

Cilia/Ift protein and motor-related bone diseases and mouse models

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1. ABSTRACT

Primary cilia are essential cellular organelles projecting from the cell surface to sense and transduce developmental signaling. They are tiny but have complicated structures containing microtubule (MT)-based internal structures (the axoneme) and mother centriole formed basal body. Intraflagellar transport (lft) operated by lft proteins and motors are indispensable for cilia formation and function. Mutations in lft proteins or lft motors cause various human diseases, some of which have severe bone defects. Over the last few decades, major advances have occurred in understanding the roles of these proteins and cilia in bone development and remodeling by examining cilia/lft protein-related human diseases and establishing mouse transgenic models. In this review, we describe current advances in the understanding of the cilia/lft structure and function. We further summarize cilia/lft-related human diseases and current mouse models with an emphasis on bone-related phenotypes, cilia morphology, and signaling pathways.

2. INTRODUCTION

Normal vertebrate skeletons, composed of cartilage and bone, are critical for locomotion, respiration, and protection of the vital organs (1). Both cartilage and bone are derived from embryonic mesenchyme (2, 3). Hyaline cartilage is the most widespread cartilage and prevails in the fetal skeleton, while bone is predominant at a later stage and persists postnatally throughout life (1). Most bones are formed by replacement of existing cartilage rudiments. This process is known as endochondral bone formation, which starts with embryonic mesenchymal cells condensing at the sites where the skeletal elements will form. The condensed mesenchymal cells result in the formation of the cartilaginous anlagen, which later is ossified to become bone by osteoblasts (4-6). In contrast, mesenchymal condensations that prefigure the clavicles, mandible, and certain bones of the skull, directly differentiate into osteoblasts. This process does not require a cartilaginous template and is called intramembranous ossification (1-3). Skeletal formation and homeostasis are strictly controlled

by systemic and local regulators, such as Wnt, Hedgehog (Hh) (7), Notch (8), bone morphogenetic protein 2 (BMP2) (9), and thyroid hormones (10, 11). It has uncovered over 300 genetic diseases with disorders in the growth and development of the skeleton (12). Bone development abnormalities often lead to bony deformities that severely impact patients' quality of life and general health and sometimes lead to premature death (1, 2, 13, 14). Among these genetic diseases, a certain groups are caused by disorders of the primary cilia (15).

Primary cilia are small microtubule (MT) based organs that are projected from the surface of nearly every mammalian cells and function as antennae. They sense a diverse range of extracellular signals, such as growth factors and hormones, through various surface receptors localized on the ciliary membrane, and transmit the signals back into the nucleus. Thus cilia are important for embryonic and postnatal development of many vital organs (16-18). Defects of cilia or cilia-related protein cause a wide variety of human disease symptoms, including polycystic kidneys, hydrocephalus, vision and hearing loss as well as severe bone abnormalities, suggesting that the role of cilia in multiple organ development is indispensable (18-25).

Intraflagellar transport (lft), a bidirectional transport system run by lft protein complexes and lft motors, is found to be essential for building cilia and maintaining cilia function (26, 27). lft protein complexes are formed with complex A and complex B containing about 20 lft proteins (28). Mutations in lft proteins (such as lft80, lft122, and lft43) or lft motors (such as DYNC2H1) usually lead to cilia defects and human diseases (29-31). Those groups of human diseases with primary cilia malfunction display disorders of multiple organs, including kidney, liver, brain, and limb (30) and are called ciliopathies (29, 32). A certain portion of ciliopathies, such as Jeune asphyxiating thoracic dystrophy (JATD) and short rib polydactyly syndromes (SRP), result in several bone defects, suggesting the important role of cilia in bone development (29, 33-35).

During the 1960s, scientists found cilia present in chondrocytes (36). Subsequent work has revealed that cilia are also present in osteoblasts, osteocytes, and mesenchymal stem cells (37-41). Bone is a sensitive and dynamic tissue that constantly remodels to meet the demands of its environment (42, 43). Cilia are considered to function as chemosensors and mechanosensors for bone formation and maintenance (43-45). Multiple bone developmental signaling pathways, such as Hh and Wnt signaling pathways, have been linked to primary cilia, lft proteins and motors (46-49). Both Indian hedgehog (Ihh) and Sonic hedgehog (Shh) are required for skeletal development (50-53). Ihh is an important signaling molecule in endochondral ossification because Ihh null mice display shortened face and tail with extremely short limbs and narrow rib cages (51). Shh is required for the early stage of patterning of the limb (54, 55). Genetic screens for embryonic patterning mutations first uncovered the connection between mammalian Hh signaling and primary cilia (56). Since then, the Hh pathway has become the most intensively studied pathway among the cilia-related signaling pathways (see review (57) and (58)). It is now clearly established that the core components of Hh signaling pathway, including Hh receptor Patched 1 (Ptch1), *Smoothed* (Smo) protein, and Gli transcription factors are localized to cilia (57). Ptch1 is found at the base of primary cilia in the absence of Hh. Upon exposure to Hh, Ptch1 moves out of cilia (59, 60), and then Smo moves into the cilia and shifts the Gli process from a transcriptional repressor to transcriptional activator (61). Functional intact cilia are required for the Hh signaling transduction, and mutations in cilia/lft proteins and motors disrupt Hh signaling pathway. However, mutations in different genes lead to different phenotypes, suggesting the distinct roles of individual lft proteins and motors in Hh signaling or their involvement of other signaling pathways (57). Wnt pathway is another critical pathway for skeletal development and homeostasis (62, 63). However, the relationship between Wnt and primary cilia is controversial (58, 64). Wnt signaling cascades can be broadly divided into two general categories: 1) canonical Wnt/ β -catenin pathway (stabilizing the transcription of co-activator β -catenin and therefore turning on gene expression) and 2) the non-canonical Wnt/planar cell polarity (PCP) pathway (65). Different mutations in cilia/lft proteins cause different changes in Wnt signaling pathway (64, 66, 67). However, the role of cilia/lft proteins and motors in Wnt signaling transduction remains largely unknown.

To study the fundamental biology of cilia/lft proteins and motors, various animal models, including tetrahymena thermophila, zebrafish, and mouse, have been developed (68-70). Mouse is genetically and pathophysiologically similar to human and relatively easy to be genetically manipulated. Therefore, mouse has become the favorite and reliable mammalian model for studies of bone biology (71, 72). Within the last decade, significant progress in the understanding of the cilia/lft proteins and motors in bone development and diseases has been made using various mouse genetic models and we summarize those advances here.

3. STRUCTURE OF IFT PROTEINS/ MOTORS AND CILIA

The structure of the cilia is conserved throughout evolution. When cell is in G_0 stage, primary cilium is generated from a "mother" centriole of the centrosome, which becomes the basal body in the cilia structure (73). Centriole is a cylinder comprising nine triplet MTs that provide the template for the formation of an MT-based internal structure (the axoneme) (74). Two of the most common axoneme architectures are "9 + 2" for motile cilia (9 peripheral MT doublets and 2 single central MTs) and "9 + 0" for primary cilia (only 9 peripheral MT doublets) (75). Since ribosome and membrane vesicles are absent from the cilia, ciliary proteins have to be synthesized in the cytoplasm and actively transported into the cilia along MTs (73). This highly conserved process is called lft (76), which moves macromolecules from the base to the tip of a cilium (anterograde transport) and from the tip of the cilium back to the cell body (retrograde transport) (77). Effective lft involves the lft complexes and lft motors. It is believed that the heterotrimeric kinesin-II motor protein drives the anterograde transport (78), whereas cytoplasmic dynein-2 moves the rafts back in the retrograde direction (79).

lft complexes serve as adaptors to mediate the contacts between cargo proteins and motors (80). lft was first reported in *Chlamydomonas* in 1993 by Kozminski *et al* (76, 81), and later lft particles were identified in *Chlamydomonas* by biochemical purifications (82, 83). By biochemical purification, Cole *et al* first reported that lft complex has two sub-complexes named complex A (lft-A) and complex B (lft-B) (82). Subsequently, four proteins in complex A (lft144, lft140, lft139 and lft122) and eleven proteins in complex B (lft172, lft88, lft81, lft80, lft74, lft72, lft57/55, lft52, lft46, lft27 and lft20)

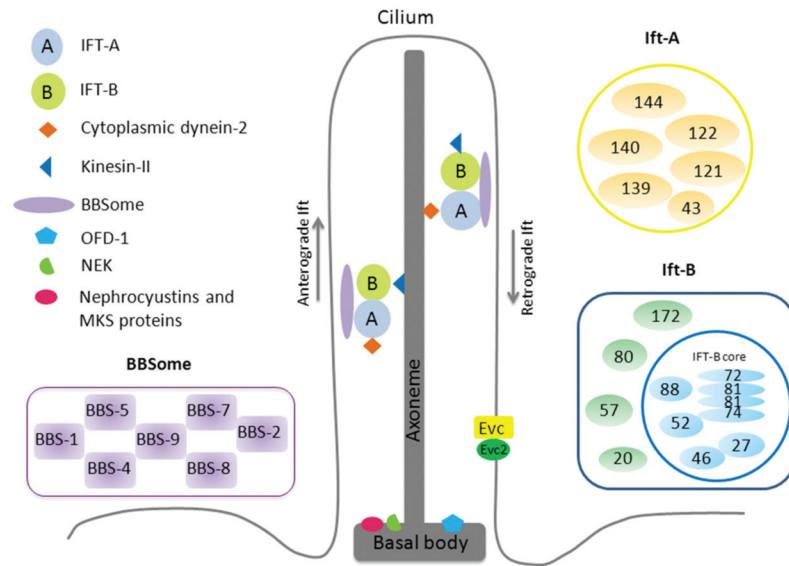


Figure 1. The primary cilium and ciliary proteins (adapted from (28, 87, 234, 235)). The structure of primary cilia is conserved throughout evolution with a basal body (“mother” centriole) and 9 peripheral MT doublets (the axoneme). The Ift-A contains 6 known proteins (Ift144, Ift140, Ift139, Ift122, Ift121 and Ift43) and is in charge of retrograde transport (from ciliary tip back to the cell body) with cytoplasmic dynein-2 as motor. Ift-B contains 14 known proteins (Ift20, Ift22, Ift25, Ift27, Ift46, Ift52, Ift54, Ift57, Ift70, Ift74/Ift72, Ift80, Ift81, Ift88 and Ift172) and is involved in anterograde transport (from the base to ciliary tip) with heterotrimeric kinesin-II as motor. Ift81 and Ift74/72 form a tetrameric complex and interact with Ift88, Ift52, Ift46 and Ift27 formed the core of complex B. Besides Ift motors and Ift complexes, BBS proteins are also important for cilia formation and maintenance. Seven highly conserved BBS proteins (BBS1, BBS2, BBS4, BBS5, BBS7, BBS8 and BBS9) formed BBSome. Evc and Evc-2 are located at the basal body and the base of the axoneme. Other proteins like OFD, NEK, Nephrocytins and MKS proteins are found localized in the basal body. And mutation of these proteins caused diseases.

have been identified (82). The proteins were named according to their size (in kD). Later, Ift43 and then Ift121 were identified in the complex A (82, 84, 85). Thus complex A is thought to be composed of six subunits including Ift144, Ift140, Ift139, Ift122, Ift43 and Ift121.

Ift-B complex is more complicated and contains 14 known proteins (Ift20, Ift22, Ift25, Ift27, Ift46, Ift52, Ift54, Ift57, Ift70, Ift74/Ift72, Ift80, Ift81, Ift88 and Ift172) (18, 82, 85, 86). Luckert *et al* used increased ionic strength to dissociate the Ift-B of *Chlamydomonas reinhardtii* and found that the biochemically intact core of Ift-B contains Ift88, Ift81, Ift74/72, Ift52, Ift46, and Ift27 (87). Ift72 and Ift74 are derived from the same gene and present in a 1:1 stoichiometry. So it is usually combined into a single term, Ift74/72 (87, 88). The other subunits including Ift172, Ift80, Ift57, and Ift20 are not required for the core and can be dissociated with the core by using a high salt concentration. Among those core proteins, Ift81 and Ift74/72 are believed to form a tetrameric complex to serve as a scaffold for the interaction with other subunits (87).

All of the Ift proteins are conserved in mammals. Ift proteins are typically found along the cilia and in the base of the cilia (89). Anterograde transport is regulated by Ift-B, while the retrograde transport is related to Ift-A (28, 90). Mutations in Ift-B usually block cilia assembly (91, 92), similar to kinesin-II mutation (93, 94), while defects in Ift-A protein(s) typically result in malformed cilia with accumulation and function defects of Ift proteins (95-97) that resemble the mutations in cytoplasmic dynein-2 (48) (Figure 1).

Besides Ift motors and Ift complexes, sixteen identified Bardet-Biedl syndrome proteins (BBS1-16) are also important for cilia formation and maintenance. Seven highly conserved BBS proteins (BBS1, BBS2, BBS4, BBS5, BBS7, BBS8 and BBS9) and one newly defined interacting protein BBIP10 formed BBSome (98, 99). BBSome is indispensable for ciliary membrane biogenesis, microtubule stability, and acetylation (99). Other BBS proteins have a variety of functions including guanosine triphosphate hydrolases (GTPase) and ubiquitin ligase (See review (100)).

