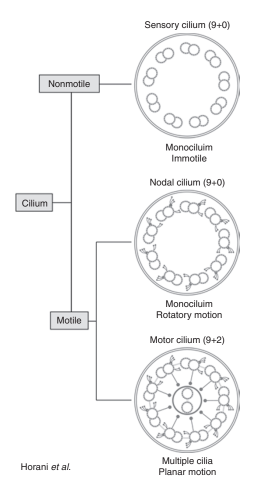


Figure 1. General classification of cilia.

The regulation of motile cilia movement can be influenced by changes in the ciliary microenvironment, such as the depth and viscosity of the apical surface fluid, redox conditions, and various infections and pollutants (including cigarette smoke) that can lead to an acquired ciliopathy (8–10). Thus, acquired or genetic disruption to the coordinated movement of motile cilia can result in ineffective mucociliary clearance and lung disease.

Distinct from motile cilia are primary (sensory) cilia, which are solitary structures present during interphase on most cell types. In mammals, sensory cilia are present on well-differentiated epithelia of sensory organs, biliary ductules, renal tubules, chondrocytes, and astrocytes. For years, these structures were considered vestigial remnants of little or no physiological significance (11). However, over the past 15 y, it has been shown that primary cilia are important signaling organelles and sense the extracellular environment. Cilia detect mechanical stimulation, chemosensation, and in specialized cases, changes in light, temperature, and gravity (12,13). In addition, development, growth, and repair functions are mediated by primary cilia through surface receptors, including sonic hedgehog, epidermal growth factor receptor, and platelet-derived growth factor receptor (14,15). Given this range of function, genetic defects in primary cilia have been associated with a growing number of clinically diverse pediatric conditions, collectively known as ciliopathies (16).

Motile cilia also have sensory functions, and syndromes caused by mutated proteins that overlap between motile and primary cilia are reported. For example, blindness due to mutations in the retinitis pigmentosa GTPase regulator gene (*RPGR*) is associated with motile cilia dysfunction (17). Further overlap in function is suggested by finding the expression of the polycystic kidney disease genes in motile cilia and an association with bronchiectasis in those individuals with cystic kidney disease (18,19). Motile cilia on human respiratory epithelia possess several members of the family of bitter taste receptors, identical to those in the tongue and nose (20). These sensory properties likely allow the motile cilia to adjust their movement in response to changes in their immediate environment.

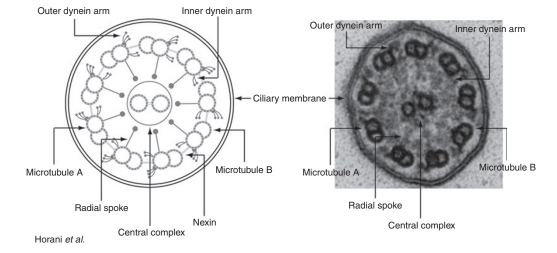


Figure 2. Ultrastructural features of the cilia. Schematic diagram (left) and transmission electron photomicrograph (right) of a normal motor cilium in cross-section, which shows the structural elements of the ciliary axoneme.

### Horani et al.

The nodal cilia are a third distinct class of cilia, but exist only transiently in the ventral node of the gastrula during embryonic development. Nodal cilia share a similar "9+0" microtubule arrangement as primary cilia but contain dyneins to provide motility (21,22). The leading hypothesis as to the function of the nodal cilia posits that the node includes two populations of cilia: motile and sensory (23,24). Through this arrangement, the motile cilia spin and generate leftward flow of extracellular fluid across the nodal surface that is then detected by sensory cilia and transformed to a biochemical signal that activates a cascade of transcription and growth factors to establish body sidedness (25). Defects within the nodal cilia pathways cause left-right laterality defects, such as situs inversus totalis, situs ambiguus, and heterotaxy associated with congenital heart disease, asplenia, and polysplenia (26,27). These findings have led to searches for ciliary anomalies in children with congenital heart disease to uncover the genetic cause of these defects (28).

#### DIAGNOSIS OF PCD: CHANGING STRATEGIES

Advances in cilia biology has been accompanied by rapid improvements in the diagnosis of the motile cilia dysfunction syndrome of PCD. In this regard, existing diagnostic studies, including transmission electron microscopy and cilia beat frequency, are expected to give way to nasal nitric oxide (NO) measurements and genetic testing. Recognition of clinical manifestations, which include neonatal respiratory distress, laterality defects, persistent infantile rhinitis, and recurrent upper and lower respiratory tract infections, continues to be the most important indication for diagnostic testing. Ciliary dysfunction leads to impaired mucociliary clearance and chronic airway infections, which results in progressive airway obstruction, atelectasis, and bronchiectasis, even early in life. Other manifestations of PCD include infertility and rarely, perinatal hydrocephalus. For a broader review, the reader is referred to several recent reviews that summarize the different clinical aspects of PCD (17,29).

