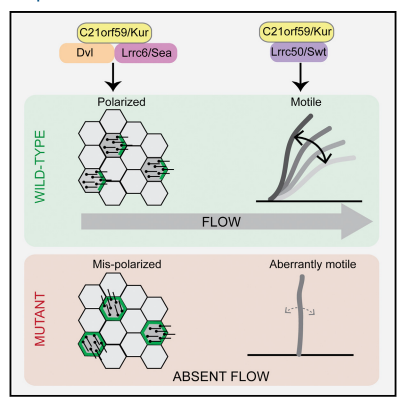
Cell Reports

c21orf59/kurly Controls Both Cilia Motility and **Polarization**

Graphical Abstract



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In Brief

Jaffe et al. report a dual role for the C21 orf59 protein in cilia motility and polarization. Zebrafish and Xenopus mutants revealed motility and planar cell polarity (PCP) as well as cilia positioning defects. C21orf59 made several interactions with PCP components and was required for proper Prickle2 localization.

Highlights

- · kurly (kur) mutants exhibit defects characteristic of motile cilia dysfunction
- c21orf59 is mutated in kur and is needed for dynein arm localization/cilia motility
- CRISPR/Cas9 with homologous recombination in Xenopus shows C21orf59 regulates PCP
- C21 orf59 interacts with various PCP components to correctly polarize motile cilia









c21orf59/kurly Controls Both Cilia Motility and Polarization

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http://dx.doi.org/10.1016/j.celrep.2016.01.069

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SUMMARY

Cilia are microtubule-based projections that function in the movement of extracellular fluid. This requires cilia to be: (1) motile and driven by dynein complexes and (2) correctly polarized on the surface of cells, which requires planar cell polarity (PCP). Few factors that regulate both processes have been discovered. We reveal that C21orf59/Kurly (Kur), a cytoplasmic protein with some enrichment at the base of cilia, is needed for motility; zebrafish mutants exhibit characteristic developmental abnormalities and dynein arm defects. kur was also required for proper cilia polarization in the zebrafish kidney and the larval skin of Xenopus laevis. CRISPR/Cas9 coupled with homologous recombination to disrupt the endogenous kur locus in Xenopus resulted in the asymmetric localization of the PCP protein Prickle2 being lost in mutant multiciliated cells. Kur also makes interactions with other PCP components, including Disheveled. This supports a model wherein Kur plays a dual role in cilia motility and polarization.

INTRODUCTION

In multicellular organisms, motile cilia are present on the surface of many cell types. Their primary function is the generation of fluid flow across epithelial surfaces and, in so doing, motile cilia are required for a variety of developmental and physiological processes. As such, abnormalities in cilia-driven flow generation result in primary ciliary dyskinesia (PCD), a condition that involves situs abnormalities, airway clearance defects leading to bronchiectasis, and infertility (Fliegauf et al., 2007; Norris and Grimes, 2012). In order to generate robust fluid flow, cilia must be properly motile. This requires a poorly understood pathway in which dynein arms are loaded into the cilium (Kobayashi and Takeda, 2012; Omran et al., 2008). Cilia must also be correctly positioned on the apical surface of cells (single-cell polarity), and they must be coordinately polarized across the entire tissue (tissue polarity). These roles involve the planar cell polarity (PCP) mechanism (Boutin et al., 2014; Hashimoto et al., 2010; Ohata et al., 2014; Park et al., 2008) and there also exists a feedback between cilia position and flow (Wallingford and Mitchell, 2011). To further our understanding of this link, it will be critical to discover novel components that impact cilia motility and ciliated cell PCP.

The zebrafish and Xenopus model systems have been used to understand the components and mechanisms required for ciliagenerated fluid flow (Becker-Heck et al., 2011; Hjeij et al., 2014; Kishimoto et al., 2008; Mitchell et al., 2007; Panizzi et al., 2012; Tarkar et al., 2013; Zhao et al., 2013). Here, we describe two zebrafish alleles of a mutant called kurly (kur) that disrupt c21orf59, a gene recently implicated in human PCD (Austin-Tse et al., 2013). We show that Kur is involved in cilia-associated developmental processes and is required for initiating cilia motility via recruitment of outer dynein arms (ODAs). Moreover, Kur interacts with multiple PCP components including Disheveled (Dvl) and is needed for correct cilia positioning in the zebrafish kidney. Experiments in Xenopus, including the generation of mosaic mutants by CRISPR/Cas9 and homologous recombination-based gene targeting, further support a role for Kur in cell polarity and tissue-level cilia polarity. Together, these results demonstrate that Kur plays a dual role in cilia motility and cilia positioning.

