Genetic interaction between central pair apparatus genes CFAP221, CFAP54, and SPEF2 in mouse models of primary ciliary dyskinesia

Casey W. McKenzie & Lance Lee

Primary ciliary dyskinesia (PCD) is a genetically heterogeneous syndrome that results from defects in motile cilia. The ciliary axoneme has a 9 + 2 microtubule structure consisting of nine peripheral doublets surrounding a central pair apparatus (CPA), which plays a critical role in regulating proper ciliary function. We have previously shown that mouse models with mutations in CPA genes CFAP221, CFAP54, and SPEF2 have a PCD phenotype with defects in ciliary motility. In this study, we investigated potential genetic interaction between these CPA genes by generating each combination of double heterozygous and double homozygous mutants. No detectable cilia-related phenotypes were observed in double heterozygotes, but all three double homozygous mutant lines exhibit early mortality and typically develop severe PCD-associated phenotypes of hydrocephalus, mucociliary clearance defects in the upper airway, and abnormal spermatogenesis. Double homozygous cilia are generally intact and display a normal morphology and distribution. Spermiogenesis is aborted in double homozygotes, with an absence of mature flagella on elongating spermatids and epididymal sperm. These findings identify genetic interactions between CPA genes and genetic mechanisms regulating the CPA and motile cilia function.
Figure 1. Early mortality of mice lacking two CPA genes. (a) Schematic diagram of the CPA showing the location of CFAP221, CFAP54, and SPEF2, which are absent in the nm1054, Cfap54<sup>4018</sup>, and bgh mouse lines, respectively. Modified with permission from McKenzie et al.<sup>29</sup>. (b) Survival curve showing mortality of nm1054/ nm1054; bgh/bgh, nm1054/nm1054;Cfap54<sup>4018</sup>/Cfap54<sup>4018</sup>, and bgh/bgh;Cfap54<sup>4018</sup>/Cfap54<sup>4018</sup> double homozygotes prior to adulthood.

The gene encoding centrosomal protein 290 (CEP290), resulting in double homozygotes with a more severe and rapid progression of retinal degeneration, indicating a genetic interaction between RPGR and CEP290<sup>26</sup>. Finally, homozygous loss of CPA genes sperm associated antigen 6 (SPAG6) and sperm associated antigen 16L (SPAG16L) produce very different phenotypes. SPAG6 mutants exhibit the full spectrum of PCD phenotypes, CPA structural defects, and reduced ciliary beat frequency<sup>17–19</sup>, while mice lacking SPAG16L have only spermatic defects and male infertility but no other cilia-associated phenotypes<sup>20</sup>. However, a genetic interaction was identified in double homozygous mutants, which have hydrocephalus and abnormal airway pathology that are more severe than the SPAG6 homozygotes alone<sup>31</sup>. Despite the paucity of studies investigating genetic interactions, the phenotypes of double mutant mice uncover new information about the roles of these genes in motile cilia function.

The CPA is a complex structure that regulates ciliary motility and waveform<sup>19,23</sup>. It is comprised of protein complexes, or projections, that associate with the core microtubules. We have previously shown that mouse models with mutations in CPA genes ciliary and flagellar associated protein 221 (CFAP221), ciliary and flagellar associated protein 54 (CFAP54), and sperm flagellar protein 2 (SPEF2) each have a PCD phenotype that includes hydrocephalus, male infertility, and airway abnormalities<sup>26–30</sup>. The PCD in mice homozygous for the nm1054 mutation, a large deletion that removes CFAP221, also known as primary ciliary dyskinesia protein 1 (PCDPI), results from cilia with reduced ciliary beat frequency (CBF) but no detectable ultrastructural abnormalities<sup>26</sup>. The *Chlamydomonas reinhardtii* homolog of CFAP221 associates with the C1d projection, a calcium-dependent CPA complex required for proper flagellar motility<sup>31–33</sup>. A gene trapped allele of CFAP54, which also encodes a member of the C1d projection complex in *C. reinhardtii*, results in reduced CBF perturbed cilia-driven fluid flow, and loss of the C1d projection<sup>34</sup>. The ultrastructural defect observed in cilia from mice lacking CFAP54 indicates that CFAP54 and CFAP221 are functionally distinct despite both proteins associating with the same complex in *C. reinhardtii*. Finally, the spontaneous big giant head (bgh) mouse line has a nonsense mutation in SPEF2 that, like mice lacking CFAP221, does not result in axonemal ultrastructural defects but does reduce the CBF<sup>27</sup>. Unlike CFAP221 and CFAP54, however, SPEF2 localizes to the C1b projection in *C. reinhardtii* rather than the C1d projection<sup>34</sup>. Figure 1a shows a schematic diagram of the CPA and indicates the location of the CFAP221, CFAP54, and SPEF2 proteins. Since nodal cilia lack a CPA, none of these models develop the lateral-flow, and loss of the C1d projection<sup>29</sup>. The ultrastructural defect observed in cilia from mice lacking CFAP54 indicates that CFAP54 and CFAP221 are functionally distinct despite both proteins associating with the same complex in *C. reinhardtii*. Finally, the spontaneous big giant head (bgh) mouse line has a nonsense mutation in SPEF2 that, like mice lacking CFAP221, does not result in axonemal ultrastructural defects but does reduce the CBF<sup>27</sup>. Unlike CFAP221 and CFAP54, however, SPEF2 localizes to the C1b projection in *C. reinhardtii* rather than the C1d projection<sup>34</sup>. Figure 1a shows a schematic diagram of the CPA and indicates the location of the CFAP221, CFAP54, and SPEF2 proteins. Since nodal cilia lack a CPA, none of these models develop the laterality defects often associated with PCD. Recently, CFAP221 and SPEF2 mutations were identified in human PCD patients<sup>35–37</sup>, highlighting the importance of these CPA genes.