The use of electron microscopy for the diagnosis of PCD dates to the original description of dynein arm defects in four subjects (1). Since then, ultrastructural defects have been the cornerstone for diagnosing PCD (29). Dynein arm defects account for 90% of cases with defined ultrastructural abnormalities. However, there are many limitations to the use and interpretation of electron microscopy. Ciliary defects can be acquired, and careful interpretation of the ultrastructural findings is necessary, since nonspecific changes may be seen related to exposure to environmental pollutants or infections. Chronic inflammation and infection common in patients with PCD often complicates the collection of an adequate epithelial biopsy with abundant ciliated cells required for analysis. Even when this limitation is overcome, high-quality tissue processing and ultrastructural analyses require sophisticated imaging equipment and expertise that are only available in few centers. Finally, newly discovered disease-causing gene mutations cause functional defects but are not associated with a consistent ultrastructural defect (30). Thus the presence of normal axonemal ultrastructure does not exclude PCD, which

underscores the difficulties of using electron microscopy as a sole diagnostic criterion.

Videomicroscopy has been advocated as a diagnostic test for PCD for measurement of cilia beat frequency and waveform analysis. The normal beat frequency of cilia obtained from healthy individuals ranges between 8 and 14 Hz, which varies with testing temperature and handling (31). Waveform analysis is an emerging method that may be reliable for diagnosing PCD (32), especially as more automated methods are introduced, but this approach currently lacks objective measures that can reliably be employed by clinicians. Ciliary motion in airway epithelial samples from PCD subjects can widely vary, from near immotility to subtle changes in beat pattern or frequency, all leading to ineffective mucociliary clearance. High-speed videomicroscopy is available only at a limited number of research centers and requires sophisticated software and expertise in analysis. This approach should be relegated to research only. Most importantly, inspection of motor cilia motion using standard light microscopy is never sufficient to support the diagnosis of PCD.

Perhaps, the most important diagnostic advance for PCD is the use of nasal NO measurement as a screening test. Nasal NO level measurements are sensitive and specific for the diagnosis of PCD and are currently part of the diagnostic criteria routinely employed by several centers in Europe and the United States (33,34). Low levels are also observed in some individuals with cystic fibrosis, thus emphasizing the importance of excluding this diagnosis when considering PCD. The precise relationship between motile cilia and nasal NO level is unclear, though the proximity of several regulatory enzymes to the ciliary basal bodies provides clues regarding their involvement in regulating cilia motility (35,36). Experimentally, NO can be detected in cells during normal cilia beating (37). Nitric oxide synthase-2 and nitric oxide synthase-3 have been found to be markedly reduced in subjects with PCD, and related mechanisms of NO metabolism have been implicated (38,39).

# GENETICS OF PCD: AN EMERGING CLASSIFICATION SYSTEM

In addition to the identification of cilia-related proteins, advances in techniques of genetic screening and sequencing are leading the way for the use of genetic testing as a diagnostic tool. PCD is usually inherited as an autosomal recessive trait, though rare cases of autosomal dominant and X-linked transmission have been reported (40,41). PCD is highly heterogenic owing to the large number of proteins involved in cilia assembly, structure, and function, and a growing number of genes have been implicated in disease. These mutant genes encode proteins that are involved in axonemal motors, ciliary structure and regulation, or ciliary assembly and preassembly (Table 1). The rate of discovery of new PCD genes has accelerated during the past 2 y. It is estimated that known PCD-causing mutations account only for about 60% of known PCD cases, but considering the pace of new discoveries, it is reasonable to expect that this percentage will rapidly increase.