RESULTS AND DISCUSSION

kur Mutants Exhibit Defects Associated with Improper **Cilia-Mediated Flow Generation**

kur^{tj271} and kur^{tm304}, two zebrafish mutants isolated from forward genetic screens (Haffter and Nüsslein-Volhard, 1996), exhibit body curvature defects and kidney cysts in larval stages (Figure 1A) (Sullivan-Brown et al., 2008). The two mutations fail to complement in trans heterozygotes, indicating that they impact the same gene. Intriguingly, the penetrance of $\mathit{kur}^{\mathit{tm304/im304}}$ mutants varied with temperature, displaying progressively more severe defects when embryos were raised at 25°C, 28°C, and 32°C, respectively (Figures 1A, 1B, and 1D). Although kur^{tj271/tj271} mutants died as larvae, kur^{tm304/tm304} mutants raised at the permissive temperature (25°C) often resolved their mild body curvature defects by 5 days post fertilization (dpf) or earlier (Figures 1A and 1B). Though kidney cysts were apparent in kur^{tm304/tm304} embryos raised at permissive temperature, they



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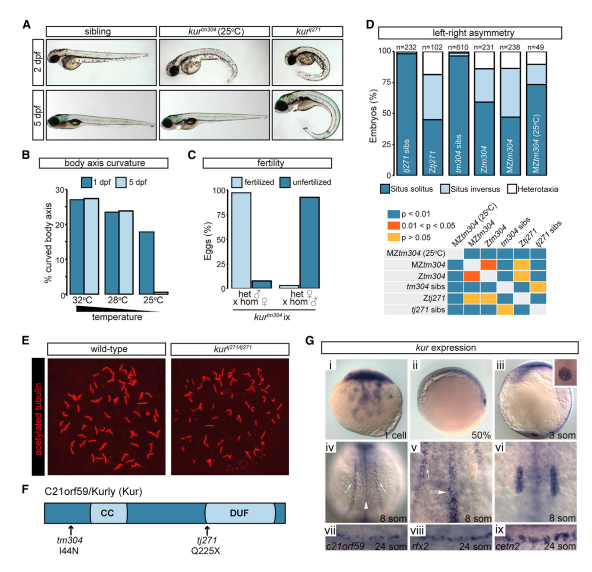


Figure 1. kur Mutants Harbor Mutations in c21orf59 and Exhibit Defects Characteristic of Defective Cilia Motility

(A) Lateral views of zebrafish sibling control embryos and kur^{tm304} and kur^{tl271} mutants at 2 dpf and 5 dpf. At 25°C, kur^{tm304} mutants exhibit early body axis defects that resolve by 5 dpf, whereas kur^{tl271} mutants do not resolve their body curvatures. Both mutants exhibit kidney cysts (white arrows), but the penetrance is variable in $kur^{tm304}ttm^{304}$ at 25°C.

- (B) Embryos from $kur^{*/tm304}$ intercrosses were raised at 32°C, 28°C, or 25°C and were scored for body curvature defects at 1 dpf and 5 dpf. While abnormalities were present in around 25% of embryos at 32°C and 28°C, mutants raised at 25°C exhibited fewer incidences of axis curvature at 1 dpf and almost all mutants had recovered at 5 dpf. The total number of embryos examined was 32°C, n = 292; 28°C, n = 332; and 25°C, n = 341. Shown is the average percentage of embryos with curved body axis from four biological replicates.
- (C) Fertility of kur^{tm304} adult mutants was assessed by scoring the number of fertilized eggs following homozygous female crosses to heterozygous males and vice versa. Shown is the average percentage of embryos fertilized from three biological replicates.
- (D) Percentage of embryos showing the indicated situs phenotypes. The embryos were processed by ISH for my17, ins, and fkd2 to mark the heart, pancreas, and liver, respectively, at 2 dpf. The statistical comparison of different genotypes was performed using chi-square analysis with p < 0.05 being considered to report significant differences.
- (E) Early somite-stage (ss) embryos were processed for immunostaining using acetylated alpha-tubulin antibodies to mark the ciliary axoneme. The confocal imaging of KV revealed the presence of cilia in sibling control embryos as well as kurtj271^{tj271} mutants.
- (F) Schematic diagram of C21orf59/Kur protein showing the approximate positions of the predicted CC domain and the DUF. The position and nature of the kur^{tm304} (l44N) and kur^{tj271} (Q225X) mutations are shown.
- (G) *kur* expression in zebrafish embryos at the stages indicated. *kur* is provided maternally at the 1-cell stage (i), followed by expression in the DFCs at 50% epiboly (ii). The expression is observed in KV (iii), pictured here at the 3-ss with a close-up of KV in the inset. During somitogenesis, *kur* is expressed in tissues with motile cilia including the pronephros (white arrows), the floorplate of the NT (white arrowheads), and the otic vesicles (iv–vi). Expression in the pronephros at the 24-ss overlaps with MCC markers *rfx2* and *cetn2* (vii–ix).