Here, we investigate potential genetic interaction between CFAP221, SPEF2, and CFAP54 by crossing the nm1054, bgh, and Cfap54<sup>4018</sup> mouse lines. Double heterozygotes show no detectable PCD phenotypes, but the nm1054/nm1054; bgh/bgh, nm1054/nm1054;Cfap54<sup>4018</sup>/Cfap54<sup>4018</sup>, and bgh/bgh;Cfap54<sup>4018</sup>/Cfap54<sup>4018</sup> double homozygotes all exhibit early mortality and often severe hydrocephalus and airway phenotypes. Double homozygous cilia are generally intact with a normal morphology and distribution. Severe defects in spermatogenesis were also observed, although marker expression analysis suggests that some ciliary components are still being assembled. These data underscore the critical role of CPA proteins in regulating proper motile cilia function and unveil genetic interactions between CPA genes.

Results

Double homozygosity results in early lethality. Mice homozygous for the nm1054, Cfap54<sup>4018</sup>, and bgh mutants lack CPA projection proteins CFAP221, CFAP54, and SPEF2, respectively (Fig. 1a). Each single mutant was shown to exhibit early mortality on the C57BL/6J (B6) background due to severe hydrocephalus but typically live a normal life span on the 129S6/SvEvTac (129) background or a mixed background without gross hydrocephalus<sup>26,27,29</sup>. In contrast, early mortality was common for double homozygotes on a mixed background (Fig. 1b). The most severe mortality phenotype was observed in nm1054/nm1054; bgh/bgh double mutants, with all mice dying or requiring euthanasia due to severe hydrocephalus by weaning age (3 weeks). The bgh/bgh;Cfap54<sup>4018</sup>/Cfap54<sup>4018</sup> double homozygotes also demonstrated substantial mortality, with less than half surviving to adulthood (8 weeks). Survival was best for nm1054/nm1054;Cfap54<sup>4018</sup>/Cfap54<sup>4018</sup> double mutants, with most mice surviving to adulthood. Gross hydrocephalus was common in mice that died prior to 8 weeks, regardless of genotype. All double heterozygotes (nm1054/+; bgh/+ , nm1054/+;Cfap54<sup>4018</sup>/+, and bgh/+;Cfap54<sup>4018</sup>/)
survived to adulthood and were analyzed at 8 weeks or older, along with wild type (WT) controls (Supplementary Table S1).

**Double homozygosity results in severe hydrocephalus.** To assess the extent of damage in the brains of double mutant mice on a mixed background, we performed histological analyses on coronal sections through the lateral ventricles, through which cerebrospinal fluid (CSF) flows and accumulates under hydrocephalic conditions. The lateral ventricle is appropriately narrow in WT and double heterozygous mice with no evidence of tissue damage (Fig. 2a–d). However, a severe hydrocephalic phenotype was observed in all three double homozygotes (Supplementary Fig. S1). The nm1054/nm1054;bgh/bgh brain shows enlarged ventricles but not extensive tissue damage, and ependymal cilia lining the ventricular wall are intact (Fig. 2e,h). It is important to note, however, that only one mouse survived long enough for analysis (Fig. 1b), so the phenotype may have been more severe in those animals that died earlier. The nm1054/nm1054;Cfap54gt/gt;Cfap54gt/gt double homozygotes typically showed mild ventricular dilatation without extensive tissue damage, and ependymal cilia remained intact (Fig. 2f,i). The most severe phenotype was observed in bgh/bgh;Cfap54gt/gt;Cfap54gt/gt brains, which commonly exhibited severe ventricular dilatation, as well as extensive damage to the white matter and cerebral cortex that was often accompanied by intraventricular hemorrhaging with a dramatic influx of red blood cells (Fig. 2g,j). While nm1054, bgh, and Cfap54gt/gt single mutants were each shown to exhibit severe hydrocephalus, this phenotype was only observed on the more susceptible B6 background and not on a mixed background.