Table 1. Currently known genes that are involved in skeletal ciliopathies

Disease	lft complex A	lft complexes B	lft motors	Others	References
JATD	<i>TTC21B, WDR19, lft140</i>	<i>lft80</i>	<i>DYNC2H1</i>		(35, 108, 109, 111, 233)
MSS	<i>lft140</i>				(111)
BBS				<i>BBS1-BBS16</i>	(114)
EVC				<i>Evc and Evc2</i>	(117-120)
WAD				<i>Evc2</i>	(121)
OFD				<i>OFD1</i>	(126)
SRP type II			<i>DYNC2H1</i>	<i>NEK1</i>	(128, 131)
SRP type III		<i>lft80</i>	<i>DYNC2H1</i>		(35, 130)
SRP severe type	<i>WDR35</i>				(34)
CED	<i>lft122, lft43, WDR35, WDR19</i>				(109, 132, 135-137)
MKS				<i>BBS2, BBS4, BBS6, MKS1/FLJ20345, MKS3/TMEM67, TMEM216, CEP290, RPGRIP1L, CC2D2A, TCTN2, B9D1, B9D2, and NPHP3</i>	(114)
Jeune asphyxiating thoracic dystrophy (JATD), Intraflagellar transport (lft), Mainzer-Saldino-syndrome (MSS), Bardet-Biedl syndrome (BBS), Ellis-van Creveld syndrome (EVC), WAD (Weyers acrofacial dysostosis), orofacialdigital syndrome (OFD), short rib polydactyly syndromes (SRP), cranioectodermal dysplasia (CED), Meckel syndrome (MKS)					

A large amount of cargo proteins have been found in the cilia or associated with cilia structures (75). Those proteins include the signaling components such as Gli and Smo in the Hh signaling pathway. Some of them are proteins located at a specific region of the cilium to regulate cilia formation or cilia-related signaling transduction such as Ellis-van Creveld syndrome protein homolog (*Evc*) and oral-facial-digital syndrome 1 (*OFD1*) (101, 102). Mutations of those proteins also cause ciliopathies.

4. CILIA/IFT PROTEIN AND MOTOR-RELATED HUMAN DISEASES

Cilia were first found in the kidneys and the thyroid gland one century ago (103). Subsequently, cilia have been found in many other organs and cell types (31). Further work revealed a dozen human genetic diseases caused by cilia-related gene mutation, which are commonly called as ciliopathies (31, 104). Those disorders includes BBS, OFD, Meckel-Gruber syndrome (MKS), Joubert syndrome (JBTS)/JBTS-related disorder (JSRD), JATD, Leber congenital amaurosis (LCA), Primary ciliary dyskinesia (PCD)/Kartagener syndrome (KS), Ellis-van Creveld syndrome (EVC), Nephronophthisis

(NPHP), Senior-Loken syndrome (SLS), autosomal dominant and recessive polycystic kidney disease (ADPKD and ARPKD), Usher syndrome (US), and Alström syndrome (ALMS) (104, 105). Those ciliopathies share common clinical features and involve many major organs, including kidney, brain, limb, retina, liver, and bone (29, 32, 104, 105).

More than 30 different genes have been linked to the skeletal-related ciliopathies. These are dynein-2 motor (*Dync2h1*), lft genes (*lft80, lft122, lft43, lft140, WDR35, WDR19, and TTC21B*) as well as the genes responsible for the basal body (*NEK1, Evc, Evc2, BBS1-16, OFD1, MKS1, MKS3, TMEM216, CEP290, RPGRIP1L, CC2D2A, TCTN2, B9D1, B9D2, and NPHP3*). Mutations of those genes result in common skeleton defects, including polydactyly, shortened bones, and craniofacial dysmorphology (Table 1) (29). Here, we only summarize the ciliopathies with bone phenotypes and their underlying genetic defects.

4.1. JATD and Mainzer-Saldino-Syndrome (MSS)

JATD, also known as asphyxiating thoracic dystrophy (ATD) was first reported in 1955 by Jeune

et al. The patient with JATD had a narrow thoracic cavity and multiple cartilage anomalies, and died due to respiratory insufficiency during the early neonatal period (106). JATD is a rare autosomal recessive skeletal dysplasia with short limbs, narrow thorax, trident acetabular roof, and occasional polydactyly (107). Studies have suggested that mutations of *Ift80*, *DYNC2H1*, *TTC21B* (encoding Ift139) and *WDR19* (encoding Ift144) are associated with JATD (33, 35, 40, 108-110). Respiratory failure is the usual cause of death in JATD in early infancy (110). MSS is related to JATD but has a milder rib phenotype and additional renal and eye defects (111).

Very recently, *Ift140* mutation was identified in JATD and MSS (111, 112). It has been found that Ift140 patients display mild chest narrowing, end-stage renal failure, and ocular dysfunction (112). Ciliary abundance and localization of IFT-A proteins are changed in the fibroblasts from *Ift40* mutant patients (111).

4.2. BBS

BBS is a pleiotropic genetic disorder with the phenotypes of retinal dystrophy, obesity, post-axial polydactyly, renal dysfunction, learning difficulties and hypogonadism (113). Polydactyly, hepatic fibrosis, and kidney, genital and heart malformations can be observed at birth (114). Sixteen BBS gene (*BBS1* to *BBS16*) mutations account for 80% of BBS. The majority of mutations occur in *BBS1* and *BBS10* (114).

4.3. EVC and Weyers acrofacial dysostosis (WAD)

EVC was first reported by Richard Ellis and Simon van Creveld in 1940. Patients had retarded growth, narrow rib cage, supernumerary fingers and toes, orofacial abnormalities and heart abnormalities (115, 116). WAD was named after Helmut Weyers, who reported it in 1952. EVC and WAD are allelic disorders, but WAD has less severe symptoms and a different pattern of inheritance (117). They are both caused by the loss-of-function mutations of *Evc* or *Evc2* genes (117-121). Valencia *et al* further demonstrated that Hh signal transduction is impaired when a mutant murine *Evc2* Weyer variant is expressed in NIH 3T3 cells (121).

4.4. OFD

OFD is an X-linked dominant inheritance disorders with 13 different described types (type I to type XIII) classified by their patterns of signs and symptoms (122, 123). The phenotype includes oral,

facial, and digital anomalies (124, 125). OFD type I, which is also known as Papillon-Leage and Psaume syndrome, has the highest incidence (122). Ferrante *et al* identified the responsible gene *OFD1* (formerly named *Cxorf5/71-7a*) for this syndrome in 2001 (126). *OFD1* is found missense, nonsense, splice, and frameshift mutation in *OFD1* patients (126). Further studies showed *OFD1* is a centrosomal protein and localized in the basal body of primary cilia (102, 127). Whether *OFD1* mutation is responsible for other types of OFD is not clear now.

4.5. SRP

SRP is a group of lethal skeletal dysplasia that are characterized by short limbs, narrow thorax, polydactyly, and multiple anomalies of major organs (128). Four lethal types have been described based on radiological findings, including Saldino-Noonan syndrome (type I), Majewski syndrome (type II), Verma-Naumoff syndrome (type III) and Beemer-Langer syndrome (Type IV) (34, 128). In addition, JATD and EVC are phenotypically related to SRP but are compatible with life. Recently, Ho *et al* suggested that JATD and SRP type III are variants of the same disorder and SRP type III has more severe phenotype in bone and other organs (129). Mutation of *Ift80* or *DYNC2H1* causes those diseases (35, 70, 130). Most recently, mutation of either *DYNC2H1* or *Never-in-mitosis Kinase 1* (*NEK1*), which regulates cell cycle-associated ciliogenesis, was identified in SRP type II (128, 131). A severe form of SRP was reported with in-frame homozygous deletion of exon 5 in *WDR35* (orthologous of *Ift121* in mammals) (34).

4.6. Cranioectodermal dysplasia (CED, or sensenbrenner syndrome)

CED is a rare autosomal recessive skeletal dysplasia (132). The syndromes of CED patients are similar to JATD patients as they have short stature, limbs and ribs, narrow chest, brachydactyly, renal failure, and hepatic fibrosis (132). They also show dolichocephaly with frontal bossing and craniosynostosis (133, 134). CED is genetically heterogeneous and is caused by mutations in *Ift122* (132), *Ift43* (135), *WDR35* (136, 137) or *WDR19* (109).

4.7. Meckel syndrome (MKS, or Meckel-Gruber Syndrome, Gruber Syndrome, Dysencephalia Splanchnocystica)

MKS is a severe fetal developmental disorder characterized by occipital encephalocele, cystic kidney dysplasia, fibrotic changes of the

liver and polydactyly (138). MKS is genetically heterogeneous and at least ten genes have been associated with MKS, including *MKS1* (*Meckel syndrome, type 1*)/*BBS13* (encoding MKS1, localized at the ciliary body), *MKS3/TMEM67* (*Transmembrane protein 67*, encoding MECKELIN, which localized to the cilium base), *TMEM216* (encoding TMEM216, which localized to the base of the primary cilia), *CEP290/BBS14* (encoding centrosomal protein CEP290/nephrocystin-6, which is involved in chromosome segregation and localized around the centrosome), *RPGRIP1L* (encoding RPGRIP1L, which colocalizes to the basal body and centrosomes), *CC2D2A* (encoding CC2D2A, which localizes to basal body and the transition zone), *TCTN2* (encoding TCTN2, which localizes to transition zone), *B9D1* (encoding B9D1, which localizes to the transition zone complex), *B9D2* (encoding B9D2, which localizes to the transition zone complex), and *NPHP3* (nephrocystin-3, encoding NPHP3, which localizes to the primary cilia) (139-144).

Karmous-Benailly discovered mutations in *BBS2*, *BBS4*, and *BBS6* in MSK patients. They suggested that prenatal presentation of BBS may mimic MKS (114).

5. FUNCTION OF IFT PROTEINS OR MOTORS IN EXISTING MOUSE MODELS

To deeply understand the function of cilia-related proteins, a variety of mouse genetic models have been generated in which lft proteins or motors are inactivated by using different genetic mutant approaches. To make the summary of lft proteins and motors clearly understood, we first briefly introduce the most commonly used techniques to generate gene mutation models in mice as described in section 5.1., and then further summarize the published mouse models that have mutations in lft proteins and motors. We focus on the bone-related phenotypes, ciliogenesis, and altered signaling pathways.

5.1. Methods to generate gene mutational mouse models

There are two main methods to generate gene mutational mouse models. One is directed and disease driven; and the other is non-directed, randomly generated mutation. The common directed techniques include most powerful method of transgenesis. The non-directed, random mutation-driven method is the application of radiation and

chemicals such as N-ethyl-N-nitrosourea (ENU) to induce mutations (145).

Transgenesis can be accomplished by two methods. One is pronuclear injection of fertilized oocytes, which is predominantly used to overexpress endogenous or mutant proteins. The other approach is using homologous recombination to modify embryonic stem cells (ESCs). The modified ESCs are then injected into mouse blastocysts to create mutant mice (knockout mice) (146, 147). Knockout mouse models can be divided into two categories: conventional and conditional knockout. In the conventional knockout model, the target gene is permanent inactivation in all cells of the body. However, this usually causes prenatal death, which makes identification of phenotype difficult. The development of the Cre-LoxP recombination system for conditional gene knockout limits the deletion of a target gene to the desired tissue and has dramatically improved knockout mouse studies (148, 149). Osteoblast-, osteoclast-, chondrocyte-, and osteocyte-specific Cre expression has been created with various promoters to turn on the Cre expression at various differentiation stages and in different cell lineages (See reviews (150) and (72)). Although transgenic mouse models provide a powerful approach to study gene function, the Cre-LoxP recombination system is not perfect. The generation of these models is time and money consuming. Moreover, Cre expression leaking to other cell types and the toxicity of high Cre levels are still the issues that need to be resolved (148, 149, 151).

ENU-induced mutation is one of those approaches that allow genome-wide, high-throughput research (146). ENU is the most potent germ line mutation inducer in rodents (152). ENU treatment of male mice results in random point mutations in the main targets of ENU in the spermatogonial stem cells. By breeding these male mice with normal female mice, the generated F1 population carries unique random ENU-induced mutations. Both genotype-driven and phenotype-driven approaches can be used for screening (146). In addition, the point mutation introduced by ENU resembles the most common form of human genetic variation (146), which makes it a powerful tool to study the specific genes. Large ENU screens have already yielded several mutations in lft-related genes showing skeletal abnormalities.

Gene trapping is another technique that randomly generates loss-of-function mutations,

which takes the middle path between random (such as ENU induced) and molecularly defined mutations (such as knockout mutation) (153). Gene trapping is performed with gene-trap vectors, which usually contain a promoterless reporter and/or selectable marker gene and a downstream polyadenylation sequence. When an expressed gene is inserted into the gene-trap vector, the reporter/selectable marker gene is expressed with the upstream exons, but the gene is prematurely terminated due to the inserted polyadenylation sequence (153, 154). In this way, the gene inactivation and transcriptional activity report can be achieved at the same time. The libraries of ESC lines containing mutations in single genes have been created and about 70 percent of protein-coding genes in the mouse genome have been inactivated in this way (154).

5.2. Mouse models of lft complex A proteins

Currently, there are six known lft-A proteins, which are lft139, lft144, lft122, WDR35, lft143 and lft140. The *in vitro* and *in vivo* studies of these proteins are described in the following sections (Table 2).