could still be raised to adulthood. Crosses between adult homozygous kurtm304/tm304 mutants suggested that male mutants are largely sterile, whereas female mutants are fertile (Figure 1C). Both kur alleles also exhibited defects in the establishment of left-right (L-R) asymmetry. In situ hybridization (ISH) for genetic markers of the heart (myl7), liver (fkd2), and pancreas (ins) at 48 hr post fertilization (hpf) revealed incidences of reversed or randomized organ positioning (heterotaxia) with respect to normal L-R asymmetry (Figures 1D and S1D-S1I). kurtj271/tj271 and kurtm304/tm304 (raised at 28°C, as is always the case unless otherwise specified) exhibited similar L-R defects, but embryos deficient for maternal and zygotic contributions of kur (MZtm304) were slightly more affected (Figure 1D), implying a partial role for maternally supplied kur gene product in the control of L-R patterning. Similar to body curvature defects, lowering the incubation temperature to 25°C resulted in less severe L-R defects in MZkur^{tm304/tm304} mutants (Figure 1D). Organ asymmetry is governed by motile cilia-driven fluid flow within a spherical and transient structure called Kupffer's vesicle (KV) (Amack, 2014). Asymmetric flow acts to initiate a left side-restricted Nodal cascade in which expression of the zebrafish Nodal gene southpaw (spaw) and its inhibitor lefty2 (lft2) are induced on the left (Nakamura and Hamada, 2012). In line with the defects in organ asymmetry, both kur mutants exhibited abnormalities in the asymmetric expression of spaw and Ift2 (Figures S1A-S1C and S1J). Together, these data place the primary L-R defect in kur mutants upstream of asymmetric gene expression, likely within the KV itself. Immunostaining demonstrated that cilia were present within the KV of kur^{tj271/tj271} and MZkur^{tm304/tm304} mutants (Figure 1E; data not shown). At the 8 somite stage (ss), we quantified the length of cilia in $kur^{tj271/tj271}$ mutants (n = 4 embryos) and found that they were not statistically different to wild-type (n = 3 embryos) controls (wild-type: $4.4 \pm 0.3~\mu m$ and $kur^{tj271/tj271}$: $4.4 \pm$ 0.1 μm). Moreover, the total number of cilia per KV was unchanged in mutants (wild-type: 60 \pm 12 and $kur^{tj271/tj271}$: 69 \pm 15). Thus, the generation of cilia themselves is not perturbed in kur mutants. However, since L-R patterning defects, kidney cysts, infertility, and body curvature abnormalities are all hallmarks of abnormal cilia function in zebrafish (Sullivan-Brown et al., 2008; Yuan and Sun, 2013), we hypothesized that kur was required for proper flow generation by impacting cilia motility and/or cilia polarity.

kur Disrupts c21orf59, a Human PCD Locus

We mapped the *kur* mutations to a region on chromosome 10 (Figure S2A). Sequencing of candidate genes revealed mutations in *c21orf59*, a gene of unknown function that is causative in some human PCD cases (Austin-Tse et al., 2013). PCD comprises disorders resulting from aberrant cilia motility (Zariwala et al., 2011), further supporting the hypothesis that *kur* is required for motile cilia flow generation. *c21orf59* encodes a highly conserved 290-amino acid protein with an N-terminal coiled coil (CC) domain and a C-terminal domain of unknown function (DUF) (Figure 1F). *kur*^{tm304} harbors a point mutation at amino acid 44 (I44N) that causes either decay or instability of the protein at higher temperatures (Figures 1F and S2B), thereby serving to explain the temperature-sensitivity exhibited by *kur*^{tm304}/tm304 mutants. By contrast, the *kur*^{ti271} mutation introduces a stop

codon at amino acid 225 (Q225X) that severely truncates the protein and results in a loss of *c21orf59* RNA (data not shown), most likely due to nonsense-mediated decay (Figure 1F).

Three lines of evidence confirmed that mutations in *c21orf59* cause the *kur* phenotypes. First, antisense morpholino oligonucleotide (MO)-mediated knockdown (KD) of endogenous *c21orf59* resulted in morphant embryos that phenocopied *kur* mutants. Morphants exhibited body curvature defects, kidney cysts, and L-R defects (Figures S2C and S2D). Second, injection of either human or zebrafish *c21orf59* RNA at the 1-cell stage rescued the body curvature defects of both *kur* mutants and *c21orf59* morphants; variable rescue of kidney cysts was also apparent (Figure S2C). Third, an independent study, which also knocked down *c21orf59*, reported similar gross phenotypes to those exhibited by *kur* mutants (Austin-Tse et al., 2013). Thus, loss-of-function (LOF) mutations in *c21orf59* underlie the phenotypes observed in *kur* mutants. We hereafter refer to *c21orf59* as *kur*.

Kur Is a Cytoplasmic Protein Required for the Initiation of Cilia Motility and Incorporation of ODAs in Cilia

In line with the phenotypic evidence that *kur* is needed for cilia function, *kur* RNA was present in tissues that exhibit motile cilia including the pronephric tubules and duct (PD), the dorsal forerunner cells (DFCs), KV, as well as the floor plate of the neural tube (NT) and otic vesicles (Figure 1G). In the pronephric tubules, *kur* exhibited the same staining pattern as *rfx2* and *cetn2*, markers of multiciliated cells (MCCs) (Figure 1G). Moreover, *kur* RNA was present at the 1-cell stage (Figure 1G), showing that the early embryo receives a maternal contribution of *kur*, a finding which can serve to explain the slightly more severe defects in MZ*kur*^{tm304/tm304} compared to Z*kur*^{tm304/tm304} mutants (Figure 1D).