**Double homozygosity results in severe airway abnormalities.** Since motile ciliary dysfunction typically results in airway abnormalities, we assessed the pathology of the maxillary sinus cavity. WT and double heterozygous mice have a clear sinus cavity with little to no evidence of mucus (Fig. 3a–d). In contrast, all three double homozygotes exhibit a more severe defect in mucociliary clearance and airway pathology (Supplementary Fig. S1). The one surviving nm1054/nm1054;bgh/bgh mouse shows severe accumulation of mucus and extensive infiltration of neutrophils, which is common in sinusitis pathology and is indicative of the tissue repair process (Fig. 3e,h). Despite the mucus and white blood cell accumulation, some motile cilia remain intact on the epithelial surface (Fig. 3h). The nm1054/nm1054;Cfap54gt/gt;Cfap54gt/gt double homozygotes typically showed some areas of mild mucus accumulation but no evidence of extensive neutrophil infiltration, and epithelial cilia remain intact (Fig. 3f). Like the nm1054/nm1054;bgh/bgh mouse, the bgh/bgh;Cfap54gt/gt;Cfap54gt/gt sinus cavity typically shows severe accumulation of mucus and areas with mild infiltration of neutrophils, sometimes accompanied by red blood cells (Fig. 3g,j). Motile cilia remain largely intact on bgh/bgh;Cfap54gt/gt;Cfap54gt/gt sinus epithelia (Fig. 3i). The sinus phenotype in double homozygous mice is consistent with mucus accumulation in the single nm1054, bgh, and Cfap54gt/gt mutants on a mixed background, as well as neutrophil infiltration in bgh mice, but the presence of red blood cells was never observed in single mutants.

The morphology and distribution of motile cilia lining the sinus cavity was further assessed by immunohistochemistry (IHC). Ciliary marker acetylated tubulin is expressed normally in WT cilia (Fig. 4a,a′), while basal body marker γ-tubulin localizes to the ciliary base (Fig. 4f,f′). Staining of acetylated tubulin confirms that the intact cilia on all three double homozygotes have a generally typical morphology and distribution consistent with WT mice, and there is no statistically significant difference in acetylated tubulin levels (Fig. 4b–e,b′–d′). In addition, γ-tubulin is normally expressed and properly localized to the ciliary base in all three double homozygotes, suggesting that cilogenesis is not perturbed (Fig. 4g–j,g′–j′). Known CPA protein SPAG6 is expressed throughout the axoneme of WT cilia (Fig. 4k,k′), and relatively weak SPAG6 staining in cilia from double homozygous sinus epithelia suggests that defects in proper CPA assembly are possible, although there are no statistically significant differences (Fig. 4l–n,l′–n′). Analysis of known radial spoke marker RSPH4A shows little difference in axonemal expression or protein level between WT and double homozygous mice (Fig. 4p–t,p′–t′), further confirming a largely normal ciliary assembly. Consistent with the normal pathology of the sinus cavity in double heterozygous mice, the morphology and distribution of motile cilia appears normal in all three double heterozygotes, also with no statistically significant difference in acetylated tubulin levels (Supplementary Fig. S2). In addition, tracheal CBF was measured for double heterozygotes using a high speed video microscopy approach, and levels were not statistically different from WT for any of the double heterozygous mice, confirming that the motile cilia are functionally normal (Supplementary Fig. S3). CBF analysis could not be performed efficiently or accurately on double homozygotes due to the low number of mice surviving to adulthood and technical challenges associated with measurement using the small neonatal or young postnatal trachea.

**Double homozygosity impairs spermatogenesis.** Although male mice lacking CFAP221, SPEF2, or CFAP54 are fertile, fertility tests were not performed for double homozygotes due to the low number of mice surviving to sexual maturity. Double heterozygotes are fertile and were used routinely for breeding to obtain double homozygotes. Effect of double homozygosity on spermatogenesis was investigated using a histopathological approach. As elongating spermatids develop into mature spermatozoa during spermiogenesis, the final step of spermatogenesis, WT flagella extend into the lumen of the seminiferous tubule of the testis (Fig. 5a). In contrast, nm1054/nm1054;Cfap54gt/gt;Cfap54gt/gt and bgh/bgh;Cfap54gt/gt;Cfap54gt/gt double homozygotes both show an absence of flagella in the lumen of the seminiferous tubule (Fig. 5b,c), indicating that flagellar formation is perturbed. While this is different from the presence of motile cilia on the sinus epithelia from these mice, it is not dissimilar to the spermatogenesis phenotype observed in single nm1054, bgh, or Cfap54gt/gt mutants. No testis data was obtained for nm1054/nm1054;bgh/bgh double homozygotes, as none survived to sexual maturity, and the only one that survived long enough to collect tissues was a female. Normal spermiogenesis was observed in the testes from nm1054/+;bgh/+; nm1054/+;Cfap54gt/gt/+; and bgh/+;Cfap54gt/gt/+ double heterozygous mice (Fig. 5d–f), which is consistent with their fertility.
To further investigate the ability of double homozygous testes to assemble flagella, we performed IHC analysis. An antibody to ciliary and flagellar marker acetylated tubulin effectively detects the elongating flagella in the lumen of the WT seminiferous tubule (Fig. 6a,a'). While no flagella are present in the lumen of nm1054/nm1054;Cfap54gt/gt or bgh/bgh;Cfap54gt/gt double homozygous testes, there are defined structures detected by the acetylated tubulin antibody in the developing spermatids from both mutants (Fig. 6b,c,b',c'), suggesting that some rudimentary axonemal structures possessing acetylated tubulin may be assembled in those cells. The antibody to basal body marker γ-tubulin primarily stains immature spermatogonia and developing spermatids in the WT seminiferous tubule (Fig. 6d).
spermatocytes within the WT seminiferous tubule and both double homozygotes (Fig. 6d–f,d′–f′), suggesting that early stages of spermatogenesis are unaffected. However, while CPA protein SPAG6 is detected throughout the flagellar axoneme in WT seminiferous tubules (Fig. 6g,g′), low SPAG6 levels without a clear staining pattern in double homozygotes suggests that CPA assembly may be impaired (Fig. 6h,h′,i,i′). The defined axonemal structures detected by the acetylated tubulin antibody in nm1054/nm1054;Cfap54gt/gt and bgh/bgh;Cfap54gt/gt spermatids are not detected by the SPAG6 antibody, with only occasional sperm heads observed. An antibody to known dynein marker DNAI1 also stains the length of the flagellar axoneme in WT testis (Fig. 6j,j′). In contrast to SPAG6, DNAI1 shows strong expression in the developing spermatids surrounding the lumen of the nm1054/nm1054;Cfap54gt/gt and bgh/bgh;Cfap54gt/gt seminiferous tubules (Fig. 6k,k′,l,l′), indicating that dynein pre-assembly may occur in the spermatid cytoplasm even in the absence of mature flagella. Acetylated tubulin staining in nm1054/+;bgh/+, nm1054/+;Cfap54gt/gt/+ and bgh/+;Cfap54gt/gt/+ double heterozygotes highlights their morphologically normal elongating spermatid flagella (Supplementary Fig. S4).