5.2.1. lft139

lft139, also termed THM1 (Tetratricopeptide Repeat Containing Hedgehog Modulator 1), is a newly identified ciliary protein (96). The gene encoding this protein is named *TTC21B* (*tetratricopeptide repeat domain 21B*), or *Thm 1* (*tetratricopeptide repeat-containing hedgehog modulator-1*) (96). lft139 contains 1,317 amino acids (about 150 kDa) and is predicted to contain 11 tetratricopeptide repeat (TPR) domains (96). Davis *et al* reported that the *TTC21B* mutation in humans causes JATD and isolated NPHP. By performing *in vitro* and *in vivo* complementation assays, they further found there is a significant enrichment of pathogenic alleles in ciliopathy cases, suggesting that *TTC21B* mutation accounts for 5% of ciliopathies cases (108).

lft139 is highly expressed in the maxillary prominence, branchial arches, limb buds, somites, and spinal cord at embryonic day 10.5. (E10.5). A null mutant of *TTC21B/lft139* named alien (aln) (*lft139^{alien}*) in A/J mice by ENU mutagenesis (95, 96) results in embryonic defects in limbs, eyes, skull, and brain development (95), which are correlated with its expression pattern. *In vivo* and *in vitro* studies showed that lft139 accumulates in cilia and colocalizes with acetylated α -tubulin, which is similar to other lft proteins such as lft88 (96). Cilia in the *lft139^{alien}* limb cells display

short and bulb-like structures in the distal tips. Interestingly, lft88 largely accumulates at this bulb-like structure while undetectable at the proximal ends (96). However, Smo and full-length Gli proteins (key factors of Hh pathway) still can be transported into cilia. Those data suggest that mutation of lft139 only impairs retrograde lft, which leads the accumulation of complex B proteins in the cilia tip and forms the bulb-like structure (96).

Further studies show that Shh pathway was inappropriately activated in *lft139^{alien}* and that overall phenotype of *lft139^{alien}* was very similar to *Rab23* (a negative modulator of Shh signaling) mutants (96). The phenotype of Smo null mice was partially rescued by lft139 deletion. Additionally, Gli2 was overactive in *lft139* mutant, and the double mutant of *Gli2* and *lft139* resembled the phenotype of *Gli2* mutant. Those studies suggested that lft139 is a negative regulator in Shh pathway downstream of Smo and upstream of Gli2 (96). Consistence to this study, Stottmann *et al* reported that Shh signaling was up-regulated in the rostral portion of the *lft139^{alien}* embryo, and the reduction of Shh ligand level (heterozygous for a null allele of Shh) partially rescued the phenotype (95). Besides, the levels of the β -catenin protein were not significantly altered in the mutant mice (95).

5.2.2. lft122

lft122 has a molecular weight of approximately 140 KD and contains seven WD40 domains at the N-terminus (132, 155). The missense mutation of *lft122* is found in CED patients, and the fibroblasts from CED patients had significantly reduced both the numbers of primary cilia in and their length (132). Currently, two lft122 mutant mouse models have been generated (97, 156). One mouse model named lft122-null was generated by deletion of exon 1-3 of DNA repair gene-mediator of RNA polymerase II transcription subunit 1 (*Med1*). Due to the genomic overlap of *lft122* and *Med1*, mutation in lft122-null abrogates the expression of both *Med1* and *lft122*. (97, 157). lft122-null embryos had defects as early as E9.5 and died from E10.5 to E13.5 due to heart defect and diffuse hemorrhagic lesions. Other phenotypes expressed by these mice included exencephaly, rostral neural tube defect, delay in limb development, defects in the ventral portion of the head with altered development of the eyes and branchial arches. Cilia loss was found in the homozygous mutant and malformed cilia were observed in heterozygous embryos, which

Table 2. Current mouse models to study the function of lft-A proteins in bone development

Gene	Mouse models	Mutation	Cilia	Phenotype	Pathways	Ref
lft139	lft139 ^{alien}	ENU mutagenesis, complete loss-of-function	Short and bulb-like structure cilia with accumulation of complex B proteins in the cilia tip	Defects in limbs, eyes, skull and brain development	Overactivated Shh pathway lft139 is the negative regulator in the Shh pathway downstream of Smo and upstream of Gli2, , no significant change in Wnt pathway	(95, 96)
lft122	lft122- null	Exon 1-3 mutation of <i>MED1</i> , null mutation	Cilia loss in the homozygous and malformed cilia in heterozygous	Usually die between E10.5 to E13.5. Exencephaly, delay in limb development, defects of the ventral portion of the head, rostral neural tube, eye and branchial arches	Impaired Shh due to reduced Gli2/Gli3 and Gli3 repressor functions	(97)
	lft122 ^{sopb}	ENU- mutagenesis, complete loss-of-function	Short and swollen cilia with complex B proteins accumulation	Die around E13.5. Neural defects, pre-axial polydactyly, enlarged bronchial arches, and ocular defects	Enhanced Shh with Gli2 and Gli3 accumulation in the cilia tips but absent of TULP3 in cilia	(158)
lft144	lft144 ^{tw}	ENU- mutagenesis, partial loss-of-function	Relatively normal cilia structure but reduced frequency	Die at E11.0. Polydactyly, small rib cages, cleft palate and severe craniofacial anomalies	lft144 ^{tw} MEFs respond weakly to stimulation with upstream Hh agonists, a general increase or ectopic activation of Hh in lft144 ^{tw} embryos, particularly in the facial prominences and limbs	(160)
	lft144 ^{dmhd}	ENU- mutagenesis complete loss-of-function	Short and had highly disrupted axoneme	Die after E13.0. Polydactyly and craniofacial abnormalities. Smaller forelimb paddle, with truncation in the PD and to a lesser extent the AP axes	Less activation of the Shh pathway in the caudal neural tube	(160, 161)
	lft144 ^{tw} lft122 ^{sopb}	Double mutation	Similar to lft144 ^{dmhd}	Similar to lft144 ^{dmhd}	Less Shh pathway activity	(161)
	lft144 ^{tw} Dync2h1 ^{mmi}	Double mutation	Reduced cilia number, cilia swollen in Dync2h1 ^{mmi}	Partially rescued most ventral neural cell type, and the floor plate	Partial rescue of Shh signaling	(161)
WDR35 (lft121)	WDR35 ^{yet}	ENU- mutagenesis, complete loss-of-function	Cilia loss with lft88 accumulated in the basal body	Usually die at E12.5. Severe defects in cardiovascular, embryo turning, and limb outgrowth and patterning. Failure of the somite derivatives	Not reported	(34)
	WDR ^{tm2a}	“targeted trap” null allele	Not reported	Phenocopies WDR35 ^{yet}	Not reported	(34)

(Contd..)

Table 2. Contd...

Gene	Mouse models	Mutation	Cilia	Phenotype	Pathways	Ref
<i>lft140</i>	<i>lft140^{null}</i>	Germline deletion with C57Bl/6 Prm-Cre	Not reported	Die at mid-gestation	Not reported	(162)
	<i>lft140^{cauli}</i>	ENU-mutagenesis, “possibly damaging” mutation	Cilia morphology was severely disrupted with broader and bulbous appearance	Start to die at E13.5. Exencephaly, anophthalmia, severely disorganised ribs with extensive exostoses, vertebral and palatal defects, agenesis/hypoplasia of the craniofacial skeleton, and polydactyly of the hindlimbs	Ectopic Hh signaling activation and reduced Gli3 repressor production.	(163, 164)
	<i>lft140^{-/-}</i>	Germline deletion with CMV-Cre	Not reported	Similar to <i>lft140^{cauli}</i>	Not reported	(164)

associated with defects in left–right asymmetry. Additionally, mutation of *lft22* reduced Gli2/Gli3 and Gli3 repressor functions, suggesting that Shh signaling is impaired (97).

Another mouse model *lft122^{sopb}* was generated with ENU mutagenesis and named *sister of open brain (sopb)* due to the similar phenotype to *Rab23/open brain* mutants (158). *Sopb* is a null allele of *lft122* with a T-to-C transition mutation in the start codon of *lft122*. *lft122^{sopb}* embryos died around E13.5, displayed neural defects, pre-axial polydactyly, enlarged bronchial arches, and ocular defects. Cilia frequency was normal in the neural tube of *lft122^{sopb}* mice but reduced in *lft122^{sopb}* MEFs. Cilia in *lft122^{sopb}* were shorter but swollen along their lengths and tips, which accumulated with complex B proteins as stained with lft88 and lft57. The complex A protein lft140 was undetectable in the cilia of *lft122^{sopb}* cells, suggesting a role of lft122 in facilitating complex A stability and retrograde lft (158). Suppressor of Fused (SUFU) and Smo transportation to cilia were not affected while Gli2 and Gli3 accumulated in the cilia tips after disruption of lft122. Ciliary localization of TULP3, a repressor of Shh pathways, was also blocked in *lft122^{sopb}*. These data suggested that lft122 negatively regulates the Shh pathway downstream of Smo through differentially controlling the ciliary localization of the Hh pathway regulators (158). The different phenotypes observed in those two models may be caused by different mouse strain backgrounds (*lft122^{sopb}* of C3Heb/FeJ and lft122-null of C57BL/6, respectively) and/

or by the effect of MED1 deletion in lft122-null mutant (158).

5.2.3. lft144

lft144, coded by the gene *WDR19* (*WD repeat domain 19*), has WD40 regions and TPR protein–protein interaction domains (159). *lft144* localizes at the base and tip of the cilium in a punctate pattern along the axoneme in the mouse embryo fibroblasts (MEFs) (160). Mutation of *lft144* causes CED. The fibroblasts from the CED patients show undetectable *lft144* and the reduction in cilia abundance and length (109).

Currently, two ENU induced mouse mutant models of *lft144* (*lft144^{tw}* and *lft144^{dmhd}*) have been generated (160, 161). Mice with the *Twinkle-toes* mutation (*lft144^{tw}*), which introduces a leucine-to-proline substitution at amino acid position 750 (L750P, within the first TPR repeat), are partial loss-of-function mutations. Similar to the phenotype of CED patients, *lft144^{tw}* embryos die at E11.0 and display polydactyly, small rib cage, and severe craniofacial anomalies. The number of ciliated cells was reduced in *lft144^{tw}* mouse limbs and isolated MEFs, while the cilia structure appeared relatively normal. In addition, *lft144^{tw}* cells and tissues have enhanced ligand-independent Hh signaling, but reduced response to Hh ligands.

A more severe allele of the *lft144* mutation is known as *diamondhead* (*lft144^{dmhd}*), which represents a functional null mutation (161). These mutant embryos displayed different phenotypes

when compared to *lft144^{tw}* mice. The cilia were shorter and showed highly disrupted axoneme in *lft144^{dmhd}* embryos. Unlike *lft144^{tw}*, *lft122^{sopb}* and *lft139^{alien}*, *lft144^{dmhd}* had less activity in Shh signaling, but greater numbers of cells with an intermediate level of Shh activity in the caudal neural tube (161). *lft144^{dmhd}* showed smaller forelimb paddle, with truncation in the proximal-distal (PD), and to a lesser extent, the anterior-posterior (AP) axes when compared to those *lft144^{tw}* mice revealed a dose-dependent effect of *lft144* on limb outgrowth (160).

The cilia in double mutation of *lft144^{tw}* and *lft122^{sopb}* (*lft144^{tw}lft122^{sopb}*) were shorter and more bulbous cilia than those in either single mutation and were similar to those in *lft144^{dmhd}* embryos (161). Also the phenotype of *lft144^{tw}lft122^{sopb}* is close to that in *lft144^{dmhd}*, suggesting that both *lft144^{tw}* *lft122^{sopb}* and *lft144^{dmhd}* result in a stronger disruption of the IFT-A complex. Double mutations of *lft144^{tw}* and *Dync2h1^{mmi}* (*lft144^{tw}Dync2h1^{mmi}*) partially rescued the axoneme structure and reduced the level of cilia swollen in *Dync2h1^{mmi}* (161). The authors concluded that *lft144* is also required for normal anterograde trafficking (161). Further studies on the Hh pathway showed that transportation of the Shh pathway proteins Gli2, SUFU, and Kif7 were normal in *lft144^{tw}*, *lft144^{tw}lft122^{sopb}* and *lft144^{dmhd}*. But the membrane proteins Arl13b, adenylyl cyclase III (ACIII), and Smo failed to localize to primary cilia in the more complete lft-A mutation (*lft144^{tw}lft122^{sopb}* and *lft144^{dmhd}*). Reduced ACIII level in cilia led to the decrease in cAMP and protein kinase A (PKA) activity, which leads to the ligand independent activation of Shh. This can explain the up-regulated Shh observed in *lft144^{tw}*, but loss of Shh activity in *lft144^{dmhd}* with the null mutation of *lft144* due to severe disruptions of the cilia structure and membrane protein trafficking (161).

5.2.4. WDR35 (lft121)

WDR35 (WR repeat domain 35), which is orthologous to *lft121* in mammals, has seven closely spaced WD40 repeats at the N-terminal and a TRP-like motif at the C-terminus (34). WD40 repeats are believed to have the functions of intracellular trafficking, cargo recognition, and binding. Mutation of this structure caused SRP with a ciliogenesis defect (34). Mutations in the C-terminal domains are more likely associated with CED and result in impairment, but not in complete inhibition of retrograde transport (34, 136).