Copyright © 2014 International Pediatric Research Foundation, Inc

Table 1. Ge	enes mutated i	n primary ciliary dyskin	esia	
Gene	Axonemal component	Ciliary ultrastructural defects	OMIM	Reference
DNAH5	ODA-HC	ODA defects	603335	(58)
DNAH11	ODA-HC	Normal	603339	(34,66)
DNAI1	ODA IC	ODA defects	604366	(59,65)
DNAI2	ODA IC	ODA defects	605483	(60)
DNAL1	ODA-LC	ODA defects	610062	(62)
TXNDC3	ODA LC/IC	Partial ODA defects	607421	(61)
RSPH4A	RSH	CP defects	612647	(63)
RSPH9	RSH	CP defects or normal	612648	(63)
CCDC39	DRC	Microtubule disorganization	613798	(64,67)
CCDC40	DRC	Microtubule disorganization	613799	(64,68)
CCDC164	DRC	DRC links defects	TBD	(73)
CCDC103	ODA docking	ODA defects	614677	(69)

CCDC114

HYDIN

DNAAF1

(LRRC50)

DNAAF2

DNAAF3

HEATR2

LRRC6

ODA docking

Central pair

Cytoplasmic

Cytoplasmic

Cytoplasmic

Cytoplasmic

Cytoplasmic

Mendelian Inheritance in Man; RSH, radial spoke; TBD, to be determined. The identification of PCD-associated genes has relied on a

**ODA** defects

**ODA+IDA** defects

**ODA+IDA** defects

**ODA+IDA** defects

**ODA+IDA** defects

**ODA+IDA** defects

CP, central pair; DRC, dynein regulatory complex; HC, heavy chain; IC, intermediate

chain; IDA, inner dynein arm; LC, light chain; ODA, outer dynein arm; OMIM, Online

**CP** defects

615038

610812

613190

612517

614566

614864

614930

(56,57)

(70)

(47)

(48)

(49)

(55)

(71, 72)

combination of experimental models, proteomic analysis, and sequencing of candidate genes (17,29,42-44). The best-studied model of motile cilia is the alga, C. reinhartii (45,46). Screening algae with defective motility or abnormal flagellar structure has been routinely used to study the function of orthologous ciliary proteins, many of which were linked to human disease (17,29,42-44,47). The second major advance leading to PCD gene identification was the collection of ciliogenesis-related transcriptomes and cilia proteomes that provided lists of cilia genes that could be linked to newly found disease mutations (48). More recently, massive parallel sequencing to analyze regions of interests in the genome has allowed more rapid identification of multiple new mutations in cohorts of PCD subjects without prior knowledge of candidate genes (49). Approaches that previously took months can now be completed in weeks or days. In addition, whole-exome sequencing has the potential to unravel the genetic causes of rare diseases and was recently successfully used to identify new candidate genes associated with PCD (50–52).

outer dynein arm: DNAH5, DNAI1, DNAL1, DNAI2, TXNDC3,

n particular, dynein axonemal intermediate chain 1 (DNAII; /IIM 604366) and dynein axonemal heavy chain 5 (DNAH5; /IM 603335), which encode components of the outer dynein rm and account for more than 30% of all cases (17,69). DNAI1 vas the first disease-associated gene to be identified using a candidate gene approach, relying on screening algae for abnormal flagellar beat and outer dynein arm defects. DNAI1 mutations were found in a large cohort at a prevalence of 9% of all identified PCD subjects (70). Mutations in DNAH5, another component of the outer dynein arm, were discovered using homozygosity mapping in large affected endogamous families (71), and later identified by sequencing in PCD subjects (53). Similarly, homozygosity mapping in consanguineous families has been used to identify mutations in other structural proteins, including the radial spoke head proteins 9 (RSPH9; MIM 612648) and 4A (RSPH4A; MIM 612647), both associated with absence of the central pair and motility defects (58,72). Other examples of mutated axonemal proteins associated with PCD discovered using the candidate gene approach include intermediate dynein chain DNAI2 (MIM 605483) (60), which comprise about 2% of PCD patients (55), and TXNDC3 (MIM 607421) (56).

Mutations in the dynein axonemal heavy chain 11 (DNAH11; MIM 603339) gene, that encodes an outer dynein arm protein, present with an intriguing phenotype (30,73). Unlike other PCD-causing mutations, it is not associated with an ultrastructural defect and cilia have normal (or more rapid) beat frequency. It is presumed that an abnormal waveform results in ineffective mucociliary clearance. Mutations in other genes, such as coiled-coil domain containing proteins CCDC39 (MIM 613798) and CCDC40 (MIM 613799), produce inconsistent ultrastructural abnormalities characterized by disordered microtubules in some but not all cilia, which underscores the clinical observation that current diagnostic testing will miss PCD cases.

Several nonstructural cilia-associated proteins have been found to be mutated in PCD individuals and result in the absence of outer and inner dynein arms. These mutations involve proteins that are considered to function in cilia assembly or "preassembly" pathways. Identification of the involvement of these proteins has spurred research into the mechanisms of cilia biogenesis. To understand the role of these mutations, cilia biogenesis can be conceptually divided

Review Genetics of primary ciliary dyskinesia

into the following processes: intra-axonemal transport, transfer into the cilium, and cytoplasmic preassembly.