In addition to testis pathology, the morphology of mature sperm from the epididymis was also assessed using light microscopy. WT sperm have a hook-shaped head and a long flagellum (Fig. 7a). In contrast, sperm were rarely present in the nm1054/nm1054;Cfap54gt/gt or bgh/bgh;Cfap54gt/gt double homozygous epididymides, and those that were present have normal heads but severely shortened flagellar stubs that would likely impair proper sperm motility (Fig. 7b,c). As described above, sperm data could not be obtained for nm1054/nm1054;bgh/bgh double homozygotes, as only one female mutant survived long enough to collect
tissues. Consistent with the testis data and normal fertility, all double heterozygous mice have epididymal sperm that are indistinguishable from WT sperm (Fig. 7d–f).

Discussion

In this study, we investigated potential genetic interaction between CPA genes CFAP221, SPEF2, and CFAP54 by crossing the nm1054, bg/h, and Cfap54<sup>486<sup>gt</sup>/Cfap54<sup>486<sup>gt</sup></sup> mouse models with mutations in those genes. We identified an absence of ciliary phenotypes in double heterozygotes, but nm1054/nm1054:bg/h, nm1054/nm1054:Cfap54<sup>486<sup>gt</sup>/>Cfap54<sup>486<sup>gt</sup></sup>, and bg/h:bg/h:Cfap54<sup>486<sup>gt</sup>/>Cfap54<sup>486<sup>gt</sup></sup> double homozygotes all exhibit PCD phenotypes of hydrocephalus,
mucociliary clearance defects, and spermatogenesis abnormalities. Double mutants also exhibit severe early mortality on a mixed background that is not observed in single mutants. Double homozygous cilia appear intact and normally distributed, with the exception of the ependymal cilia, where the severe ventricular dilatation observed in \( bg/h;Cfap54^{gt/gt}/Cfap54^{gt/gt} \) double mutants can cause extensive tissue damage. In addition, spermiogenesis is aborted in double homozygous males, and there is an absence of mature flagella on elongating spermatids and epididymal sperm.

The absence of histopathological or ciliary phenotypes in any of the double heterozygous mice (\( nmt1054/++;bg/h++;nmt1054/++;Cfap54^{gt/gt}/++;, bg/h++;Cfap54^{gt/gt}/++) indicates that haploinsufficiency for two different CPA genes does not result in a ciliary phenotype. While this is expected given the autosomal recessive inheritance associated with these mutations, it demonstrates that a dose-dependent reduction in two different CPA proteins does not put a significant enough strain on the CPA to noticeably affect its function or the function of the motile cilium. The PCD phenotypes associated with the double homozygous mutants, however, are striking and generally more severe than single mutants. Most notable is the early mortality associated with double homozygotes on a mixed genetic background (Fig. 1). Homozygotes with either the \( nmt1054, bg/h, \) or \( Cfap54^{gt/gt} \) mutation exhibit early mortality on the B6 background due to severe hydrocephalus, presumably due to genetic modifiers segregating in that strain that influence susceptibility to severe hydrocephalus. On the 129 background or a mixed background, single homozygotes do not develop severe hydrocephalus and live a normal life span, despite exhibiting airway ciliary phenotypes and male infertility. All three double homozygotes (\( nmt1054/nmt1054;bg/h;bg/h, nmt1054/nmt1054;Cfap54^{gt/gt}/Cfap54^{gt/gt}, \) and \( bg/h;bg/h;Cfap54^{gt/gt}/Cfap54^{gt/gt} \)) develop hydrocephalus on a mixed background (Fig. 2), suggesting that the defect in CSF flow in double homozygotes may be primarily due to severe ciliary dysfunction and highlighting genetic interactions. Given that several double homozygotes had to be euthanized in the early post-natal period due to severe hydrocephalus, the hydrocephalus is the most likely cause of the early mortality.