Mill *et al* isolated a WDR35 null mutant mouse line *yeti* (WDR35^{yet}) from a recessive ENU mutagenesis screen that was looking for genetic effects on embryogenesis (34). They found that a single G>A mutation in the splice acceptor site of exon 22 of *WDR35* resulted in nonsense-mediated decay of mutant transcripts and complete loss of function (34). WDR35^{yet} embryos died at E12.5 and exhibited severe defects in cardiovascular, embryo turning, and limb outgrowth and patterning. Additionally, WDR35^{yet} displayed failure of the somite derivatives, including migration and differentiation of putative ribs (34).

WDR35 accumulates in the basal body and along the cilia axoneme. Both WDR35^{yet} MEFs and fibroblasts from SRP patient (WDR35^{r5/r5}) had cilia loss and less *lft88* accumulation around the basal body area (34). Interestingly, fibroblasts from CED patient (WDR35 mutation) displayed shorter cilia and accumulation of *lft88* and *lft57* in the distal ends of cilia (135). Those data suggest that *WDR35* mutation in SRP may interfere with anterograde transportation or other vesicular trafficking, which leads to the cilia defect, while the *WDR35* mutation in CED only affects retrograde transport and results in a typical lft-A mutation phenotype like the *lft122* mutation (34). Mill *et al* also described WDR^{tm2a} mice as an embryonic stem cell derived “targeted trap” null mutation (34). The loss of WDR35 by targeted mutation phenocopies the WDR35^{yet}, but the details have not yet been revealed (34).

5.2.5. lft43

lft43 is the smallest member in lft-A complex and it does not contain any identifiable domains (28). *lft43* was the third gene identified as mutation in patients of CED with a homozygote N-terminal deletion (135). A recent study suggested that there is the direct interaction between *lft43* and *lft121*, and *lft121* is another gene responsible for CED (90). Fibroblasts from an *lft43* mutant CED patient had shorter cilia and an accumulation of complex B proteins in the distal ends of cilia indicated by *lft88* and *lft57* staining, which is similar to other lft-A mutation (135). No *lft43* mutant mouse model has been published yet.

5.2.6. lft140

The structure of *lft140* is similar to *lft144* (28). Mutation of *lft140* has been correlated with MSS and JATD (111). Fibroblasts from some MSS patients had fewer ciliated cells and a normal distribution of *lft140*, but altered distributions of *lft88*

and lft46 (evenly distributed along the cilium instead of localizing in the base and tip), suggesting a defect in retrograde ciliary transport (111).

Jonassen *et al* generated lft140^{flox/flox} mice with two Loxp sites flanking exon 7 of *lft140* (162). lft140^{null} was generated by crossing these mice with C57Bl/6 Prm-Cre (Cre recombinase expressed in the germline). These mice died at mid-gestation, but the phenotype has not been described yet (162). The function of lft140 in kidney development has been studied by crossing lft140^{flox/flox} mice with HoxB7-Cre to delete *lft140* in the collecting ducts. Loss of lft140 from the collecting ducts disrupted cilia assembly and caused renal cystic disease. Significant increases in tissue fibrosis and the expression of canonical Wnt pathway genes and Hh mediators was also found in highly cystic, but not precystic, kidneys in these mice, suggesting that abnormalities in these different pathways may influence cyst progression (162).

Another lft140 mouse model was identified from a comprehensive phenotype-driven ENU screen and was named *cauliflower* (*cauli*) (163, 164). *Cauli* mutation is a recessive missense mutation with a T-to-A substitution in exon 19 at position 2564 (leading to an isoleucine to lysine amino acid change at position 855, p.I855K). lft140^{cauli} embryos died at E13.5 and exhibited exencephaly, anophthalmia, severely disorganized ribs with extensive exostoses, vertebral and palatal defects, agenesis/hypoplasia of the craniofacial skeleton, and polydactyly of the hindlimbs (163, 164). Cilia morphology in lft140^{cauli} limb buds was severely disrupted with a broader and bulbous appearance. Similar to other lft-A mutants, ectopic Hh signaling activity has been observed in lft140^{cauli} embryos. The loss of lft140 resulted in reduced Gli3 repressor production, and thus led to the polydactylous phenotype and rib exostoses (164). The null mutation of *lft140* (lft140^{-/-}) was also generated by Miller *et al* by crossing the lft140^{flox/flox} mice with CMV-Cre mice (Cre recombinase is expressed in all tissues including germ cells) (164). lft140^{-/-} embryos died at about E11.5 with the identical phenotype to that observed in lft140^{cauli} (164).

5.3. Mouse models of lft complex B proteins

lft complex B contains 14 known proteins named lft20, lft22, lft25, lft27, lft46, lft52, lft54, lft57, lft70, lft74/lft72, lft80, lft81, lft88, and lft172. *In vitro* and *in vivo* studies examining these proteins are described in the following sections (Table 3).

5.3.1. lft80

lft80 is a 777-residue protein, which contains seven WD40 domains (165) and is the first lft protein that was identified with mutation in human diseases -JATD and SRP type III, (130). Knockdown of lft80 in zebrafish resulted in cystic kidneys with down-regulated Ptch1, suggesting reduced Hh activity (165). Rix *et al* generated the mouse model named lft80^{gt/gt} using gene trap strategy (insert gene-trap cassette pGT0lrx in the intronic region downstream of exon 9 in lft80) (70). The homozygotes lft80^{gt/gt} are hypomorphs with the low level of wild-type transcript production, but most died perinatally. Two percent of homozygotes survived to postnatal stages and had severe growth retardation with short long bones, a narrow rib cage and polydactyly. MEFs from lft80^{gt/gt} E14.5 embryos had normal cilia but reduced Hh activity in response to smoothened agonist purmorphamine. These data suggested that a low level of lft80 expression is enough for ciliogenesis, but not sufficient for Hh signaling transduction (70).

The results from our laboratory showed that lft80 was highly expressed in mouse long bone and skull, as well as during osteoblast and chondrocyte differentiation (166, 167). We used RNA interference (RNAi) to silence *lft80* in the murine mesenchymal progenitor cell line C3H10T1/2 and bone marrow derived stromal cells (BMSCs) and found that loss of lft80 led to shorten or loss of cilia and significantly blocked osteogenesis. Gli2 expression was significantly decreased in lft80-silenced cells, suggesting the impaired Hh pathway activity (167). Silence of lft80 in BMSCs also impaired the chondrogenic differentiation by down-regulating Hh signaling, whereas up-regulating Wnt signaling (166). Those data suggested the important role of lft80 in skeletal development. Currently, we have generated lft80^{flox/flox} conditional knockout mouse alleles, and are identifying the role and mechanism of lft80 in different bone cells by using specific bone cell Cre transgenic lines to delete lft80 in various bone cells.

5.3.2. lft88

lft88 (Tg737 or Polaris), a TPR-containing protein, was first identified in ARPKD (168). The lft88^{Tg737Rpw} (also known as ORPK, FVB/N strain) mouse model was generated 20 years ago as a model for human recessive polycystic kidney disease (168). lft88^{Tg737Rpw} has a transgene insertion-induced hypomorphic allele of lft88 (continued expression

Table 3. Current mouse models to study the function of Ift-B proteins in bone development

Gene	Mouse models	Mutation	Cilia	Phenotype	Pathways	Ref
Ift80	Ift80 ^{gt/gt}	Gene trap, hypomorph with low level wild-type transcript production	Normal cilia	Usually die perinatally Severe growth retardation with short long bones, narrow ribcage and polydactyly	Less Hh activity	(70)
Ift88	Ift88 ^{Tg737Rpw} (ORPK)	Transgene insertion-induced hypomorphic allele	Stunted and malformed cilia	Survive to young adulthood Scruffy fur, severe growth retardation, cystic renal phenotype, preaxial polydactyly, defects in skeletal patterning and growth with craniofacial abnormalities, cleft palate, unfused tibia and fibula, and significantly smaller growth plates	Defects in PDGFR- α / PDGF-A signaling	(168-171)
	Ift88 ^{Δ2-3βGal} (Ift88 ^{tm1Rpw})	A Null mutation	Cilia loss	Arrested at mid-gestation with neural tube defects, enlargement of the pericardial sac and left-right asymmetry defects	Reduced expression of <i>Shh</i> and <i>Hnf3β</i> in midline No changes in <i>Shh</i> or its downstream genes in limb bud	(91, 170)
	Ift88 ^{fxo} (Ift88 ^{hypo})	ENU-mutagenesis, null mutation	Lack nodal cilia	Dies at E12.5. Sharp angle of the mesencephalic flexure, abnormal brain morphology and preaxial polydactyly	Disrupted Hh signaling with loss of both Gli activator function and Gli repressor function in the limbs	(56, 173)
	Ift80 ^{flox/flox} , Pr \times 1-Cre	Deletion in limb mesenchyme	Cilia loss	Severe polydactyly and defects in endochondral bone formation	Aberrant Shh and Ihh signaling	(174)
	Ift80 ^{flox/flox} , Ms \times 2-Cre	Deletion in limb ectoderm	Cilia loss	No overt effect on limb patterning	Not reported	(174)
	Ift80 ^{flox/flox} , Col2 α -Cre	Deletion in committed chondrocyte	Cilia loss	Postnatal dwarfism and premature loss of the growth plate with down-regulated proliferation and up-regulated hypertrophic differentiation of the chondrocytes. Thicker articular cartilage with increased cell density and OA symptoms	Reduced Hh but elevated Wnt/ β -catenin (reduced Sfrp5) signaling. Upregulated Hh signaling in articular cartilage with reduced Gli3 repressor/ Gli3 activator	(93) (176) (175)
Ift172	Ift172 ^{wim}	ENU-mutagenesis, a null mutation	Lack nodal cilia	Arrested at E10.5-11.5. Open neural tube in the head that lacked the normal groove on the ventral midline.	Strong defect in Shh signaling Normal Wnt signaling	(56) (178)
	Ift172 ^{avc1}	ENU-mutagenesis, partial loss of function	Defects in cilia	Die at birth. Anomalies in vertebral, tracheoesophageal and limb, defects in cardiac, anal atresia, renal dysplasia and hydrocephalous	Defects in Hh signaling with reduced Gli3 processing and Gli2 accumulation	(92)
	Ift172 ^{Sib}	Conventional knockout, a null mutation	Short and without visible microtubules	Die between E12.5 and 13.0. Severe craniofacial malformations, defects in brain patterning, failure to close the neural tube and exencephaly	<i>Shh</i> and <i>Gli1</i> decreased in the developing forebrain and midbrain	(179)

(Contd..)

Table 3. Contd...

Gene	Mouse models	Mutation	Cilia	Phenotype	Pathways	Ref
	<i>lft172^{Δ/Δ}</i>	CMV-Cre-mediated germline deletion	Not reported	Embryonic lethality. Neural tube defects (Similar to <i>lft172^{wim}</i> and <i>172^{Sib}</i>)	Not reported	(180)
	<i>lft172^{flt}</i> ; <i>Prx1-Cre</i>	One allele is germline deletion and another is only deleted in the developing limb bud mesenchyme	Not reported	Eight digits on each forelimb and a single extra digit on each hind limb	Not reported	(180)
<i>lft20</i>	<i>lft20^{Flox}</i> ; <i>Prm-Cre</i>	Germline deletion	Not reported	Die before birth	Not reported	(183)
<i>lft25</i>	<i>lft25^{neo}</i> and <i>lft25^{null1}</i>	Null mutation	Normal cilia	Die at birth. Omphaloceles, polydactyly with a preaxial digit duplication, micrognathia, cleft palate and malalignment of the sternal vertebrae	Ciliary accumulation of Shh components and attenuated Shh	(185)
<i>Ngd5</i> (<i>lft52</i>)	<i>lft52^{hypo}</i>	Partial loss-of-function mutations	Not reported	Tight mesencephalic flexure, left-right and ventral midline defects, polydactyly and craniofacial defect	Defects in Hh pathway in limb buds	(173)
<i>Traf3ip1/MIP-T3</i> (<i>lft54</i>)	<i>Traf3ip1^{GT}</i>	Gene Trap	Cilia disrupt	Die before E13.5. Neural developmental defects, cardiac edema and polydactyly	Defects in Hh pathway. Unchanged Wnt pathway. Elevated mTor pathway	(193)
<i>Hippi</i> (<i>lft57</i>)	<i>Hippi^{-/-}</i>	Conventional knockout	Defects in nodal cilia	Die predominantly prior to E10.5. Defects in left-right patterning and nervous system development	Down-regulated Shh pathway	(196)

of alternatively spliced transcripts and presence of low levels *lft88*), which allows homozygous mutant mice to survive into young adulthood. *lft88^{Tg737Rpw}* mice have scruffy fur, severe growth retardation, cystic renal phenotype, and preaxial polydactyly in all limbs (168, 169). *lft88^{Tg737Rpw}* mice also have defects in skeletal patterning and growth with craniofacial abnormalities, cleft palate, unfused tibia and fibula (170). The cilia are not completely abolished in *lft88^{Tg737Rpw}* but are stunted and malformed (169). Fibroblasts isolated from *lft88^{Tg737Rpw}* mice do not respond to platelet-derived growth factor A (PDGF-A) and fail to conduct chemotaxis toward PDGF-A (169). The growth plates of *lft88^{Tg737Rpw}* mice are significantly smaller in length and width, suggesting the role of *lft88*/cilia in chondrocyte differentiation and hypertrophy (171). Chondrocytes from *lft88^{Tg737Rpw}* mice have a defect in the compression-induced Ca^{2+} signaling but not ATP release, suggesting that cilia are involved in ATP reception instead of initiation of mechanotransduction (172).