Next, we analyzed cilia motility in kur mutants. Cilia in the kidney tubules and NT were almost entirely immotile in kur^{tj271/tj271} mutants, whereas kurtm304/tm304 mutants exhibited variable motility defects with greater numbers of immotile cilia observed when embryos were raised at higher temperatures (Movies S1, S2, and S3). Changing the incubation temperature of kur^{tm304/tm304} mutants from 25°C to 32°C did not hinder the motility of cilia that had already begun beating in the NT or kidney after 2-24 hr at the restrictive temperature (n = 4/4 embryos for each location). By contrast, lowering the temperatures from 32°C to 25°C did allow immotile cilia to become motile after 12 hr (Movie S4). A subset of immotile cilia in the KV of 28°C-raised kurtm304/tm304 mutants became motile after shifting to 25°C for 70 min (n = 2/6 embryos). After 240 min at 25°C, 50% of embryos harbored motile cilia in the KV (n = 3/6 embryos). Furthermore, while embryos raised at 32°C had immotile cilia in the NT and kidney, all embryos exhibited motile cilia in these locations after an overnight incubation at 25°C (n = 4/4 embryos). Together, these findings suggest that Kur is required for the initiation of motility, but not its maintenance. It is noteworthy that in Chlamydomonas reinhardtii, an upregulation of FBB18 (the Kur homolog) was observed upon deflaggelation (Li et al., 2004), while FBB18 was stabilized upon impairment of cilia motility (Austin-Tse et al., 2013), as might be expected for a protein whose function is to aid in the initiation of cilia motility. Ultra-structural analysis revealed a lack of ODAs in kur^{tj271/tj271}



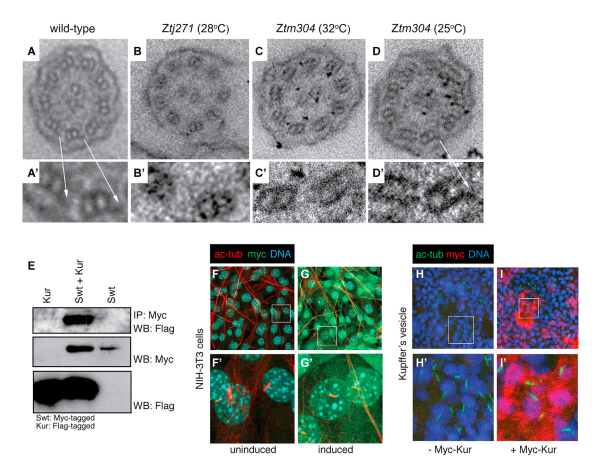


Figure 2. Kur Is a Cytoplasmic Factor Required for the Localization of ODAs to the Cilium

(A–D') Transmission electron micrographs (TEMs) showing the ultrastructure of pronephric cilia in cross-section. The arrows in (A) and (A') indicate ODAs that are missing in mutants (B–C'), but are restored in kur^{tm304/tm304} mutants raised at permissive temperature (D and D').

(E) IP of Myc-tagged Swt and subsequent Myc-Swt and FLAG-Kur western blot (WB) reveals physical association between Swt and Kur.

(F-G') NIH 3T3 cells transfected with inducible Myc-tagged Kur were immunostained for the ciliary axoneme (acetylated alpha tubulin [ac-tub]; red), DNA (Hoechst; blue), and myc (Myc-Kur; green). The Kur is not localized to the cilium, but instead localizes to the cytoplasm.

(H–I') Immunostaining of KV in control embryos or embryos injected with Myc-Kur. The cilia were marked by ac-tub (green) staining, DNA with Hoechst (blue), and Myc-Kur was localized with an anti-Myc antibody (red). The Myc-Kur, which rescues kur mutants, was present in cells of the KV, but appeared absent from cilia themselves.

and *kur*^{tm304/tm304} mutants (Figures 2A–2D), which can explain the motility defects. In accordance, *kur* morphants also exhibit ODA defects (Austin-Tse et al., 2013). Interestingly, human PCD patients harboring *C21ORF59* mutations exhibit both ODA and IDA defects.

Given these data, one possibility is that Kur controls cilia motility by participating in the assembly of ODAs in the cytoplasm and/or by assisting in the transport of ODAs into the cilium as is known to be the case for a small number of other proteins (Kobayashi and Takeda, 2012). Several lines of evidence support this view. First, epitope-tagged Kur localized to the cytoplasm in puncta, but not the cilium, in both NIH 3T3 fibroblasts (primary cilia) and KV of zebrafish embryos (motile cilia) (Figures 2F–2I), similar to what has recently been described for rat tracheal cells (Austin-Tse et al., 2013). The finding that the *Chlamydomonas* Kur homolog FBB18 was stabilized upon loss of cilia motility (Austin-Tse et al., 2013) may explain our observations that Kur was undetectable in normal KV motile cilia. Second.

kur^{tj271/tj271} mutants strongly phenocopy zebrafish switch hitter (swt) mutants that harbor a mutation in leucine-rich repeat-containing 50 (Irrc50), a homolog of Chlamydomonas oda7 that acts to transport intact dyneins into the cilium (Freshour et al., 2007). Third, immunoprecipitation (IP) experiments revealed a physical interaction between Kur and Lrrc50/Swt, suggesting that the proteins function in a common mechanism (Figure 2E). Thus, Kur is a cytoplasmic factor that, along with Lrrc50/Swt, most likely plays a role in either the cytoplasmic pre-assembly and/or transport to the cilium of dynein complexes, potentially acting as a chaperone in the transport of dynein complexes to cilia to initiate cilia motility. Indeed, like FBB18, several other chaperones are upregulated quickly upon deflagellation in Chlamydomonas including Hsp70, Hsp90, and Hsp22 (Li et al., 2004).