Despite the presence of motile cilia in the brain and airway of double homozygous mutants, there is an absence of mature flagella on elongating spermatids and epididymal spermatozoa from \( nmt1054/nmt1054;Cfap54^{gt/gt}/Cfap54^{gt/gt} \) and \( bg/h;bg/h;Cfap54^{gt/gt}/Cfap54^{gt/gt} \) double homozygous mutants (Figs. 5, 7). Reproductive data was not obtained for \( nmt1054/nmt1054;bg/h;bg/h \) double homozygotes due to mortality prior to sexual maturity. The difference between motile cilia and sperm flagella is consistent with single \( nmt1054, bg/h, \) and \( Cfap54^{gt/gt} \) homozygous mutants, as well as mutations in other axonemal genes that prevent proper flagellar formation without affecting ciliogenesis. Presence of acetylated tubulin staining in developing spermatids

**Figure 5.** Histological analysis of double mutant testes. Sections of WT (a), double homozygous (b,c), and double heterozygous (d–f) testes showing an absence of elongating spermatid flagella in double homozygotes. Sections are stained with H&E. Arrowheads indicate the lumen of the seminiferous tubule, where elongating spermatid flagella are generated during spermiogenesis.
Figure 6. Immunohistochemical analysis of ciliary markers in double homozygous testes. Sections of WT, nm1054/nm1054;Cfap54gt/gt, and bgh/bgh;Cfap54gt/gt testis stained with antibodies to flagellar marker acetylated tubulin (a–c, a′–c′), basal body marker γ-tubulin (d–f, d′–f′), CPA protein SPAG6 (g–i, g′–i′), and dynein protein DNAI1 (j–l, j′–l′). The arrowheads indicate rudimentary axonemal structures (b, b′) and developing sperm heads (h′, i′), and the arrows indicate the spermatogonium cell type (e, e′).
from nm1054/nm1054;Cfap54gt/gt and bgh/bgh;Cfap54gt/gt double homozygotes suggests that the spermatids start to make axonemes, but the process aborts early in spermiogenesis (Fig. 6). In addition, DNAI1 staining in spermatids also indicates that cytoplasmic dynein pre-assembly is occurring in the double homozygotes. However, low staining without a clear staining pattern for CPA marker SPAG6, along with absence of detectable axonemal structures, suggest that CPA assembly may be perturbed. These data underscore fundamental differences in the mechanisms that drive and regulate motile ciliogenesis and sperm flagellar formation.

Interestingly, the nm1054/nm1054;Cfap54gt/gt double homozygous phenotype is noticeably less severe than the phenotypes of nm1054/nm1054;bgh/bgh and bgh/bgh;Cfap54gt/gt double homozygotes. The nm1054/nm1054;Cfap54gt/gt double mutants exhibit a much higher survival rate (Fig. 1), show mild ventricular enlargement in the brain (Fig. 2, Supplementary Fig. S1), and display only modest amounts of mucus accumulation in the sinus cavity without extensive white blood cell infiltration (Fig. 3, Supplementary Fig. S1). This mild phenotype may be due to CFAP221 and CFAP54 both associating with the C1d projection of the CPA (Fig. 1). If disrupting both proteins still largely only perturbs the single complex, the phenotype may not be substantially more severe than single nm1054 or Cfap54gt/gt mutants. SPEF2, however, localizes to the C1b projection, so loss of SPEF2 and either CFAP221 or CFAP54 would likely disrupt two different protein complexes, put additional strain on the CPA, and have a more substantial effect on ciliary function. As a result, nm1054/nm1054;bgh/bgh and bgh/bgh;Cfap54gt/gt double homozygotes have more severe brain and airway phenotypes, further highlighting genetic interactions between SPEF2 and CFAP221 or CFAP54. These findings are consistent with previous studies identifying genetic interactions between RPGR and CEP29016 and between SPAG6 and SPAG16L21, both due to particularly severe ciliary phenotypes.

Because of the early mortality associated with double homozygosity, combined with the low rate of double homozygote birth (1/16 of offspring from two double heterozygotes), mouse numbers for phenotypic analysis were limited and prevented detailed physiological analysis of ciliary motility. However, histopathological analyses demonstrate severe PCD-associated phenotypes and genetic interactions. Additional cell biological and biochemical studies are required to determine how these protein complexes regulate mammalian ciliary motility, as well as how the mechanisms regulating spermiogenesis differ from those driving motile ciliogenesis. As a result, these studies will enable a better understanding of the genetic and molecular mechanisms underlying PCD inheritance and pathogenesis.