As an organelle, the cilium projects into the environment and is uniquely isolated from the cell. Transfer of proteins from the cytoplasm into the cilium is limited by the periciliary diffusion barrier, which separates the ciliary membrane from the cell membrane (74,75). Direct access to the cilium is guarded by a ciliary pore complex, a structure analogous to nuclear pores (76,77). Located at the base of the cilium, the complex surrounds the basal body and its associated proteins. It is believed that proteins targeted to the cilium from the Golgi gain access by employing a ciliary localizing sequence, similar to the nuclear localizing sequence, as was described for KIF17 (78). Once through the ciliary pore, proteins move to a second regulatory region called the transition zone in the proximal part of the ciliary axoneme. Providing transport through this compartment and into cilia are numerous well-known intraflagellar transport proteins, which move essential protein cargos anterograde and retrograde along the length of the cilium (5). Precisely, how specific proteins are coded for precise delivery and assembly within the cilium and how the transition zone functions as a gate is not yet known.

Several gene mutations in preassembly proteins have been found to cause PCD. The first PCD-associated preassembly protein found was DNAAF2 (MIM 613190) (43), which was identified in mutant Oryzias latipes and later in PCD subjects who had complete absence of outer dynein arms. Localized within the apical region of the cell cytoplasm, DNAAF2 belongs to the proteins interacting with Hsp90 family and interacts with DNAI2 and the chaperone heat shock protein HSP70 to facilitate assembly or dynein complex transport into the cilia (43,79). Similar to DNAAF2, MOT48 is a protein interacting with Hsp90 localized in the cell body that was identified in the C. reinhartii ida10 strain that display motility defects and is associated with preassembly of both outer dynein arms and a subset of inner dynein arms, likely needed for the stability of dynein heavy chain components. Mutations in the related proteins DNAAF1 (LRRC50; MIM 612517) (42,56) and DNAAF3 (MIM 614566) (44) caused outer dynein arm defects in subjects with PCD. Evidence for assembly roles of these proteins is substantiated in PCD-mutant cells where components of the inner dynein arm were found to accumulate in the cytoplasm of mutated cells and fail to move into the cilium. These findings indicate the existence of a multistep cytoplasmic assembly pathway.

Another preassembly protein with mutations causative of PCD is HEATR2 (MIM 614864), which was recently implicated in dynein arm assembly (50). Similar to other preassembly factors, HEATR2 is localized to the cytoplasm of ciliated cells. However, HEATR2 is present throughout the cytoplasm rather than in the apical region, which suggests that it either functions at different stages of dynein assembly or is part of a chaperone complex that facilitates transfer of different dynein complexes along an "assembly line." This role was supported by the finding that HEATR2 mutations are associated with mislocalization of inner arm proteins. Other proteins with

HEAT-containing repeats are implicated in ciliary and nuclear import (78), suggesting that HEATR2 could potentially have similar mechanisms.

Mutations in *LRRC6* (MIM 244400) also lead to PCD (66,67). A member of a protein family with diverse functions, including splicing factors and nuclear transport (80), LRRC6 is found in the cytoplasm (67) and colocalizes with basal body markers. Mutations in LRRC6 were shown to downregulate expression of other dynein arm proteins and indicate an additional regulatory role. Thus, dynein arm preassembly is intricately regulated by both positive and negative feedback mechanisms.

#### AREAS FOR FUTURE STUDY IN PCD AND CONCLUSION

As described previously, newer techniques hold promise for discovery of additional PCD-associated mutations and genetic screening for PCD. Parallel sequencing has been used to analyze regions of interests and in the absence of candidates, whole-exome sequencing have been used to successfully identify new candidate genes associated with PCD (50-52). These advances, together with the rapid identification of genes associated with PCD seen during the last 2 y, have the potential to revolutionize the diagnostic testing and lead to earlier identification and treatment of affected children. There is every expectation that soon we will be able to identify the genetic cause for PCD in most suspected cases, which would potentially allow massive genetic screening of all PCD-associated genes in suspected individuals. The use of emerging technologies such as DNA microchips may facilitate the commercialization of such tests. It is envisioned that these approaches will be used in combination with clinical symptoms and nasal NO levels to diagnose patients with PCD.