Kur Is Required for Cilia Positioning and MCC Polarity

During our analysis of *kur* mutants, we noted that cilia within kidney tubules were disorganized (Figures 3A-3F). The

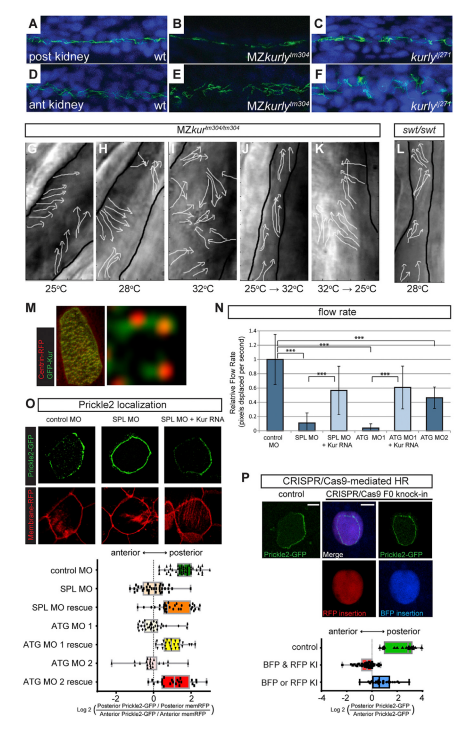


Figure 3. kur Mutants and Morphants Exhibit Polarity Defects in Zebrafish and Xenopus

(A-F) Staining of cilia axonemes (ac-tub; green) and nuclei (Hoechst; blue) in zebrafish pronephric tubules show cilia are present, but appear disorganized and mispolarized in mutants.

(G-L) Stills from live imaging of cilia. The cilia directionality is traced by the arrows in the MZkurtm304/tm304 (G-K) and the lokto237b/to237b mutants (L). The black outlines mark the edges of the tubules. The cilia in MZkurtm304/tm304 are progressively more disorganized at higher temperatures (G-I). Temperature shift experiments in kurtm304/tm304 mutants show that cilia do not become disorganized after shifting from permissive to restrictive temperatures (25°C \rightarrow 32°C) (J) and that cilia are able to recover their alignment defects by shifting from restrictive to permissive temperatures (32°C \rightarrow 25°C) (K). The swt/swt mutants exhibit dilated tubules, but normal cilia alignment (L), suggesting that tubule dilation does not cause alignment defects.

(M) Cytoplasmic localization of GFP-Kur in a Xenopus MCC showing an enrichment at the basal body and rootlets (inlay).

(N) Quantification of cilia generated fluid flow in control embryos and kur morphants. The average flow velocity measured by the displacement of dyed micro-beads along the skin of stage 29 Xenopus embryos is shown. The data represent three independent MOs and rescue experiments were performed with GFP-Kur RNA. The values are averages of at least ten flow lines from five embryos from two independent experiments. The error bars indicate SD and ***p value < 0.0002.

(O) Representative images and quantification of Prickle2-GFP (green) and membrane-RFP (red) localization in control and kur morphants (posterior is right). The box and whisker plots represent the log2 of the posterior fluorescent intensity divided by the anterior fluorescent intensity normalized for memRFP intensity.

(P) Representative images and quantification of Prickle2-GFP localization in mosaic F0 CRISPR/ Cas9-edited MCCs. The control embryos were injected with Cas9 protein and homology arms without kur gRNA. RFP positive MCCs indicate integration of RFP into the short kur allele, and BFP positive MCCs indicate integration of BFP into the long kur allele. The box and whisker plot represent the log2 of the posterior Prickle2-GFP fluorescent intensity divided by the anterior fluorescent intensity.