Methods

Mice. The nm1054, bgh, and Cfap54gt/gt mouse lines were maintained on the C57BL/6J (B6) and 129S6/SvEvTac (129) backgrounds as previously described26,27,29. Each mutation results in loss of transcript or protein expression, and each line was genotyped as previously described26,27,29. Since previous studies of the single mutations on B6, 129, and mixed backgrounds indicated that the only phenotype influenced by background strain was hydrocephalus, which was more severe and resulted in early mortality on the B6 background26,27,29, all analyses in this study were performed on double heterozygous or double homozygous mice on a mixed background. (B6 × 129)F1 double heterozygotes were generated by crossing a single heterozygote on the B6 background to a different single heterozygote on the 129 background. (B6 × 129)F2 double homozygotes were generated by intercrossing (B6 × 129)F1 double heterozygotes, and occasional (B6 × 129)F3 double homozygotes were generated by intercrossing (B6 × 129)F2 double heterozygotes. Double heterozygous (nm1054/+;bgh/+,
for all analyses at that age or later. Double homozygous animals dying prior to 8 weeks due to phenotype severity were used for histological and immunohistochemical analysis at the age of death when available. Wild type (WT) animals from the same generations were used as controls. Mouse numbers are detailed in Supplementary Table S1. Due to early mortality associated with double homozygotes, it was often difficult to collect or analyze samples, but the pathological phenotype was still assessable in the available samples. Other than male infertility, no sex-specific differences have ever been observed for the phenotypes associated with nm1054/+;Cfap54<sup>gt/gt</sup> or samples, but the pathological phenotype was still assessable in the available samples. Other than male infertility, no sex-specific differences have ever been observed for the phenotypes associated with nm1054/, bgh, or Cfap54<sup>gt/gt</sup>/CD<sup>R0</sup>. Therefore, male and female mice were pooled for analyses in this study. All experiments involving animals were performed in accordance with the Animal Welfare Act and National Institutes of Health (NIH) policies and were approved by the Sanford Research Institutional Animal Care and Use Committee.

**Histology.** Heads were immersion fixed in Bouin’s fixative until the bones were decalcified. Coronal slices were then cut through the maxillary sinuses and the lateral ventricles of the brain. Testes were immersion fixed in 10% buffered formalin. Fixed testes, brain slices, and sinus slices were embedded in paraffin, sectioned, and stained with Hematoxylin and Eosin (H&E) as previously described<sup>28–30</sup>. Stained tissue sections were analyzed by light microscopy using an upright Leica DM6000B microscope and Leica Aperio VERSA slide scanner. Spermatozoa were collected from the epididymis and diluted in phosphate buffered saline (PBS), following which they were spread onto slides, dried, fixed in methanol, and stained with the Camc differential stain kit (Cambridge Diagnostic Products, Inc.) as previously described<sup>27</sup>. Stained spermatozoa were analyzed by light microscopy on an Olympus IX71 inverted microscope.

Because some double homozygous mice died or required euthanasia at earlier ages, thereby resulting in different tissue and organ sizes, severity of brain and sinus phenotypes were quantitatively compared through a scoring system. Severity of the brain phenotype was scored based on the following criteria: 1—no ventricular dilatation; 2—slight dilatation or opening of the ventricle with no visible damage to adjacent tissue; 3—extensive dilatation into the center of the hemisphere with no visible damage to adjacent tissue; 4—extensive dilatation into the center of the hemisphere with damage to adjacent tissue (loss of ependyma, loss of white matter, cortical thinning, and/or hemorrhaging). Severity of the sinus phenotype was scored based on the following criteria: 1—no visible mucus in the maxillary sinus cavity; 2—traces of mucus accumulating in the maxillary sinus cavity; 3—extensive mucus accumulation filling large portions of the sinus cavity; 4—extensive mucus accumulation filling large portions of the sinus cavity with infiltration of neutrophils and/or red blood cells. Statistical significance was determined by one-way ANOVA.

**Immunohistochemistry.** Testes, brain slices, and sinus slices were fixed and prepared as described above for histology. The fixed tissues were embedded in paraffin, sectioned, and stained using the BenchMark XT automated slide staining system (Ventana Medical Systems, Inc.) as previously described<sup>28–30</sup>. Primary antibodies included a mouse acetylated tubulin antibody (Sigma Aldrich T6793) at a 1:10,000 dilution, a mouse γ-tubulin antibody (Sigma Aldrich T6557) at a 1:500 dilution, a rabbit SPAG6 antibody (Invitrogen PA5-58389) at a 1:500 dilution, a rabbit RSPH4A antibody (Sigma Aldrich HPA031198) at a 1:100 dilution, and a rabbit DNA11 antibody (Sigma Aldrich HPA021649) at a 1:300 dilution. Biotin SP-conjugated AffiniPure donkey anti-mouse (Jackson ImmunoResearch 715-065-151) and goat anti-rabbit (Jackson ImmunoResearch 111-065-144) secondary antibodies were both used at a 1:1,000 dilution. The slides were visualized by light microscopy using the upright Leica DM6000B microscope and Leica Aperio VERSA slide scanner, and high magnification images were acquired using the upright Nikon Eclipse Ni-E microscope with a 100 × objective and a Nikon DS-2MV camera.