The lack of an extensive mapping of mutations in PCD has hampered our ability to define a genotype–phenotype relationship. Moreover, the heterogeneity of disease and compound allelic mutations further complicate pinning specific phenotypes on unique mutations. The creation of specialized clinics in pediatric academic centers for the diagnosis and management of affected individuals with PCD in the United States and Europe will aid in the collection of genetic and clinical data. In parallel, analysis of the protein affected by specific gene mutations using biochemical and physiologic approaches will be an essential component of this work.

Despite progress in the genetics and diagnosis of PCD and related cilia disease, we currently lack effective or well-tested therapies. Instead, current treatments for PCD are directed at symptoms, largely extrapolated from experience with other conditions associated with bronchiectasis. Ultimately, it is our hope that the investigation of the mechanisms of cilia assembly and function, together with careful genetic and biochemical assessment, will provide better therapeutic options for individuals with PCD.

#### STATEMENT OF FINANCIAL SUPPORT

The authors were supported by the Children's Discovery Institute of St. Louis Children's Hospital and Washington University School of Medicine (to A.H., S.L.B., and T.W.F.) and National Institutes of Health (NIH, Bethesda, MD) awards HL056244 (to S.L.B.), R21 HL082657 (to T.W.F.), and U54 HL096458

(to T.W.F.). The last grant supports the Genetic Disorders of Mucociliary Clearance Consortium, part of the NIH Rare Diseases Clinical Research Network (RDCRN), with programmatic support from the NIH Office of Rare Diseases Research (ORDR). The views expressed do not necessarily reflect the official policies of the Department of Health and Human Services; nor does mention by trade names, commercial practices, or organizations imply endorsement by the US government.

#### REFERENCES

- 1. Afzelius BA. A human syndrome caused by immotile cilia. Science 1976;193:317–9.
- Eliasson R, Mossberg B, Camner P, Afzelius BA. The immotile-cilia syndrome. A congenital ciliary abnormality as an etiologic factor in chronic airway infections and male sterility. N Engl J Med 1977;297:1–6.
- Fliegauf M, Benzing T, Omran H. When cilia go bad: cilia defects and ciliopathies. Nat Rev Mol Cell Biol 2007;8:880–93.
- Gerdes JM, Davis EE, Katsanis N. The vertebrate primary cilium in development, homeostasis, and disease. Cell 2009;137:32–45.
- Pedersen LB, Rosenbaum JL. Intraflagellar transport (IFT) role in ciliary assembly, resorption and signalling. Curr Top Dev Biol 2008;85:23–61.
- Goodenough UW, Heuser JE. Substructure of inner dynein arms, radial spokes, and the central pair/projection complex of cilia and flagella. J Cell Biol 1985;100:2008–18.
- Heuser T, Raytchev M, Krell J, Porter ME, Nicastro D. The dynein regulatory complex is the nexin link and a major regulatory node in cilia and flagella. J Cell Biol 2009;187:921–33.
- Johnson NT, Villalón M, Royce FH, Hard R, Verdugo P. Autoregulation of beat frequency in respiratory ciliated cells. Demonstration by viscous loading. Am Rev Respir Dis 1991;144:1091–4.
- Hirst RA, Sikand KS, Rutman A, Mitchell TJ, Andrew PW, O'Callaghan C. Relative roles of pneumolysin and hydrogen peroxide from Streptococcus pneumoniae in inhibition of ependymal ciliary beat frequency. Infect Immun 2000;68:1557–62.
- Simet SM, Sisson JH, Pavlik JA, et al. Long-term cigarette smoke exposure in a mouse model of ciliated epithelial cell function. Am J Respir Cell Mol Biol 2010;43:635–40.
- 11. Sorokin SP. Reconstructions of centriole formation and ciliogenesis in mammalian lungs. J Cell Sci 1968;3:207–30.
- 12. Pazour GJ, Witman GB. The vertebrate primary cilium is a sensory organelle. Curr Opin Cell Biol 2003;15:105–10.
- Davenport JR, Yoder BK. An incredible decade for the primary cilium: a look at a once-forgotten organelle. Am J Physiol Renal Physiol 2005;289:F1159–69.
- 14. Caspary T, Larkins CE, Anderson KV. The graded response to Sonic Hedgehog depends on cilia architecture. Dev Cell 2007;12:767–78.
- Christensen ST, Pedersen SF, Satir P, Veland IR, Schneider L. The primary cilium coordinates signaling pathways in cell cycle control and migration during development and tissue repair. Curr Top Dev Biol 2008;85:261–301.
- Baker K, Beales PL. Making sense of cilia in disease: the human ciliopathies. Am J Med Genet C Semin Med Genet 2009;151C:281–95.
- Leigh MW, Pittman JE, Carson JL, et al. Clinical and genetic aspects of primary ciliary dyskinesia/Kartagener syndrome. Genet Med 2009;11: 473–87.
- Driscoll JA, Bhalla S, Liapis H, Ibricevic A, Brody SL. Autosomal dominant polycystic kidney disease is associated with an increased prevalence of radiographic bronchiectasis. Chest 2008;133:1181–8.
- Jain R, Javidan-Nejad C, Alexander-Brett J, et al. Sensory functions of motile cilia and implication for bronchiectasis. Front Biosci (Schol Ed) 2012;4:1088–98.
- Shah AS, Ben-Shahar Y, Moninger TO, Kline JN, Welsh MJ. Motile cilia of human airway epithelia are chemosensory. Science 2009;325:1131–4.
- Nonaka S, Tanaka Y, Okada Y, et al. Randomization of left-right asymmetry due to loss of nodal cilia generating leftward flow of extraembryonic fluid in mice lacking KIF3B motor protein. Cell 1998;95:829–37.
- Essner JJ, Vogan KJ, Wagner MK, Tabin CJ, Yost HJ, Brueckner M. Conserved function for embryonic nodal cilia. Nature 2002;418:37–8.