non-canonical Wnt or PCP mechanism has been implicated in the polarization of cilia (Wallingford and Mitchell, 2011). No gross convergent-extension (CE) defects, hallmarks of aberrant PCP, were observed in kur mutants, a result likely explained by kur not being expressed in a spatiotemporal manner that would be consistent with a role during gastrulation. We therefore hypothesized that Kur plays a role in PCP in ciliated cells where kur is expressed at high levels. To test this prediction, we first focused on the larval kidney. The kidney tubules of kur mutants were dilated and the cilia within were improperly oriented (Figures 3G-3I and S3). Tubule dilation is common in mutants in which cilia motility is disrupted, however, other mutants that exhibit cilia motility defects and tubule dilation, such as swt (Figures 3L and S3) and lok (Figure S3) (Sullivan-Brown et al., 2008), do



not exhibit polarization defects. Thus, polarization and motility are mechanistically separable, and the aberrant polarization of cilia in kur kidney tubules is not likely a secondary consequence of motility defects or tubule dilation. As with other phenotypes reported above, cilia polarization defects in kidney tubules were temperature sensitive in kurtm304/tm304 mutants, with more severe defects observed when embryos were raised at 32°C compared to 25°C (Figures 3G-3I and S3). Embryos raised at 28°C had an intermediate phenotype (Figures 3H and S3). Raising the temperature from 25°C to 32°C did not impact the polarization of cilia (compare Figure 3J to Figures 3G and 3I) even after an extended period at 32°C. By contrast, lowering the temperature from 32°C to 25°C did allow some recovery in cilia polarization, with more cilia being correctly polarized after the temperature was lowered (compare Figure 3K to Figures 3G and 3l). Thus, similar to cilia motility, kur function appears to be required to establish, but not maintain, cilia polarity.

Next, we asked whether Kur has a conserved role in cilia polarity and so turned to the ciliated skin of the Xenopus laevis embryo, where cilia polarization is particularly well characterized (Mitchell et al., 2009). The larval skin contains numerous MCCs, which possess cilia that beat in a coordinated fashion to drive directional fluid flow across the surface of the tissue. Although a GFP-tagged version of Xenopus Kur was found to be primarily cytoplasmic, it was also enriched at the ciliary basal body and adjacent rootlet that projects away from the basal body (Figure 3M). This localization was strikingly reminiscent of Disheveled (Dvl2), an intracellular mediator of PCP signals (Park et al., 2008). Xenopus contains two pseudo alleles of kur, designated as "long" and "short" (Matsuda et al., 2015). MO-mediated KD of kur in Xenopus using both splice site and start site MOs that target both pseudo alleles of kur (SPL MO and ATG MO1, respectively), resulted in a loss of cilia-driven fluid flow along the anterior-posterior embryonic axis, consistent with a defect in cilia motility (Figure 3N). This loss of flow was partially rescued by the addition of GFP-Kur-encoding RNA that was resistant to targeting by the MO (Figure 3N).

Next, to investigate the potential effect of Kur KD on PCP, we scored the localization of the PCP component Prickle2 by imaging an exogenously supplied Prickle2-GFP, which is known to asymmetrically accumulate on the posterior cell boundary of MCCs (Figure 30) (Butler and Wallingford, 2015). Depletion of Kur using three different MOs all resulted in uniform distribution of Prickle2-GFP relative to membrane-associated red fluorescent protein (memRFP) (Figure 3O), suggesting a loss in proper PCP. Importantly, this loss of Prickle2-GFP asymmetry was largely rescued by the expression of kur RNA (Figure 30). Polarity of MCC cilia arises from a combination of PCP coupled to a positive feedback mechanism driven by flow-mediated hydrodynamic forces (Mitchell et al., 2007, 2009; Park et al., 2008). Interestingly, a third MO (ATG MO2), targeting only one kur allele (the long allele), resulted in a weaker flow phenotype (Figure 3N) and maintained normal cilia beat frequency (control, 17.8 \pm 1.3 Hz and Kur ATG MO2, 18.4 \pm 2.8 Hz). Thus, in this morphant, the decrease in flow could result from either a loss of cilia polarity (within the cell) or a loss of MCC polarity (within the tissue). To distinguish between these possibilities, we assessed cilia polarity in ATG MO2 morphants, finding that while cilia were coordinately polarized within individual MCCs, there was a loss of coordinated polarity between MCCs (Figures S4D and S4E). In control embryos, 95% of cells orient their cilia in the posterior direction, thus facilitating the generation of a robust, posteriorly directed fluid flow. By contrast, *kur* KD (ATG MO2) led to a significant number of cells mispolarizing in the anterior direction (Figure S4E). This phenotype is consistent with a defect in PCP signaling and is similar to what has been reported for manipulations of the Wnt receptor Frizzled-3 and the PCP component Vangl2 (Figure S4E) (Mitchell et al., 2009).

To further validate the Kur depletion phenotype, we used CRISPR/Cas9-mediated genome editing in combination with homologous recombination (HR)-induced integration (Auer and Del Bene, 2014) to generate mosaic F0 mutant animals. A single guide (sg)RNA was designed to induce double strand breaks in both long and short alleles of the kur gene. We co-injected the gRNA/Cas9 with allele-specific DNA targeting constructs containing a fluorescent marker such that cells edited for the long allele integrated blue fluorescent protein (BFP) with a terminal codon and cells edited for the short allele integrated RFP with a terminal codon (Figures S4A and S4B). Gene targeting was confirmed by PCR (Figure S4C). In MCCs containing both BFP and RFP, we observed a complete loss of Prickle2-GFP asymmetric localization, similar to what was observed in the kur morphants (Figure 3P). Interestingly, cells that contained either BFP or RFP, indicating editing of only a single allele, gave an intermediate phenotype in which the Prickle2-GFP asymmetry was still present but considerably weaker (Figure 3P).