To further study the function of *lft88*, Murcia *et al* (91) generated a null allele (*Tg737^{Δ2-3βGal}* or *lft88^{tm1Rpw}*, FVB/N strain) by homologous recombination in ESCs. The initial coding exons (exon 2 and part of exon 3) were replaced with the β-galactosidase (β-Gal) reporter gene. *Tg737^{Δ2-3βGal}* embryos that are arrested in development at mid-gestation exhibit neural tube defects, enlargement of the pericardial sac, and significant left-right asymmetry defects. Cilia were absent in *Tg737^{Δ2-3βGal}* ventral node cells. Expression of both *Shh* and *Hnf3β* is reduced in the midline in E8.0 *lft88^{Δ2-3βGal}* embryos, contributing to the alterations in midline development (91). However, Zhang *et al* found that no changes in Shh or its downstream genes occurred in the limb buds of *Tg737^{Δ2-3βGal}* embryos (170).

Huangfu *et al* identified *lft88^{fxo}* (*Flexo*, also called *lft88^{hypo}*) from the screen of ENU-induced embryonic abnormalities (56). *lft88^{fxo}* is a null allele of *lft88* with deletion of exon 16 (including part of

one of the ten predicted TRP motif). *Ift88*^{fxo} embryos lack nodal cilia and display an unusually sharp angle of the mesencephalic flexure at E9.5-10.5. Those embryos are arrested at E12.5-13.5 with abnormal brain morphology and preaxial polydactyly (56, 173). Liu *et al* studied the Hh signaling in the limbs of *Ift88*^{fxo} embryos and proposed that both Gli activator function (down-regulation of *Ptch1* and *Gli1* expression) and Gli repressor function (*Hand2* misexpression and polydactyly) are compromised in the limbs of *Ift88*^{fxo} (173).

Haycraft *et al* generated *Ift88*^{lox/lox} conditional allele with LoxP sites flanking exons 4-6 (174). The deletion of these exons with Cre-line is predicted to result in a translational frame shift and loss of all *Ift88* function (a null mutation). *Prx1*-Cre (*Ift88*^{lox/lox}; *Prx1*-Cre, limb mesenchyme with both chondrocytes and perichondrium) and *Msx2*-Cre (*Ift88*^{lox/lox}; *Msx2*-Cre, limb ectoderm) were used to delete *Ift88* on different cell populations within the developing limb (174). Disrupted *Ift88* using both Cre lines caused cilia loss (93, 174). *Ift88*^{lox/lox}; *Msx2*-Cre mice did not exhibit an overt effect on limb patterning. However, *Ift88*^{lox/lox}; *Prx1*-Cre displayed severe polydactyly and defects in endochondral bone formation with aberrant Shh and Ihh signaling (174). Song *et al* crossed *Ift88*^{lox/lox} mice with *Col2a1*-Cre to delete *Ift88* in committed chondrocyte lineage (*Ift88*^{lox/lox}; *Col2a1*-Cre) (93). The phenotype of *Ift88*^{lox/lox}; *Col2a1*-Cre is similar to *Kif3a*^{lox/lox}; *Col2a*-Cre with postnatal dwarfism and complete loss of the growth plate, resulting from down-regulated the proliferation and up-regulated hypertrophic differentiation of the chondrocytes. They further demonstrated that *Ift88* regulated chondrocyte rotation to maintain the columnar organization of the growth plate (93). Chang *et al* studied the signaling pathways involved in the growth plate abnormality by comparing gene expression profiles in normal and *Ift88*^{lox/lox}; *Col2a*-Cre growth plates (175). They found that the expression of Hh and secreted frizzled-related sequence protein 5 (*Sfrp5*) (an extracellular antagonist of Wnt signaling pathways) was significantly lower in *Ift88*^{lox/lox}; *Col2a*-Cre. These data support the increase of Wnt/ β -catenin signaling in flat columnar cells of the growth plate in *Ift88*^{lox/lox}; *Col2a*-Cre as measured by expression of *Axin2* and *Lef1*, as well as nuclear localization of β -catenin (175). In addition, *Ift88*^{lox/lox}; *Col2a*-Cre mice showed thicker articular cartilage with increased cell density, which probably results from the reduced apoptosis rate in cartilage remodeling (176). The mechanical properties of cartilage from *Ift88*^{lox/lox};

Col2a-Cre mice significantly decreased, particularly in the deeper zones (172). Up-regulation of Hh signaling as measured by expression of *Ptch1* and *Gli1* was observed in *Ift88*^{lox/lox}; *Col2a*-Cre articular cartilage, which contributes to the osteoarthritis (OA) symptoms. A reduction in the Gli3 repressor to activator ratio also occurred, which is an explanation for the elevated Hh signaling (176).

5.3.3. *Ift172*

Ift172 is the largest lft protein. It is predicted to have N-terminal WD-40 repeats and C-terminal α -helical TPRs (28).

Huangfu *et al* identified a *wimble* (*wim*) mutation of *Ift172*, a null mutation, in a screen for recessive ENU-induced mutations (56). The T to C mutation in the *wim* allele causes a leucine to proline substitution (Leu1564Pro) near the C-terminus. *Ift172*^{wim} embryos have morphological abnormalities at E9.5 and arrest at E10.5-11.5 due to strong defect in Shh signaling. *Ift172*^{wim} mice lack nodal cilia and have an open neural tube in the head, which does not have the normal groove on the ventral midline (56). *Ptch1* mutant mice (177) (partial exon 1, and exon 2 of *ptch1* were replaced with lacZ and a neomycin (neo) resistance gene through homologous recombination in ESCs) were crossed with *Ift172*^{wim} to generate double mutants, which showed the similar phenotype as *Ift172*^{wim}. Loss of *Ptch1* caused the activation of Hh pathway, but *Ift172* mutation blocked this activation, suggesting *Ift172* acts downstream of *Ptch1* in Shh signaling pathway (56). The loss of cilia did not affect the response of the *Ift172*^{wim} mid-gestation embryos or MEFs to Wnt ligands (178).

Friedland-Little *et al* identified *Ift172*^{avc1} from a screen of ENU-induced mutations (92). This mutation is an A-to-G transition in the splice donor site downstream of exon 24 of *Ift172* to alter *Ift172* splicing. *Ift172*^{avc1} allele maintains some wild-type transcript, which is a partial loss of function mutation and dies at birth. *Ift172*^{avc1} mice show anomalies in vertebral, tracheoesophageal and limb, and defects in cardiac, anal atresia, renal dysplasia and hydrocephalous. But left-right axis formation is normal in *Ift172*^{avc1} (92). *Ift172*^{avc1} cells have truncated cilia (including disrupted IFT88 localization) both *in vivo* and *in vitro*. *Ift172*^{avc1} embryos display a global reduction in the expression of *Ptch1* and *Gli1*. A 70.7% reduction of Gli3 processing (from full-length to transcriptional repressor) in *Ift172*^{avc1} embryos and disrupted Hh-dependent Gli2 localization in *Ift172*^{avc1}

MEFs has been observed, which contributes to the Hh signaling defect in *lft172^{avc1}*. Quantitative analysis wild-type, *lft172^{avc1/avc1}* and *lft172^{wim/avc1}* revealed the dose-dependent requirement for *lft172* in ciliogenesis and Hh signaling (92).

Another mouse model named *Slb* mutant (*lft172^{Slb}*) was generated by Gorivodsky *et al* (179). A *neo* cassette was inserted to replace the first three exons of *lft172* and cause the loss of function mutation. *lft172^{Slb}* mice die between E12.5 and E13.0 with severe craniofacial malformations, defects in brain patterning, failure to close neural tube and exencephaly. Cilia are initiated but shorter and without visible microtubules. Expression of *Shh* and *Gli1* is lower in the developing forebrain and midbrain of *lft172^{Slb}*. In addition, *lft172* is required for early regulation of *Fgf8* at the midbrain-hindbrain boundary (179).

To further study the role of *lft172* in the later stage of development, Howard *et al* generated mice with a conditional knockout of *lft172* (*lft172^{fl/fl}*) using LoxP sites flanking exons 2-3 (180). Deletion with the Cre line can remove about a third of a WD40 repeat domain and cause a frame shift, resulting in a stop codon. Similar to *lft172^{wim}* and *lft172^{Slb}*, germline deletion with CMV-Cre (*lft172^{Δ/Δ}*) leads to embryonic lethality due to neural tube defects. To examine tissue-specific function of *lft172*, *Prx1*-Cre was used to delete *lft172* in the developing limb. *lft172^{fl/fl}; Prx1*-Cre mice (one allele is a germ line deletion and another is only deleted in the developing limb bud mesenchyme) have eight digits on each forelimb and a single extra digit on each hind limb (180).

5.3.4. *lft20*

lft20 is the smallest lft protein in the lft family and has several unique features. It is the only lft protein that localizes to the Golgi complex in mammalian cells, suggesting its role in the ciliary protein sorting (165, 166). *lft20* is anchored to the Golgi complex by the Golgi protein-Golgi Microtubule Associated Protein 210 (GMAP210)/thyroid hormone receptor interacting protein 11 (TRIP11) (166). Without GMAP210, *lft20* is no longer localized to the Golgi structure and cilia are shorter with reduced ciliary localization of membrane protein polycystin-2. GMAP210 mutant mice die at birth and display growth restriction, ventricular septal defects of the heart, omphalocele, and lung hypoplasia (166). Strong knockdown of *lft20* in mammalian cells blocks cilia formation while moderate knockdown does not block cilia assembly, but reduces the

amount of polycystin-2 to localize to the cilia (165). Those data demonstrated GMAP210 and *lft20* work together at Golgi in sorting or transporting proteins like polycystin-2. Besides GMAP210/ TRIP11, *lft20* also exhibits a strong interaction with *lft57*/Hippi and kinesin II subunit, *Kif3b* (181). *lft20* is tethered by *lft57* to the lft particle, physically interacts with *Kif3b* and mediates the ATP-dependent dissociation of kinesin. Loss of *lft57* leads to the disassociation of *lft20* from lft particles but does not affect the association of kinesin to the lft particle (182).

In addition, *lft20* is expressed in lymphoid and myeloid cells, which do not have primary cilia. *lft20* takes part in the polarized recycling of the TCR/ CD3 complex, which is the first connection of lft proteins and membrane trafficking in cells without cilia (167). But whether *lft20* functions in membrane trafficking in the ciliated cells remains unknown.

The mouse floxed allele of *lft20* (*lft20^{Flox}*), which has LoxP sites flanking exons 2 and 3 of *lft20*, has been established by Jonassen *et al* (183). Germline deletion of *lft20* using *Prm*-Cre led to embryonic lethality (183). So far, *HoxB7*-Cre and human red/green pigment gene promoter (HRGP)-Cre have been used to study the function of *lft20* in kidney and photoreceptor cells (183, 184). But the function of *lft20* has yet to be reported in bone, and our lab is currently trying to understand the function and mechanism of *lft20* in bone development and remodeling.

5.3.5. *lft25*

lft25 is one of the two lft proteins (*lft25* and *lft27*) that are absent from *C. elegans* and *Drosophila*, suggesting a role of *lft25* beyond ciliogenesis (185). Keady *et al* generated the *lft25^{neo}* mice by using homologous recombination to modify ESCs containing a splice acceptor, a promoterless β-galactosidase-neo resistance gene fusion and LoxP sites flanked exon 3 (185). By crossing these mice with the germ line Cre *Prm*-Cre mice, they generated *lft25^{null1}*, which lacks exon 3 of *lft25*. Both *lft25^{neo}* and *lft25^{null1}* are null mutations without *lft25* expression. These mice die at birth with severe heart defects but have normal cilia. These are different from other lft-B null mutants that usually fail to form cilia and die around mid-gestation. In addition, mutation of *lft25* leads to omphaloceles, polydactyly with a preaxial digit duplication, micrognathia, cleft palate, and malalignment of the sternal vertebrae. *lft25* is needed for the exit of *Smo* and *Ptch1* from cilia (not import) and *Gli2* accumulation at the ciliary

tip in response to activation of the Shh pathway. Loss of *Ift25* leads to a failure to activate the Hh pathway upon stimulation (185).

5.3.6. *Ift27*

Ift27 belongs to the small G protein family (close to Rab subfamily) with Rab-like small GTPase activity (186, 187). It binds weakly with GDP and GTP, and displays very low, but significant intrinsic GTPase activity (186). In addition, it forms a sub-complex with *Ift25* in green algae, *Chlamydomonas reinhardtii*, mice and humans (186, 188). *Ift25* and *Ift27* seem to be preassembled before entering the cilia and associating with other Ift-B members, suggesting the role of *Ift25/27* in Ift initiation (186, 188). However, the roles of the *Ift27* in cilia formation and bone development are not yet known and no *Ift27* mutant mouse model has been published yet.

5.3.7. *RabL5 (Ift22)*

Ift22, another Ift protein with significant sequence similarity to Rab-like GTPases, is known as Rab-like 5 (*RabL5*) in mammals (28). Mutation of the *Ift22* ortholog in *C.elegans* (*Rab-like 5*) did not alter the cilia (189). But knock-down of *Ift22* by RNAi in the trypanosome flagellum surprisingly resulted in short flagella filled with a high amount of IFT proteins, which is a typical retrograde Ift defect phenotype (190). No mouse model or related human diseases have been reported yet.