- Takao D, Nemoto T, Abe T, et al. Asymmetric distribution of dynamic calcium signals in the node of mouse embryo during left-right axis formation. Dev Biol 2013;376:23–30.
- 24. Yoshiba S, Shiratori H, Kuo IY, et al. Cilia at the node of mouse embryos sense fluid flow for left-right determination via Pkd2. Science 2012;338:226–31.
- 25. Watanabe D, Saijoh Y, Nonaka S, et al. The left-right determinant Inversin is a component of node monocilia and other 9+0 cilia. Development 2003;130:1725–34.
- 26. Larkins CE, Long AB, Caspary T. Defective Nodal and Cerl2 expression in the Arl13b(hnn) mutant node underlie its heterotaxia. Dev Biol 2012;367:15–24.
- 27. Kennedy MP, Omran H, Leigh MW, et al. Congenital heart disease and other heterotaxic defects in a large cohort of patients with primary ciliary dyskinesia. Circulation 2007;115:2814–21.
- 28. Nakhleh N, Francis R, Giese RA, et al. High prevalence of respiratory ciliary dysfunction in congenital heart disease patients with heterotaxy. Circulation 2012;125:2232–42.
- 29. Ferkol TW, Leigh MW. Ciliopathies: the central role of cilia in a spectrum of pediatric disorders. J Pediatr 2012;160:366–71.
- Knowles MR, Leigh MW, Carson JL, et al.; Genetic Disorders of Mucociliary Clearance Consortium. Mutations of DNAH11 in patients with primary ciliary dyskinesia with normal ciliary ultrastructure. Thorax 2012;67:433–41.
- Konietzko N, Nakhosteen JA, Mizera W, Kasparek R, Hesse H. Ciliary beat frequency of biopsy samples taken from normal persons and patients with various lung diseases. Chest 1981;80:Suppl 6:855–7.
- 32. Smith CM, Hirst RA, Bankart MJ, et al. Cooling of cilia allows functional analysis of the beat pattern for diagnostic testing. Chest 2011;140:186–90.
- O'Callaghan C, Chilvers M, Hogg C, Bush A, Lucas J. Diagnosing primary ciliary dyskinesia. Thorax 2007;62:656–7.
- 34. Strippoli MP, Frischer T, Barbato A, et al.; ERS Task Force onPrimary Ciliary Dyskinesia in Children. Management of primary ciliary dyskinesia in European children: recommendations and clinical practice. Eur Respir J 2012;39:1482–91.
- Stout SL, Wyatt TA, Adams JJ, Sisson JH. Nitric oxide-dependent cilia regulatory enzyme localization in bovine bronchial epithelial cells. J Histochem Cytochem 2007;55:433–42.
- Xue C, Botkin SJ, Johns RA. Localization of endothelial NOS at the basal microtubule membrane in ciliated epithelium of rat lung. J Histochem Cytochem 1996;44:463–71.
- 37. Jiao J, Wang H, Lou W, et al. Regulation of ciliary beat frequency by the nitric oxide signaling pathway in mouse nasal and tracheal epithelial cells. Exp Cell Res 2011;317:2548–53.
- Karadag B, James AJ, Gültekin E, Wilson NM, Bush A. Nasal and lower airway level of nitric oxide in children with primary ciliary dyskinesia. Eur Respir J 1999;13:1402–5.
- Pifferi M, Bush A, Maggi F, et al. Nasal nitric oxide and nitric oxide synthase expression in primary ciliary dyskinesia. Eur Respir J 2011;37:572–7.
- Narayan D, Krishnan SN, Upender M, et al. Unusual inheritance of primary ciliary dyskinesia (Kartagener's syndrome). J Med Genet 1994;31:493–6.
- Moore A, Escudier E, Roger G, et al. RPGR is mutated in patients with a complex X linked phenotype combining primary ciliary dyskinesia and retinitis pigmentosa. J Med Genet 2006;43:326–33.
- Loges NT, Olbrich H, Becker-Heck A, et al. Deletions and point mutations of LRRC50 cause primary ciliary dyskinesia due to dynein arm defects. Am J Hum Genet 2009;85:883–9.
- Omran H, Kobayashi D, Olbrich H, et al. Ktu/PF13 is required for cytoplasmic pre-assembly of axonemal dyneins. Nature 2008;456:611–6.
- 44. Mitchison HM, Schmidts M, Loges NT, et al. Mutations in axonemal dynein assembly factor DNAAF3 cause primary ciliary dyskinesia. Nat Genet 2012;44:381–9, S1–2.
- 45. Li JB, Gerdes JM, Haycraft CJ, et al. Comparative genomics identifies a flagellar and basal body proteome that includes the BBS5 human disease gene. Cell 2004;117:541–52.
- Mitchell DR. The evolution of eukaryotic cilia and flagella as motile and sensory organelles. Adv Exp Med Biol 2007;607:130–40.