The loss of MCC cilia polarity coupled with the loss of Prickle2 asymmetry when *kur* is knocked down or mutated indicates that Kur has an important role in facilitating the establishment of cilia polarity. Furthermore, our results indicate that the function of Kur in cilia motility can be separated from its role in PCP.

Kur Makes Interactions with PCP Components and Represses the Canonical Wnt Pathway

Given the polarity defects exhibited by kur mutants and morphants, we searched for interactions between Kur and known PCP pathway components. kur zebrafish mutants strongly phenocopy inversin (inv) morphants, with both exhibiting kidney cysts, L-R patterning defects, and aberrant body curvature (Simons et al., 2005). Inv is thought to act as a molecular switch controlling the relative activities of canonical Wnt and PCP signaling by interacting with and modulating the function of DvI (Simons et al., 2005). We tested for a genetic interaction between kur and inv by injecting suboptimal doses of MOs targeting these genes. In this experiment, targeting of inv or kur alone produced minimal body curvature defects and few kidney cysts (Figures 4A and 4B). However, co-injection of these MOs at the same doses produced much more striking body curvature and kidney cyst phenotypes (Figures 4A and 4B) that were significantly different to single-injected embryos or uninjected controls (Figure 4C). This suggests that kur genetically interacts with inv to modulate common processes.

Given these genetic findings, we then asked whether Kur protein physically associates with members of the PCP pathway.

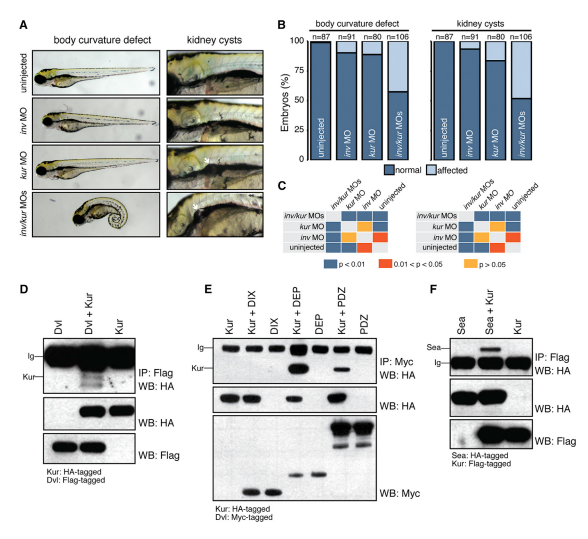


Figure 4. Kur Interacts with PCP Components

(A-C) Low concentration of inv or kur MO produce few defects, but, in combination, produce additive defects in body curvature and presence of kidney cysts. The white arrows show kidney cysts. The statistical comparison of different genotypes was performed using chi-square analysis with p < 0.05 being considered to report significant differences.

(D) IP of FLAG-tagged DvI and subsequent FLAG-DvI and HA-Kur western blot (WB) reveal physical association between DvI and Kur.

(E) IP of Myc-tagged Dvl domains and subsequent Myc-Dvl and HA-Kur western blot (WB) reveal physical association between Kur and the DEP and PDZ domains, but not the DIX domain of Dvl.

(F) IP of FLAG-tagged Kur and subsequent FLAG-Kur and HA-Sea western blot (WB) reveal physical association between Sea and Kur.

Using coimmunoprecipitation of tagged proteins, we found that Kur and Dvl physically interact (Figure 4D). Dvl consists of three distinct domains: a DIX domain that controls canonical Wnt signaling as well as DEP and PDZ domains that play roles in PCP. Utilizing Dvl deletion constructs, we found that Kur interacted with both the DEP and PDZ domains, but we could not detect an interaction between Kur and the DIX domain (Figure 4E). Moreover, Kur also associated with Leucine-rich repeat-containing 6 (Lrrc6/Seahorse [Sea]) (Figure 4F), another Dvl-interacting protein that controls the switch between canonical Wnt and PCP signaling (Kishimoto et al., 2008; Serluca et al., 2009). Intriguingly, Sea has also been implicated in the regulation of cilia motility by control of dynein arm assembly (Serluca et al., 2009; Zhao et al., 2013). In conclusion, the genetic and physical

interactions between Kur and known PCP components, together with the polarity defects of kur mutants and morphants, strongly supports a role for Kur in controlling polarity via PCP.

Finally, we reasoned that, if Kur promotes PCP signals, then it might act as a repressor of canonical Wnt signaling. Using luciferase readouts of the Super8XTopFlash reporter to assay Dvl's ability to activate canonical Wnt signaling, we found that Kur does indeed repress canonical Wnt signals (Figure S4F). While expression of Dvl alone led to strong induction of the reporter, co-expression of either Dvl/Kur or Dvl/Sea led to reduced reporter activity. Expression of Dvl, Kur, and Sea together further repressed activation of the canonical Wnt reporter (Figure S4F). Thus, Kur likely acts alongside Inv and Sea to promote PCP signaling over canonical Wnt signaling



via Dvl. Crucially, defects in these pathways upon *kur* LOF would be restricted to ciliated cells where *kur* is strictly localized, explaining the lack of gross canonical Wnt phenotypes or CE defects in *kur* mutants and morphants. Finally, we performed overexpression of *kur* in zebrafish by injection of wild-type RNA at the 1-cell stage. Injection of *kur* RNA induced moderate dorsalization (Figure S4G). In agreement with the Super8XTopFlash reporter assay, this suggests that Kur can act as a repressor of canonical Wnt.