The Aperio ImageScope software (v12.3.3.7014) was used for quantification of staining intensity on slide-scanned images of sinus sections. Regions of interest were drawn around cilia in fields where the ciliated epithelial cells were clearly in focus. Positivity, which is determined by the number of positive pixels divided by the number of total pixels (positive staining plus negative counter-stain pixels) in a selected region, represents staining intensity and was calculated for each image using the Positive Pixel Count v9 analysis algorithm with default manufacturer input settings. Statistical analysis was performed using GraphPad Prism 6 (v6.02). Outliers were identified using the ROUT analysis (Q = 1%), and any rare data point that was an order of magnitude different from all other values within a given cohort was removed from analysis. Statistical significance was determined by one-way ANOVA.

**Ciliary beat frequency (CBF) analysis.** Tracheae were dissected into Medium 199, Hank’s Balanced Salts and equilibrated at room temperature for 1 h, after which they were cut into rings using a Nikon SMZ1000 stereomicroscope. The rings were placed into fresh media and maintained at 28 °C using a thermal plate (Takai Hit TP-110RS05) for consistency throughout the experiment. CBF was analyzed on an Olympus IX71 inverted microscope using the Sisson-Ammons Video Analysis system as previously described<sup>26,27,29,40</sup>. Statistical significance was determined by one-way ANOVA.

**Data availability**

No large datasets were generated or analyzed during the current study. Representative data are included in this published article (and its Supplementary Information files), and all raw data is available from the corresponding author upon reasonable request.
References

1. Bustamante-Marin, X. M. & Ostrowski, L. E. Cilia and mucociliary clearance. *Cold Spring Harb. Persp. Biol.* 9, a028241 (2017).

2. Horani, A. & Ferkol, T. W. Advances in the genetics of primary ciliary dyskinesia: Clinical implications. *Chest* 154, 645–652 (2018).

3. Knowles, M. R., Zarirwala, M. & Leigh, M. Primary ciliary dyskinesia. *Clin. Chest Med.* 37, 449–461 (2016).

4. Lee, L. Mechanisms of mammalian ciliary motility: Insights from primary ciliary dyskinesia genetics. *Gene* 473, 57–66 (2011).

5. Lee, L. Riding the wave of ependymal cilia: Genetic susceptibility to hydrocephalus in primary ciliary dyskinesia. *J. Neurosci. Res.* 91, 1117–1132 (2013).

6. Olesc, C. et al. X-linked primary ciliary dyskinesia due to mutations in the cytoplasmic axonalen dop flag assembly factor PIH1D3. *Nat. Commun.* 8, 14279 (2017).

7. Paff, T. et al. Mutations in PIH1D3 cause X-linked primary ciliary dyskinesia with outer and inner dynein arm defects. *Am. J. Hum. Genet.* 100, 160–168 (2017).

8. Moore, A. et al. RPGR is mutated in patients with a complex X linked phenotype combining primary ciliary dyskinesia and retinitis pigmentosa. *J. Med. Genet.* 43, 326–333 (2006).

9. Wallmeier, J. et al. De novo mutations in FOXI1 result in a motile ciliopathy with hydrocephalus and randomization of left/right body asymmetry. *Am. J. Hum. Genet.* 105, 1030–1039 (2019).

10. Satir, P. & Christensen, S. T. Overview of structure and function of mammalian cilia. *Annu. Rev. Physiol.* 69, 377–400 (2007).

11. Becker-Heck, A. et al. The ciliated-coil domain containing protein CCDC40 is essential for motile cilium function and left-right axis formation. *Nat. Genet.* 43, 79–84 (2011).

12. Sugrue, K. F. & Zohn, I. E. Mechanism for generation of left isomerism in Ccdc40 mutant embryos. *PLoS ONE* 12, e0171180 (2017).

13. Brunner, S. et al. Overexpression of RPGR leads to male infertility in mice due to defects in flagellar assembly. *Biol. Reprod.* 79, 608–617 (2008).

14. Bukowy-Bieryllo, Z. et al. RPGR mutations might cause reduced orientation of respiratory cilia. *Pediatr. Pulmonol.* 48, 352–362 (2013).

15. Hong, D. H. et al. A retinits pigmentos GTPlase regulator (RPGR)-deficient mouse model for X-linked retinitis pigmentosa (RP3). *Proc. Natl. Acad. Sci. U.S.A.* 97, 3649–3654 (2000).

16. Rao, K. N. et al. Ciliopathy-associated protein CEP290 modifies the severity of retinal degeneration due to loss of RPGR. *Hum. Mol. Genet.* 25, 2005–2012 (2016).

17. Li, X. et al. Otitis media in sperm-associated antigen 6 (Spag6)-deficient mice. *PLoS ONE* 9, e112879 (2014).

18. Sapiro, R. et al. Male infertility, impaired sperm motility, and hydrocephalus in mice deficient in sperm-associated antigen 6. *Mol. Cell Biol.* 22, 6298–6305 (2002).

19. Teves, M. E. et al. Sperm-associated antigen 6 (SPAG6) deficiency and defects in cilogenesis and cilia function: Polarity, density, and beat. *PLoS ONE* 9, e107271 (2014).

20. Zhang, Z. et al. Deficiency of SPAG16L causes male infertility associated with impaired sperm motility. *Biol. Reprod.* 74, 751–759 (2006).