### Horani et al.

- Dutcher SK. Elucidation of basal body and centriole functions in Chlamydomonas reinhardtii. Traffic 2003;4:443–51.
- Gherman A, Davis EE, Katsanis N. The ciliary proteome database: an integrated community resource for the genetic and functional dissection of cilia. Nat Genet 2006;38:961–2.
- Metzker ML. Sequencing technologies the next generation. Nat Rev Genet 2010;11:31–46.
- Horani A, Druley TE, Zariwala MA, et al. Whole-exome capture and sequencing identifies HEATR2 mutation as a cause of primary ciliary dyskinesia. Am J Hum Genet 2012;91:685–93.
- Knowles MR, Leigh MW, Ostrowski LE, et al.; Genetic Disorders of Mucociliary Clearance Consortium. Exome sequencing identifies mutations in CCDC114 as a cause of primary ciliary dyskinesia. Am J Hum Genet 2013;92:99–106.
- 52. Onoufriadis A, Paff T, Antony D, et al.; UK10K. Splice-site mutations in the axonemal outer dynein arm docking complex gene CCDC114 cause primary ciliary dyskinesia. Am J Hum Genet 2013;92:88–98.
- Olbrich H, Häffner K, Kispert A, et al. Mutations in DNAH5 cause primary ciliary dyskinesia and randomization of left-right asymmetry. Nat Genet 2002;30:143–4.
- Guichard C, Harricane MC, Lafitte JJ, et al. Axonemal dynein intermediate-chain gene (DNAI1) mutations result in situs inversus and primary ciliary dyskinesia (Kartagener syndrome). Am J Hum Genet 2001;68:1030–5.
- Loges NT, Olbrich H, Fenske L, et al. DNAI2 mutations cause primary ciliary dyskinesia with defects in the outer dynein arm. Am J Hum Genet 2008;83:547–58.
- Duriez B, Duquesnoy P, Escudier E, et al. A common variant in combination with a nonsense mutation in a member of the thioredoxin family causes primary ciliary dyskinesia. Proc Natl Acad Sci USA 2007;104: 3336–41.
- Horváth J, Fliegauf M, Olbrich H, et al. Identification and analysis of axonemal dynein light chain 1 in primary ciliary dyskinesia patients. Am J Respir Cell Mol Biol 2005;33:41–7.
- Castleman VH, Romio L, Chodhari R, et al. Mutations in radial spoke head protein genes RSPH9 and RSPH4A cause primary ciliary dyskinesia with central-microtubular-pair abnormalities. Am J Hum Genet 2009;84: 197–209.
- Blanchon S, Legendre M, Copin B, et al. Delineation of CCDC39/CCDC40 mutation spectrum and associated phenotypes in primary ciliary dyskinesia. J Med Genet 2012;49:410–6.
- Pennarun G, Escudier E, Chapelin C, et al. Loss-of-function mutations in a human gene related to Chlamydomonas reinhardtii dynein IC78 result in primary ciliary dyskinesia. Am J Hum Genet 1999;65:1508–19.
- Bartoloni L, Blouin JL, Pan Y, et al. Mutations in the DNAH11 (axonemal heavy chain dynein type 11) gene cause one form of situs inversus totalis and most likely primary ciliary dyskinesia. Proc Natl Acad Sci USA 2002;99:10282–6.
- 62. Merveille AC, Davis EE, Becker-Heck A, et al. CCDC39 is required for assembly of inner dynein arms and the dynein regulatory complex and for normal ciliary motility in humans and dogs. Nat Genet 2011;43:72–8.