Together, these findings from zebrafish and Xenopus demonstrate that Kur is involved in both cilia motility and polarization to control cilia-mediated processes such as the generation of fluid flow. As the role for the Kur homolog FBB18 in cilia motility is conserved in Chlamydomonas (Li et al., 2004), we suggest that this was the ancestral function of Kur and that its interactions with the PCP pathway evolved later along with multicellularity. Given the multitude of physical interactions Kur makes with proteins required for cilia motility and PCP, we propose that Kur acts as a scaffold to coordinate the activity of these two pathways. Indeed, by being involved in both cilia motility and ciliated cell PCP, Kur is a good candidate for a gene that mechanistically links these two processes, which are both critical for cilia function and the generation of polarized fluid flow. It will be important to find more components of these pathways, and to further characterize these processes at the molecular level, to aid in the understanding and diagnosis of human conditions such as PCD.

EXPERIMENTAL PROCEDURES

Zebrafish Mutants

Established zebrafish protocols were adhered to in accordance with Princeton University Institutional Animal Care and Use Committee. $kur^{t/271}$ and kur^{tm304} originated as part of a large-scale ENU mutagenesis screen (Haffter et al., 1996). They were generated in the TU strain and maintained by outcrossing to AB, WIK, and PWT. MZ kur^{tm304} embryos were obtained by crossing a $kur^{tm304/tm304}$ female with a $kur^{+/tm304}$ male, as homozygous males are sterile. Embryos were maintained at 28°C unless otherwise specified.

CRISPR/Cas9-Mediated HR in Xenopus laevis

Established Xenopus protocols were adhered to in accordance with Northwestern University policies (protocol number 2104-3527). kur mutant cells were generated in F0 animals using CRISPR/Cas9-mediated genome editing. A gRNA (GAGGCCACAGACCAGCTGAG[GGG]) that targeted both pseudo alleles of Xenopus kur (XI.4468) was designed (Chr2L:14433422-14433441 and Chr2S:19471649-19471627). HR facilitated "repair" was performed using \sim 400 bp homology arms such that BFP was inserted in frame into the long Chr2L pseudo allele (with a terminal stop codon) and RFP was inserted in frame into the short Chr2S allele (with a terminal stop codon) to mark cells that were edited (Figures S4A and S4B). Fertilized eggs were injected at the 1-cell stage with 500 pg Cas9 protein (PNAbio, CP01-250), 500 pg sgRNA, and 25 pg each of the circular plasmid DNA containing the homology arms and fluorescent marker (pKHR-Kur2L-BFP and pKHR-Kur2S-RFP). While the F0 animals are mosaic, the DNA containing the homology arms and the fluorescent markers has no promoter such that edited cells are the only cells that fluoresce (note: Kur is expressed in MCCs, therefore fluorescent expression specific to MCCs is further evidence of proper insertion). The presence of BFP and RFP insertions was verified using PCR to amplify fragments using BFP or RFP primers and primers outside of the homology arms (Figures S4A-S4C). Using this approach, we scored the fluorescent intensity (see below) of Prickle2-GFP in cells that were fluorescent for BFP, RFP, or both BFP and RFP.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, four figures, and four movies and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2016.01.069.

AUTHOR CONTRIBUTIONS

K.M.J., D.T.G., J.S.-R., T.-S.J.K., J.L.P., and N.F.C.M. designed and performed zebrafish and cell-culture experiments. M.E.W., S.K.K., and B.J.M. designed and performed *Xenopus* experiments. All authors interpreted the results. K.M.J., D.T.G., J.S.-R., B.J.M., and R.D.B. wrote the paper.

ACKNOWLEDGMENTS

The authors thank: Derrick Bosco, Cori Hasty, Phillip Johnson, Heather McAllister, and LAR staff for zebrafish care; Ray Habas for Dvl constructs; K. Horvath for Renilla constructs; and Dr. Gary Laevsky and the Molecular Biology Confocal Microscopy Facility, which is a Nikon Center of Excellence; and the National Xenopus Resource Centre for animals and expertise. This research was supported in part by a competitive research award to B.J.M. from the Eugene and Millicent Bell Fellowship Fund in Tissue Engineering and the Laura and Arthur Colwin Endowed Summer Research Fellowship Fund at the Marine Biological Laboratory in Woods Hole. The research was also supported by an NRSA grant to K.M.J. (#1F32HD060396-01A1), an NIH-NIGMS grant to B.J.M. (#2R01GM089970), and an NIH-NICHD grant to R.D.B. (#2R01HD048584).

Received: July 21, 2015 Revised: November 29, 2015 Accepted: January 22, 2016 Published: February 18, 2016

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