5.3.8. *Ift46*

Even though *Ift46* does not contain any recognizable domains or motifs, it directly interacts with *Ift52*, *Ift70* and *Ift88* in the core of Ift-B (28). Study showed that *Ift46* is necessary for Ift-B stability (191). Recently, *Ift46* has been identified as a BMP-2 sensitive protein in mouse chondrocytes (192). *Ift46* is located in the mouse primary cilium of chondrocytes and is preferentially expressed in early hypertrophic chondrocytes in the growth plate. Knockdown of *Ift46* in cultured chondrocytes by small interfering RNA (siRNA) oligonucleotides does not affect the expression of the *Col2a1*, *Col10a1*, BMP receptors, and Smads. Instead, it up-regulates *Msx1*, *fibroblast growth factor receptor-1 (Fgfr1)*, *type XII collagen (Col12a1)*, *biglycan (Bgn)*, *matrix metalloproteinase-10 (Mmp10)*, and *heat shock protein 47 (Hsp47)* expression (192). Antisense morpholino oligonucleotides were injected into *Danio rerio* embryos to deplete zebrafish *Ift46*, which led to severe defects in embryonic development. These data highlight the role of *Ift46* in chondrocyte function and vertebrates development (192).

However, whether *Ift46* is involved in cilia formation in chondrocytes and how it affects bone development and remodeling have not been evaluated yet.

5.3.9. *Ngd5 (Ift52)*

Ift52 is predicted to have an N-terminal 'Gltf'-domain (GldG/Ift) and proline-rich stretch (28). *Ift52* can bind *Ift70* and *Ift88* directly through the proline-rich stretch (28). Mouse *Ngd5* is the mammalian ortholog of *Ift52* (181). Liu *et al* generated *Ift52^{hypo}* from two mutant ESC lines in which the *Ngd5* gene was disrupted by replacing it with the lacZ gene (173). The C-terminal 99 amino acids of the *Ngd5* protein were replaced with lacZ in the first line (XL826). In the second allele (RRJ295), the last 47 amino acids of the *Ngd5* protein were replaced with lacZ. Mice derived from both lines are termed as *Ift52^{hypo}*. They are partial loss-of-function mutations and have similar phenotypes, including tight mesencephalic flexure, left-right and ventral midline defects, polydactyly and craniofacial defects at embryonic stage (173). In addition, *ptch1* is not detected in the limb buds of *Ift52^{hypo}*, suggesting the disruption of Hh signaling (173).

5.3.10. *Traf3ip1 (Ift54)*

The homology of *Ift54* in mammalian systems is tumor necrosis factor alpha receptor 3 interacting protein 1 (*Traf3ip1*)/microtubule interacting protein that associates with *Traf3* (MIP-T3, hereafter called *Traf3ip1*), which was initially characterized through its interactions with tubulin, actin, TNFR-associated factor-3 (*Traf3*), interleukin-13 receptor $\alpha 1$ (IL-13R1), and Disrupted-In-Schizophrenia 1 (DISC1). *Traf3ip1* inhibits IL-13-mediated phosphorylation of signal transducer and activator of transcription-6 (STAT6). *Traf3ip1* also regulates *Traf3* and DISC1 by sequestration of these factors to the cytoskeleton in mammals (193, 194).

Berbari *et al* generated the *Traf3ip1^{GT}* mouse model with embryonic stem cell line harboring a gene trap insertion in intron 11 of *Traf3ip1* (193). MEFs from *Traf3ip1^{GT}* had nearly 90% less transcript levels than the level in wild-type MEFs. *Traf3ip1^{GT}* mouse is lethal at the embryonic stage before E13.5, having neural developmental defects, cardiac edema, and polydactyly. Hh signaling was dramatically reduced in both the neural and endoderm tubes in E10.5 *Traf3ip1^{GT}* embryos as determined by crossing with *Ptch1-lacZ* Hh reporter mice. By crossing those mice with BAT-Gal Wnt reporter mouse line, the author found that there is no overt alternation in canonical Wnt signaling in the

Traf3ip1^{GT} mutant embryos. MEFs from Traf3ip1^{GT} embryos have defective cilia formation, up-regulated acetylated microtubules, and increased cell size. Interestingly, deletion of Traf3ip1 increases the activity of the mammalian target of rapamycin (mTor) pathway, which likely causes the change of cell size in Traf3ip1 mutant cells (193).

5.3.11. Hippi (lft57)

lft57 has a coiled-coil domain at the C-terminus and interacts with lft20 in a yeast two-hybrid assay (181). The ortholog of *lft57* in mouse is *Hippi* (*huntingtin interacting protein-1 protein interactor*), which is involved in neuronal apoptotic cell death in Huntington disease (195). Houde *et al* generated Hippi-null mice (Hippi^{-/-}, C57BL/6J-129S1/SvImJ genetic background) by replacing the first 95 codons of the *Hippi* open reading frame with the neo gene or neo plus β -Gal (196). Hippi^{-/-} embryos predominantly die prior to E10.5 and have defects in nodal cilia, left-right patterning, and nervous system development. In addition, the expression levels of *Shh* and *ptch1* in the neural tube are down-regulated, suggesting that deletion of Hippi likely disrupts Hh signaling pathway (196).

5.3.12. lft70

lft70 is predicted to contain a number of TPRs (28). Mutations of the lft70 orthologs *dyf-1* in *C. elegans* or *fleer* in zebrafish led to defective cilia formation (197-199). The roles of lft70 in mouse and human disease have not been reported yet.

5.3.13. lft74/72 and lft81

lft74 and lft72 are derived from the same gene and often named as lft74/72 (88). Both lft74/72 and lft81 are core proteins in complex B. lft74/lft72 interact with the lft81. They all have coiled-coil regions in their structure (28). Very recently, Bhogaraju *et al* demonstrated that lft74 and lft81 form a tubulin-binding module, which is required for ciliogenesis in human RPE-1 cells (retinal pigment epithelium cells), suggesting lft74/lft81 play a role in the tubulin transportation within cilia (200). Further studies are needed to determine whether these lft proteins are involved in bone development.

5.4. Mouse models of lft motors

Anterograde and retrograde lft are operated by two types of MT-based motors, kinesin-2 (plus end-directed MT motor) and dynein-2 (minus end-directed MT motor). Although significant efforts have been contributed to elucidate the function of lft motors (See review (201)), their roles in ciliogenesis and

signaling transduction are still poorly characterized (Tables 4=5).

5.4.1. Dynein-2

The dynein motor protein family contains axonemal dynein, cytoplasmic dynein-1, and cytoplasmic dynein-2, which are involved in a wide range of functions in eukaryotic cells (See review (202)). Dynein-2 only exists in the organisms that have cilia or eukaryotic flagella and is believed to function primarily as the retrograde motor for lft (202). Dynein-2 contains a heavy chain (DYNC2H1), an intermediated chain (D2IC), a light intermediate chain (D2LIC, or LIC3), and a light chain (LC) (201, 203). The mouse homology of LC and D2IC has not been reported, and how the Dynein-2 motor interacts with lft protein complexes is largely unknown.

5.4.1.1. DYNC2H1

Dync2h1 encodes the heavy chain of the cytoplasmic dynein-2 motor (DYNC2H1, or DHC1b and DHC2) (35). The Dync2h1 mutation has been linked to the JATD and SRP (33, 35, 131). Chondrocytes from patients with Dync2h1 mutation had short cilia with bulbous distal ends (33).

Huangfu *et al* generated two mouse lines with Dync2h1 mutation named Dnchc2^{ln} and Dnchc2^{GT} (48). Dnchc2^{ln} is ENU induced mutation with a phenylalanine-to-serine (F3890S) mutation in a conserved residue of the Dnchc2 motor domain. Dnchc2^{GT} is generated by using the ESC gene trap insertion approach, which produces a truncated Dync2h1 (lacking the entire dynein motor domain) fused to β -geo (48). Homozygous Dnchc2^{ln}, Dnchc2^{GT}, and transheterozygous Dnchc2^{ln}/Dnchc2^{GT} mice all die at approximately E12.5 and have similar phenotypes, including abnormal brain morphology, randomization of heart-looping polarity and polydactylous. Cilia were present on the Dnchc2^{ln} embryos with altered morphology. These cilia are similar to those seen in the lft-A defect embryos, with reduced length and a bulge along their length (48, 204). Among Dync2h1^{ln}, Dync2h1^{tn} and Dync2h1^{Gt}, cilia in Dync2h1^{ln} are less affected (204). Hh signaling is disrupted, and the *Ptch1* expression is reduced in Dnchc2^{ln} neural tube. Besides, Hh pathway is fully activated in Ptch1 mutants. Dnchc2^{ln} Ptch1 double mutant mice has the similar phenotypes as Dnchc2^{ln} (48). Smo, Gli2, and Ptch1 accumulate along the Dnchc2^{ln} cilia with or without Shh stimulation (204). These results suggest that Dync2h1 acts downstream of Ptch1.

Table 4. Current mouse models to study the function of lft retrograde motors in bone development

Gene	Mouse models	Mutation	Cilia	Phenotype	Pathways	Ref
Dync2h1	Dnchc2 ^{lin}	ENU-mutagenesis, missense mutation	Abnormal cilia with reduced length and bulge along their length	Die at approximately E12.5, abnormal brain morphology, randomization of heart-looping polarity and polydactylous	Defects in Hh pathways with high level of Smo, Gli2 and Ptch1 accumulation along the cilia with or without Shh stimulation	(48, 204)
	Dnchc2 ^{GT}	Gene trap insertion, null allele	Not reported	Similar to Dnchc2 ^{lin}	Not reported	(48)
	Dync2h1 ^{tn}	ENU-mutagenesis, deletion in AAA domain 4	Abnormal cilia with reduced length and bulges along the axoneme	Not reported	Transportation of activated Gli2 out of cilia is blocked. Wnt signaling is normal in midgestation embryo or embryo-derived fibroblasts.	(67, 205)
	Dnchc2 ^{Q397Stop}	ENU-mutagenesis, null mutation	Shorter and wider cilia	Die at E12.0 or earlier with pericardial edema and heart failure. Severe defects in dorsoventral patterning of the forebrain and patterning of the spinal cord.	Both Gli activator and Gli repressor activity were significantly reduced. Disruption in Gli3 proteolytic processing and Smo localization to cilia.	(207)
	Dnchc2 ^{W2502R}	ENU-mutagenesis, null mutation	Shorter and wider cilia	Similar to Dnchc2 ^{Q397Stop}	Similar to Dnchc2 ^{Q397Stop}	(207)
	Dync2h1 ^{mmi}	ENU-mutagenesis, missense mutation	Swollen and filled with electron-dense particles and defects in axonemal	Defects in neural tube and motor neurons	Not reported	(161, 206)
	Dync2h1 ^{lin} lft172 ^{wim}	Double mutation	Absent of cilia	Similar to lft172 ^{wim}	Not reported	(204)
	Dync2h1 ^{lin} lft172 ^{avc1} and Dync2h1 ^{lin/lin} lft172 ^{avc1/+}	Double mutation	Normal morphology in Dync2h1 ^{lin/lin} lft172 ^{avc1/+}	Dync2h1 ^{lin} phenotype can be rescued by reducing lft172 expression, more dramatic rescue of the Dync2h1 phenotype in Dync2h1 ^{lin/lin} lft172 ^{avc1/+} (survived to at least E16.5 with nearly normal neural patterning in the caudal neural tube)	No lft or Hh pathway proteins accumulate in cilia, and Hh target gene expression appears to be normal	(204)
	lft122 ^{sopb} Dync2h1 ^{lin}	Double mutation	Similar to lft122 ^{sopb}	Similar to lft122 ^{sopb}	Ectopic activity of the Shh pathway in the neural tube. Gli2 limited to the tips of cilia and Smo was not in cilia without Shh stimulation	(204)
	Dync2h1 ^{lin/lin} lft122 ^{sopb/+}	Double mutation	Partially rescued the cilia morphology	Partially rescued the phenotype of Dync2h1 ^{lin/lin} including ventral neural development	Partially rescued Shh-dependent protein trafficking in cilia	(204)

(Contd..)

Table 4. Contd...