- 63. Becker-Heck A, Zohn IE, Okabe N, et al. The coiled-coil domain containing protein CCDC40 is essential for motile cilia function and left-right axis formation. Nat Genet 2011;43:79–84.
- 64. Panizzi JR, Becker-Heck A, Castleman VH, et al. CCDC103 mutations cause primary ciliary dyskinesia by disrupting assembly of ciliary dynein arms. Nat Genet 2012;44:714–9.
- Olbrich H, Schmidts M, Werner C, et al.; UK10K Consortium. Recessive HYDIN mutations cause primary ciliary dyskinesia without randomization of left-right body asymmetry. Am J Hum Genet 2012;91:672–84.
- 66. Kott E, Duquesnoy P, Copin B, et al. Loss-of-function mutations in LRRC6, a gene essential for proper axonemal assembly of inner and outer dynein arms, cause primary ciliary dyskinesia. Am J Hum Genet 2012;91:958–64.
- 67. Horani A, Ferkol TW, Shoseyov D, et al. LRRC6 mutation causes primary ciliary dyskinesia with dynein arm defects. PLoS ONE 2013;8:e59436.
- 68. Wirschell M, Olbrich H, Werner C, et al. The nexin-dynein regulatory complex subunit DRC1 is essential for motile cilia function in algae and humans. Nat Genet 2013;45:262–8.
- 69. Zariwala MA, Omran H, Ferkol TW. The emerging genetics of primary ciliary dyskinesia. Proc Am Thorac Soc 2011;8:430–3.
- Zariwala MA, Leigh MW, Ceppa F, et al. Mutations of DNAI1 in primary ciliary dyskinesia: evidence of founder effect in a common mutation. Am J Respir Crit Care Med 2006;174:858–66.
- Omran H, Häffner K, Völkel A, et al. Homozygosity mapping of a gene locus for primary ciliary dyskinesia on chromosome 5p and identification of the heavy dynein chain DNAH5 as a candidate gene. Am J Respir Cell Mol Biol 2000;23:696–702.
- 72. Yang P, Diener DR, Yang C, et al. Radial spoke proteins of Chlamydomonas flagella. J Cell Sci 2006;119(Pt 6):1165–74.
- Pifferi M, Michelucci A, Conidi ME, et al. New DNAH11 mutations in primary ciliary dyskinesia with normal axonemal ultrastructure. Eur Respir J 2010;35:1413–6.
- Nachury MV, Seeley ES, Jin H. Trafficking to the ciliary membrane: how to get across the periciliary diffusion barrier? Annu Rev Cell Dev Biol 2010;26:59–87.
- Garcia-Gonzalo FR, Corbit KC, Sirerol-Piquer MS, et al. A transition zone complex regulates mammalian ciliogenesis and ciliary membrane composition. Nat Genet 2011;43:776–84.
- 76. Satir P, Christensen ST. Overview of structure and function of mammalian cilia. Annu Rev Physiol 2007;69:377–400.
- Follit JA, Tuft RA, Fogarty KE, Pazour GJ. The intraflagellar transport protein IFT20 is associated with the Golgi complex and is required for cilia assembly. Mol Biol Cell 2006;17:3781–92.
- Dishinger JF, Kee HL, Jenkins PM, et al. Ciliary entry of the kinesin-2 motor KIF17 is regulated by importin-beta2 and RanGTP. Nat Cell Biol 2010;12:703–10.
- 79. Kobayashi D, Takeda H. Ciliary motility: the components and cytoplasmic preassembly mechanisms of the axonemal dyneins. Differentiation 2012;83:S23–9.
- Kobe B, Kajava AV. The leucine-rich repeat as a protein recognition motif. Curr Opin Struct Biol 2001;11:725–32.