21. Zhang, Z. et al. Accelerated mortality from hydrocephalus and pneumonia in mice with a combined deficiency of SPAG6 and SPAG16L reveals a functional interrelationship between the two central apparatus proteins. *Cell Motil. Cytoskel.* 64, 360–376 (2007).

22. Loreng, T. D. & Smith, E. F. The central apparatus of cilia and eukaryotic flagella. *Cold Spring Harb. Persp. Biol.* 9, a028118 (2017).

23. Teves, M. E., Nagarkatti-Gude, D. R., Zhang, Z. & Strauss, J. F. 3rd. Mammalian axoneme central pair complex proteins: Broader roles revealed by gene knockout phenotypes. *Cytoskeleton (Hoboken)* 73, 3–22 (2016).

24. Zhao, L., Hou, Y., McNeill, N. A. & Witman, G. B. The unity and diversity of the ciliary central apparatus. *Philos. Trans. R. Soc. Lond. B. Biol. Sci.* 375, 20190164 (2020).

25. Zhao, L., Hou, Y., Picariello, T., Craige, B. & Witman, G. B. Proteome of the central apparatus of a ciliary axoneme. *Mol. Biol. Cell* 22, 690–701 (2011).

26. Lee, L. et al. Primary ciliary dyskinesia in mice lacking the novel ciliary protein Pcdp1. *Mol. Cell Biol.* 28, 949–957 (2008).

27. Sironen, A. et al. Loss of SPEF2 function in mice results in spermatogenesis defects and primary ciliary dyskinesia. *Biol. Reprod.* 85, 690–701 (2011).

28. McKenzie, C. W. et al. Enhanced response to pulmonary *Streptococcus pneumoniae* infection is associated with primary ciliary dyskinesia in mice lacking Pcdp1 and Spef2. *Cilia* 2, 18 (2013).

29. McKenize, C. W. et al. CFAP54 is required for proper ciliary motility and assembly of the central pair apparatus in mice. *Mol. Biol. Cell* 26, 3140–3149 (2015).

30. Finn, R., Evans, C. C. & Lee, L. Strain-dependent brain defects in mouse models of primary ciliary dyskinesia with mutations in Pcdp1 and Spef2. *Neuroscience* 277, 552–567 (2014).

31. Brown, J. M., Dipetrillo, C. G., Smith, E. F. & Witman, G. B. A FAP46 mutant provides new insights into the function and assembly of the Cid complex of the ciliary central apparatus. *J. Cell Sci.* 125, 3904–3913 (2012).

32. Dipetrillo, C. G. & Smith, E. F. Pcdp1 is a central apparatus protein that binds Cat(2+)-calmodulin and regulates ciliary motility. *J. Cell Biol.* 189, 601–612 (2010).

33. Dipetrillo, C. G. & Smith, E. F. The Pcdp1 complex coordinates the activity of dynein isoforms to produce wild-type ciliary motility. *Mol. Biol. Cell* 22, 4527–4538 (2011).

34. Zhang, H. & Mitchell, D. R. Cpc1, a Chlamydomonas central pair protein with an adenylate kinase domain. *J. Cell Sci.* 117, 4179–4188 (2004).

35. Bustamante-Marin, X. M. et al. Identification of genetic variants in CFAP221 as a cause of primary ciliary dyskinesia. *Hum. Genet.* 65, 175–180 (2020).

36. Yu, C. et al. Novel mutations in SPEF2 causing different defects between flagella and cilia bridge: The phenotypic link between MMAF and PCD. *Hum. Genet.* 139, 257–271 (2020).

37. Cindric, S. et al. SPEF2- and HYDIN-mutant cilia lack the central pair-associated protein SPEF2, aiding primary ciliary dyskinesia diagnostics. *Am. J. Respir. Cell Mol. Biol.* 62, 382–396 (2020).

38. McKenize, C. W. et al. Strain-specific differences in brain gene expression in a hydrocephalic mouse model with motile cilia dysfunction. *Sci. Rep.* 8, 13370 (2018).

39. Sironen, A., Shoemark, A., Patel, M., Loebinger, M. R. & Mitchison, H. M. Sperm defects in primary ciliary dyskinesia and related causes of male infertility. *Cell Mol. Life Sci.* 77, 2029–2048 (2020).

40. Sisson, J. H., Stoner, J. A., Ammons, B. A. & Wyatt, T. A. All-digital image capture and whole-field analysis of ciliary beat frequency. *J. Microsc.* 211, 103–111 (2003).
Acknowledgements
We gratefully thank Claire Evans for assistance with histology. The Sanford Research Histology & Imaging Core was supported by NIH CoBRE Grant P20GM103548. We also thank Carol Dao, who was supported by NIH grant R25HD097633, and Matthew Billion for technical assistance. This work was funded by Sanford Research.

Author contributions
C.W.M. bred the mouse lines, performed the experiments, and analyzed the data. L.L. designed the study, analyzed the data, and wrote the manuscript. All authors read and approved the final manuscript.

Competing interests
The authors declare no competing interests.

Additional information
Supplementary information is available for this paper at https://doi.org/10.1038/s41598-020-69359-3.

Correspondence and requests for materials should be addressed to L.L.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2020