Gene	Mouse models	Mutation	Cilia	Phenotype	Pathways	Ref
mD2LIC	mD2LIC ^{-/-}	Gene targeting, null mutation	Cilia fail to form in the node	mD2LIC ^{-/-} died before E11.5 with defects in notochord and floorplate formation and a reduction in definitive endoderm	Reduction in Hh activity	(210)

Table 5. Current mouse models to study the function of lft anterograde motors in bone development

Gene	Mouse models	Mutation	Cilia	Phenotype	Pathways	Ref
Kif3a	Kif3a-Null	Removal exon 2 in ESCs	Cilia loss	Structural abnormalities, defects in left and right asymmetry and cilia loss in embryonic node	Not reported	(94)
	Kif3a ^{flox/flox} , Wnt1-Cre	Deletion in neural crest lineage cells	Not reported	Die shortly after birth due to respiratory failure, severe craniofacial defects with clefting and widened frontonasal prominence	Enhanced Hh signaling in facial mesenchyme Impaired Hh signaling in Kif3a-deficient neural crest-derived mesenchyme	(218, 219)
	Kif3a ^{flox/flox} , Dermo1-Cre	Deletion in trunk and limb skeletal progenitor cells	Not reported	Die at birth, split sternum, polydactyly and defects in ribs, knee joints and limbs	Not reported	(218)
	Kif3a ^{flox/flox} , Prx1-Cre	Deletion in the lateral plate mesoderm	Cilia loss	Forelimb polydactyly and profound shortening of limb bones	Disrupted Hh signaling in long bones	(174, 218)
	Kif3a ^{flox/flox} , Col2a1-Cre	Deletion in cartilage	Cilia disrupted (93, 220)	Postnatal dwarfism Abnormal cranial base development and altered synchondrosis growth plate organization and function at postnatal stages	Hh signaling significantly reduced within growth plates but enhanced and widespread all along perichondrial tissues Chondrocytes proliferation reduced but the hypertrophic differentiation is unregulated in growth plate, Hh signaling is unchanged	(93, 220)
	Kif3a ^{flox/flox} , Oc-Cre	Deletion in mature osteoblast	Reduced cilia number and length	Osteopenia by 6 weeks of age and partially recovered at 16 weeks	Impaired osteoblastic differentiation, impaired intracellular calcium response to fluid flow shear stress, and reduced Hh and Wnt response	(221)
	Kif3a; Col3a; I 2.3-Cre	Deletion in osteoblast progenitor	Not reported	No obvious skeletal defects. Reduced bone formation in response to mechanical ulnar loading	Not reported	(222)

(Contd...)

Table 5. Contd....

Gene	Mouse models	Mutation	Cilia	Phenotype	Pathways	Ref
Kif3b	Kif3b-Null	Null mutation with first exon replaced by neo cassette	Cilia loss	Die at mid-gestation and exhibited randomized left-right asymmetry, growth retardation, pericardial sac ballooning and neural tube disorganization	Not reported	(225)
Kif17	Kif17 ^{-/-} (231)	Frame shift with function loss	Not reported	No gross defects	Not reported	(231)
Kif7	Kif7 ^{flox/flox} ; NLS-Cre	Ubiquitous deletion with NLS-Cre	Not reported	Die at birth with severe malformations, including exencephaly, polydactyly and sternal defects	Up-regulated full-length Gli2 while down-regulated truncated 83-kD repressor Gli3 in total embryo lysates	(215)
	Kif7 ^{maki}	ENU- mutagenesis, Loss of function	Normal cilia	Die at the end of gestation with abnormal motor neurons and preaxial polydactyly	Reduced processed Gli3 proteins and the ratio of processed Gli3/full-length Gli3	(206)
	lft172 ^{wim} Kif7 ^{maki}	Double mutants (206)	Disrupt cilia	Similar to lft172 ^{wim}	Not reported	(206)
	Dync2h1 ^{mmi} Kif7 ^{maki}	Double mutants (206)	Not reported	Similar to Kif7 ^{maki} with greater loss of ventral neural cell types	A stronger loss of Shh signaling than Dync2h1 ^{mmi}	(206)
	Kif7-KO	Knockout allele	Normal cilia	Spinal cord defects, preaxial polydactyly, exencephaly, and microphthalmia, which mimic the Gli3 mutants phenotypes	Reduced proteolytic processing of Gli3. Reduce Gli2 and Gli3 accumulation at the tip of cilia in response to Hh	(216)

Dync2h1^{ttn} was generated by Ocbina *et al* also using ENU mutation method (205). Dync2h1^{ttn} carries an A-to-C substitution in the splice donor site upstream of exon 51, which deletes 38 amino acids from the ATPase-activator (AAA) domain 4 (205). Cilia morphology is altered on the Dync2h1^{ttn} MEFs with reduced length and bulges along the axoneme (67). Smo accumulates in the primary cilia of Dync2h1^{ttn} MEFs even in the absence of Shh. But the presence of Smo in cilia is not sufficient to activate the Gli-luciferase reporter, suggesting that retrograde lft blocks activated Gli2 to move out of the cilia (205). Wnt signaling is normal in mid-gestation Dync2h1^{ttn} embryo or embryo-derived fibroblasts (67).

Besides the above mentioned models, Liem *et al* generated another ENU-induced Dync2h1 mutation mouse line Dync2h1^{mmi} (206). Dync2h1^{mmi} has a missense mutation (I1789N) in the motor domain. Dync2h1^{mmi} has defects in neural

tube and motor neurons similar to other Dync2h1 mutants (206). Transmission electron microscopy (TEM) analysis revealed that cilia in Dync2h1^{mmi} are swollen and filled with electron-dense particles (161). Axonemal microtubules in Dync2h1^{mmi} cilia are defective having a reduced number of cilia and singlets instead of doublets (161).

May *et al* identified two independent ENU-induced null mutations, Dnchc2^{Q397Stop} and Dnchc2^{W2502R} (207). Dnchc2^{Q397Stop} carries an early nonsense mutation at amino acid position 397 (an early stop codon). Dnchc2^{W2502R} carries a missense mutation in a conserved AAA domain (position 2502) that is predicted to be required for motor activity. Both Dnchc2^{Q397Stop} and Dnchc2^{W2502R} show similar developmental defects with severe defects in dorsoventral patterning of the forebrain and patterning of the spinal cord (207). Most of the mutant embryos die at E12.0 or earlier with pericardial edema and heart failure. Cilia in mutant

embryos are shorter but wider than the normal cilia. Both Gli activator (*Gli1* expression) and Gli repressor activity (*Hoxd13* and *gremlin* expression) are significantly lower. In addition, mutant embryos have a disruption of the Gli3 proteolytic processing and Smo localized to the cilia (207).

Ocbina *et al* studied the relationships between *Dync2h1* and other genes required for ciliary trafficking. To investigate the relationship between anterograde ciliary trafficking and *Dync2h1*, they generated double mutant allele of *lft172*, an lft-B gene and *Dync2h1*. The phenotype of *Dync2h1^{lin}lft172^{wim}* double mutant is similar to that of *lft172^{wim}* (a null mutation allele), showing an absent of cilia and defects in all ventral neural cell types (204). Double mutation of *Dync2h1* and *lft172* using *lft172^{avc1}*, a partial loss-of-function allele of *lft172*, can partially rescue the *Dync2h1^{lin}* phenotype. Interestingly, a more dramatic rescue in Shh signaling and cilia morphology is found in the *Dync2h1^{lin}lft172^{avc1/+}* double mutants. *Dync2h1^{lin}lft172^{avc1/+}* embryos survive to at least E16.5 without having polydactyly. Thus, these results suggest that a modest reduction of lft-B expression and anterograde trafficking can rescue ciliogenesis and Shh signaling transduction in the absence of *Dync2h1* (204). To further elucidate the relationship between retrograde motors and lft-A proteins, *lft122* from lft-A was used to generate *lft122* and *Dync2h1* double mutant mice. Interestingly, mutants that are homozygous for both *Dync2h1^{lin}* and *lft122^{sopb}*, resemble the *lft122^{sopb}* phenotype with similar cilia morphology and ectopic activity of the Shh pathway in the neural tube (204). *lft88* is enriched at the cilia, and Gli2 is limited to the tips of cilia in *lft122^{sopb}* *Dync2h1^{lin}*. However, Smo is only in cilia with Shh stimulation in *lft122^{sopb}* *Dync2h1^{lin}*. Deletion of *lft122* relieves the accumulation of proteins within the cilia caused by loss of *Dync2h1*, suggesting *lft122* or lft-A complex plays a role in anterograde as well as retrograde trafficking (204). *lft122^{sopb}* leads to the lack of *Dync2h1* along the cilia, which may explain why the phenotypes in *lft122^{sopb}* *Dync2h1^{lin}* double mutants are similar to that in the *lft122^{sopb}* single mutants. *Dync2h1^{lin}lft122^{sopb/+}*, which has lowered expression of *lft122*, can partially rescue the phenotype of *Dync2h1^{lin}*, including the cilia morphology, ventral neural development and Shh-dependent protein trafficking in cilia (204), indicating that normal Shh-dependent protein trafficking in cilia can take place in the absence of *Dync2h1* if the *lft122* level is reduced.

5.4.1.2. mD2LIC

Mouse dynein-2 light intermediate chain (mD2LIC) was recently discovered. Both reciprocal immunoprecipitations and sedimentation assays showed that mD2LIC interacts specifically with DYNC2H1, suggesting mD2LIC is also involved in retrograde transport (208, 209).

Rana *et al* generated a null *mD2LIC* mutant mice (*mD2LIC^{-/-}*) by disrupting both the transcriptional and translational start sites in *mD2LIC* (210). *mD2LIC^{-/-}* mice die before E11.5 with defects in notochord and floorplate formation, a reduction in definitive endoderm and loss of cilia in the node of *mD2LIC^{-/-}* embryos. *mD2LIC^{-/-}* embryos displayed reduction in Hh signaling activity, which is similar to the phenotype observed in *Smo^{-/-}* and *Shh^{-/-}lhh^{-/-}* double mutants.

5.4.2. Kinesin-2

Kinesins are ATP-dependent microtubule-based molecular motors that carry cargo along microtubules in an anterograde direction (211, 212). There are more than 40 members in the kinesin superfamily that can be divided into 14 distinct families (kinesin-1 to -14) (213). Accumulating evidence shows heterotrimeric kinesin-2 motor is the "core" of IFT motor while other proteins including kinesin-2, -3 and -4 function as "accessory" motors for the transport of specific cargo dependent on cell types (211). In vertebrates, two distinct heterotrimeric kinesin-2 complexes, named kinesin-2 family protein A (Kif3a)/Kif3b/ kinesin-associated protein (KAP3) and Kif3a/Kif3C/ KAP3, which may play different roles in the ciliogenesis and function. In addition, heterotrimeric kinesin-2 and homodimeric Kif17 in the Kinesin-2 family also transport exogenously expressed cyclic nucleotide-gated channels into the cilium (214). Kif7 in the kinesin-4 family is localized to cilia base and the tip in mouse embryos and plays important roles in the ciliary trafficking and the response of Hh signaling (206, 215, 216).

5.4.2.1. Kif3a

Marszalek *et al* generated the Kif3a-Null mouse by removing exon 2 in ESCs, which completely disrupted Kif3a protein expression (94). The Kif3a-Null mouse usually dies at E10 with structural abnormalities, defects in left and right asymmetry, and cilia loss in embryonic node, suggesting that a kinesin-based transport is required for cilia formation (94).

Kif3a^{fllox/flox} was generated by inserting two LoxP sites, which flanks exon 2 of *Kif3a* in the intronic DNA sequences (94, 217). By crossing these mice with Wnt1-Cre (Kif3a^{fllox/flox}; Wnt1-Cre or Kif3a CKO), *Kif3a* is inactive in neural crest lineage cells. Kif3a^{fllox/flox}; Wnt1-Cre mice die shortly after birth due to respiratory failure and display severe craniofacial defects, including clefting of the secondary palate and widened frontonasal prominence (218, 219). Kolpakova-Hart *et al* reported the Kif3a^{fllox/flox}; Wnt1-Cre mice have the similar phenotype as Smo^{fllox/flox}; Wnt1-Cre mice (Shh signaling is disrupted in neural crest cells due to deletion of Smo). In addition, Gli1 is not detectable in the mesenchymal cells. Those data demonstrate that Hh signaling is impaired in Kif3a-deficient neural crest-derived mesenchyme (218). In contrast, Brugmann *et al* found that Kif3a-deficient neural crest-derived facial mesenchyme lacks cilia and has enhanced Hh signaling, including enhanced *Gli1*, *Ptch1* and *Shh*, but reduced *Gli3*, which accounts for the excessive proliferation in the facial prominences (219). The opposite Hh signaling response after deletion of Kif3a may result from inconsistent expression of Gli proteins in different tissues.

To study the role of Kif3a in axial and appendicular skeleton development, Dermo1-Cre (mesoderm-specific) was used to inactivate Kif3a in the trunk and limb skeletal progenitor cells (Kif3a^{fllox/flox}; Dermo1-Cre) (218). Kif3a^{fllox/flox}; Dermo1-Cre mice die immediately after birth, which likely result from the defects in ribcage. Besides, Kif3a^{fllox/flox}; Dermo1-Cre mice have a split sternum, polydactyly and defects in knee joints and limbs (218).

Restricted deletion of Kif3a in the lateral plate mesoderm with Prx1-Cre (Kif3a^{fllox/flox}; Prx1-Cre), which is limb/sternum-specific and does not affect the somitic mesoderm skeletal derivatives, such as ribs and vertebrae, leads to abnormal rib pairing, forelimb polydactyly and profound shortening of limb bones (174, 218). Ihh signaling is inactive (reduced *Gli1* expression and normal *Gli3* expression) in the long bone in Kif3a^{fllox/flox}; Prx1-Cre, which contributes to defects in chondrocyte differentiation and the loss of the bone collar adjacent to the metaphysis (174, 218).

Specific deletion of Kif3a in cartilage using Col2a1-Cre (Kif3a^{fllox/flox}; Col2a1-Cre) resulted in abnormal cranial base development and altered synchondrosis growth plate organization and function at the postnatal stage. Kif3a deficient synchondroses

display a delay in chondrocyte hypertrophy and enhanced intramembranous ossification. Hh signaling is significantly reduced within mutant growth plates but enhanced and widespread in all of the perichondrial tissues in Kif3a^{fllox/flox}; Col2a1-Cre mice as detected the expression of *Ptch1*, *Gli1*, and syndecan-3 (hedgehog-binding cell-surface component) (220). Song *et al* reported that Kif3a^{fllox/flox}; Col2a1-Cre mice have postnatal dwarfism. Chondrocytes proliferation is reduced, but the hypertrophic differentiation is up-regulated, which leads to premature loss of the growth plate in Kif3a^{fllox/flox}; Col2a1-Cre mice (93). Moreover, cilia, chondrocyte rotation, actin cytoskeleton and localization of activated FAK are all disrupted, suggesting that an important role of Kif3a or cilia in maintaining the columnar organization of the growth plate and chondrocyte differentiation. In addition, *Ptch1* expression of the Hh signaling pathway is not changed (93).

To analyze the function of Kif3a in osteoblast formation and function, Qiu *et al* used Oc-Cre (osteocalcin promoter, limited in mature osteoblast) to specifically delete Kif3a in the osteoblast lineage (Kif3a^{fllox/flox}; Oc-Cre). Kif3a^{fllox/flox}; Oc-Cre mice exhibit osteopenia at six weeks of age, which is partially recovered at 16 weeks. Kif3a deficiency leads to a reduction in the number of cilia and their length. Deletion of Kif3a increases cell proliferation but impairs osteoblastic differentiation. Qiu *et al* also found that deletion of Kif3a impairs intracellular calcium response to fluid flow shear stress and reduces Hh and Wnt responses, which may account for the defect of osteoblast-mediated bone formation in Kif3a^{fllox/flox}; Oc-Cre mice (221). Temiyasathit *et al* generated another osteoblast lineage mutant mouse line (Kif3a^{fllox/flox}; Kif3a; Col1(I)) using the Col1a 2.3-Cre (2.3kb fragment of the $\alpha 1(I)$ -collagen promoter, deletion in osteoblast progenitor cells) mice (222). Kif3a; Col1(I)-Cre mice have no obvious skeletal defects. However, Kif3a; Col1(I)-Cre mice exhibited reduced bone formation in response to mechanical ulnar loading, which linked the Kif3a/cilia to mechanosensor function in the bone (222).

Kinumatsu *et al* studied the role of Kif3a/cilia in temporomandibular joint (TMJ) development (223). Three-month-old Kif3a; Col2a1-cre mice had narrow and flat condyles with an irregular bony surface that was often fused with the articular disc. Alteration of Ihh signaling topography was associated with ectopic bone formation in condyles, implicating Kif3a/cilia are required for the coordination of Hh in chondrogenic condylar growth (223).

5.4.2.2. Kif3b

Kif3b is the other motor subunit in kinesin-2. A recent study suggested the Kif3b serves as a link between polycystin-2 (PC2, encoded by *PKD2*) and fibrocystin/polyductin (FPC, encoded by *PKHD1*, which is responsible for ARPKD), and mediates the regulation of PC2 channel function by FPC (224).

Nonaka *et al* generated the Kif3b-Null mutant mouse model by replacing most of the first exon (encoding the entire kinesin motor domain) with neo cassette (225). Kif3b-Null embryos die at mid-gestation and exhibit randomized left-right asymmetry, growth retardation, pericardial sac ballooning, and neural tube disorganization. Cilia is completely absent in the nodes of Kif3b-Null embryos.

5.4.2.3. Kif7

Kif7 is the mammalian homologue of *Drosophila* Costal2 (Cos2), which plays a central role in the Hh pathway in *Drosophila* (216, 226). Kif7 belongs to the kinesin-4 family and has an N-terminal motor domain (227). Knockdown of Kif7 in zebrafish resulted in ectopic Hh signaling, suggesting that Kif7 also acts in Hh signal in vertebrates (228).

Cheung *et al* generated the conditional allele of *Kif7* by flanking exons 2-4 with LoxP sites (*Kif7^{lox/lox}*) (215). By crossing *Kif7^{lox/lox}* with NLS-Cre transgenic mice (ubiquitous Cre expression), they created the null mutation of Kif7 (*Kif7^{lox/lox}; NLS-Cre*), and found that *Kif7^{lox/lox}; NLS-Cre* mice die at birth with severe malformations, including exencephaly, polydactyly, and sternal defects, a phenotype similar to the phenotype of Gli3 mutant mice. They further found that Kif7 physically interacts with Gli transcription factors and controls their proteolysis and stability (215). The deletion of Kif7 up-regulates full-length Gli2 while it down-regulates truncated 83-kD repressor Gli3 in total embryo lysates. To fully understand the role of Kif7 in Hh signaling pathway, they further generated double mutants by using *Smo* mutant, *Gli2* mutant, or *Gli3* mutant mice to cross with *Kif7^{lox/lox}; NLS-Cre*, and compared these mutants with each single mutant. Their results suggested that Kif7 plays a critical role in the balance of Gli activators and repressors downstream of Smo (215).

Liem *et al* identified an ENU-induced *matariki* (*maki*) mutation of *Kif7* (*Kif7^{maki}*) (206). *Kif7^{maki}* is a missense mutation (L130P) in a conserved region of the motor domain of *Kif7*, which

led to the loss of Kif7 function. *Kif7^{maki}* mice die at the end of gestation with abnormal motor neurons, preaxial polydactyly and elevated Shh signaling. The results from *Smo* and *Kif7* double mutation result in Shh-dependent ventral neural cell types at E10.5, suggesting that Kif7 functions downstream of Smo and that loss of Kif7 causes ligand-independent activation of Shh pathway. Loss of Kif7 leads to a reduction in processed Gli3 proteins and in the ratio of processed Gli3/full-length Gli3, confirming the role of Kif7 in Gli3 proteolysis or stability. *Kif7^{maki}* mice have normal cilia. To study the relationship between Kif7 and cilia, *lft172^{wim}Kif7^{maki}* double mutants were generated by Liem *et al* (206). They found that *lft172^{wim}Kif7^{maki}* is similar to *lft172^{wim}* mutants, suggesting that the activity of Kif7 depends on the presence of cilia. To analyze the role of Kif7 in ciliary trafficking (206), Liem *et al* also generated *Dync2h1^{mmi}Kif7^{maki}* double mutants and found that *Dync2h1^{mmi}Kif7^{maki}* phenocopies *Kif7^{maki}* mutants with greater loss of ventral neural cell types, suggesting that Kif7 functions before retrograde lft, and that DYNC2H1 has additive and positive roles in Shh signaling (206).

Endoh-Yamagami *et al* generated *Kif7* knockout mice (*Kif7-KO*) by replacing the initiation codon and following N-terminus motor domain with a neo cassette (216). *Kif7-KO* embryos display spinal cord defects, preaxial polydactyly, exencephaly, and microphthalmia, which mimic the phenotypes of Gli3 mutants. The Gli3 repression targets *Gremlin* and *HoxD13* are elevated in *Kif7-KO*. Meanwhile, *Gli1* and *Ptch1* expression are normal in both *Kif7-KO* embryos and MEFs. Those data suggest that Kif7 regulates Hh signaling mainly through controlling the formation or function of the Gli3 repressor. Cilia morphology is normal in MEFs from *Kif7-KO* embryos. Kif7 accumulates at the tip of the cilia in an Hh-dependent manner. Kif7 physically interacts with Gli3 and is required for efficient Gli3 processing to its repressor form, indicating that Kif7 functions principally as a negative regulator of the Hh pathway in mice.

5.4.2.4. Kif17

Kif17 belongs to the kinesin-2 family, and a pair of Kif17 forms a homodimer with their head domain (229). Kif17 has diverse functions in different species and different kind of cells. Dysfunctions of Kif17 are linked to a number of pathologies (See review (229)). Kif17 or its homolog mediates lft cargos to the ciliary tip, therefore, it plays a role in ciliogenesis and function (229). Recently, a new

peptide named TTC308 has been identified in lft complex B in mouse pituitary cells that interacted with mouse Kif17 in co-immunoprecipitation studies (230). This finding confirmed the possible functional interaction between Kif17 and cargo transport. Whether Kif7 takes part in the ciliogenesis in mice is currently unknown.

Yin *et al* generated Kif7 mutant mice (Kif17^{-/-}) by replacing exon 4 of Kif17 with a poly (A)-less PGK-neo cassette (231) resulting in a frame shift, and disruption of Kif17 expression. Kif17^{-/-} mice grow normally without any gross defects. They found that Kif7 is important for maintenance of N-methyl-D-aspartate (NMDA) receptor subunit 2B (NR2B) and NR2A level in hippocampal neurons, which is involved in memory and the learning process. However, whether Kif17^{-/-} mice have the bone related phenotype has not been evaluated yet (231).

6. SUMMARY AND PERSPECTIVES

The development of transgenic and mutagenesis technology provides powerful tools to study the function of cilia, lft proteins, and motors. Some of the lft proteins and motors already have the mouse models established. A few of them have more than one model with different degrees of loss of function, allowing the study of dosage dependent effects of these genes. The loss of lft complex A protein usually leads to the “swollen” cilia, which are filled with complex B proteins, suggesting that the complex B, which is in charge of the anterograde transport, can function without the complex A. Mutations of most lft complex B proteins lead to cilia disruption due to the totally blocked anterograde transport. This raises the problems to distinguish the function of individual lft proteins and to identify whether the phenotype results from a deficient lft protein alone or is secondary to cilia loss. It is also possible that the severe consequence of loss of cilia masks the other function of lft proteins independent of cilia.

Over the past few years, much attention has been paid to elucidating the gene functions and mimicking the human diseases using mouse models. However, the treatment of lft- and cilia-related diseases in humans is still at the initial developmental stages. For the patients with the lft protein or lft motor mutation, exogenous introduction of the lft or lft-related proteins or gene to treat these diseases are worth trying. However, before doing so, it is necessary to figure out which genes are defective in each individual patient and how these genes interact

with other genes in the cells. Currently, several candidate modulators for ciliogenesis and cilium length have been found from one high-throughput functional screening (232). Further studies should elucidate whether these modulators can rescue ciliogenesis and cilia function in lft proteins/motors defective or mutant cells. Proper dosage and delivery methods could be tested with the described mouse models. We anticipate that those small molecules will provide us novel strategies to treat ciliopathies. More work is needed to achieve the comprehensive understanding of the downstream pathways of cilia. Much attention has been paid to the Hh signaling pathways, but the interactions of Hh signaling and cilia/lft, and the relationship of cilia/lft with Wnt, Notch, PDGF and many other pathways are still ambiguous or largely unknown. With better knowledge of these pathways and how they interact with cilia/lft, proper agonists or antagonists can be selected or developed to regulate the downstream signaling. These mouse models provide the needed tools to test the target candidates for the treatment of the ciliopathies patients.

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- Abbreviations:** MTs – microtubules; Ift – intraflagellar transport; JATD - Jeune asphyxiating thoracic dystrophy; SRP - short rib polydactyly syndromes; Hh – hedgehog; Ihh - Indian hedgehog; Shh - Sonic hedgehog; Ptch1

- Patched 1; Smo – Smoothed; PCP - planar cell polarity; BBS - Bardet–Biedl syndrome; Evc - Ellis-van Creveld syndrome protein homolog; OFD1 - oral-facial-digital syndrome 1; MKS - Meckel-Gruber syndrome; JBTS - Joubert syndrome; JSRD - JBTS-related disorder; LCA - Leber congenital amaurosis; PCD - primary ciliary dyskinesia; KS - Kartagener syndrome; NPHP – nephronophthisis; SLS - Senior-Loken syndrome; ,ADPKD - Autosomal dominant polycystic kidney disease; ARPKD - autosomal recessive polycystic kidney disease; US - Usher syndrome; ALMS - Alström syndrome; MSS - Mainzer-Saldino-Syndrome; ATD - asphyxiating thoracic dystrophy; WAD - Weyers acrofacial dysostosis; NEK1 - never-in-mitosis Kinase 1; CED - Cranioectodermal dysplasia; MKS - Meckel syndrome; ENU - N-ethyl-N-nitrosourea; PDGF-A - platelet-derived growth factor A; ESCs -embryonic stem cells; THM1 - Tetratricopeptide Repeat Containing Hedgehog Modulator 1; MEFs - mouse embryo fibroblasts; PD - proximal-distal; AP - anterior-posterior; ACIII - adenylyl cyclase III; SUFU - Suppressor of Fused; BMSCs - bone marrow derived stromal cells; Sfrp5 - secreted frizzled-related sequence protein 5; OA - osteoarthritis; neo - neomycin; GMAP210 - Golgi Microtubule Associated Protein 210; TRIP11 - thyroid hormone receptor interacting protein 11; HRGP - human red/green pigment gene promoter; GTPase - guanosine triphosphate hydrolases; Fgfr1-fibroblast growth factor receptor-1; Col12 α 1 - type XII collagen; Bgn – biglycan; Mmp10 - matrix metalloproteinase-10; Hsp47 - heat shock protein 47 kDa; Traf3 - TNFR-associated factor-3, IL-13R1 - interleukin-13 receptor α 1, DISC1 - Disrupted-In-Schizophrenia 1; STAT6 - signal transducer and activator of transcription-6; AAA - ATPase-activator; PC2 - polycystin-2; TMJ - temporomandibular joint; FPC - fibrocystin/polyductin; OSM-3 - osmotic avoidance abnormal protein 3; NMDA - N-methyl-D-aspartate ; NR2B - NMDA receptor subunit 2B

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