Bicaudal C, a novel regulator of Dvl signaling abutting RNA-processing bodies, controls cilia orientation and leftward flow

Charlotte Maisonneuve1,*, Isabelle Guilleret1,*, Philipp Vick2, Thomas Weber2, Philipp Andre2, Tina Beyer2, Martin Blum3 and Daniel B. Constam1,†

Polycystic diseases and left-right (LR) axis malformations are frequently linked to cilia defects. Renal cysts also arise in mice and frogs lacking Bicaudal C (BicC), a conserved RNA-binding protein containing K-homology (KH) domains and a sterile alpha motif (SAM). However, a role for BicC in cilia function has not been demonstrated. Here, we report that targeted inactivation of BicC randomizes left-right (LR) asymmetry by disrupting the planar alignment of motile cilia required for cilia-driven fluid flow. Furthermore, depending on its SAM domain, BicC can uncouple Dvl2 signaling from the canonical Wnt pathway, which has been implicated in antagonizing planar cell polarity (PCP). The SAM domain concentrates BicC in cytoplasmic structures harboring RNA-processing bodies (P-bodies) and Dvl2. These results suggest a model whereby BicC links the orientation of cilia with PCP, possibly by regulating RNA silencing in P-bodies.

KEY WORDS: Polycystic kidney disease, PCP, Flow, Nodal, SAM domain, K-homology

INTRODUCTION

In vertebrates, left-right (LR) asymmetry of the visceral situs is established by Nodal signals from the posterior notochord (PNC), also known as the ventral node in mammals (Brennan et al., 2002; Levin et al., 1995), Kupffer’s vesicle in zebrafish, or the gastrocoel roof plate (GRP) in Xenopus (Essner et al., 2002; Schweickert et al., 2007). These structures extrude motile cilia, which propel a leftward fluid flow to activate Nodal by an unknown mechanism specifically on the left side (Nonaka et al., 2002; Nonaka et al., 1998) (reviewed by Shiratori and Hamada, 2006). Scanning electron microscopy analysis established that node cilia are tilted towards the posterior pole, presumably because the basal bodies are displaced from the center of the convex apical plasma membrane to the posterior hemisphere (Cartwright et al., 2004; Nonaka et al., 2005; Okada et al., 2005). Theoretical and mechanical models suggest that the posterior tilt of the rotational axes is indispensable to the coordination of effective strokes and the generation of flow (Cartwright et al., 2004; Nonaka et al., 2005; Okada et al., 2005). Confirming this prediction, irregularities in the alignment of cilia in inv/inv mice carrying a mutation in the ankyrin repeat protein inversin (Mochizuki et al., 1998; Morgan et al., 1998; Yokoyama et al., 1993) are accompanied by a drastic reduction in nodal flow (Okada et al., 1999; Okada et al., 2005). Consistent with a role in cilia-driven flow, inversin localizes to node cilia, and situs defects in inv/inv embryos can be rescued by culture in administering an artificial leftward flow (Watanabe et al., 2003). However, an inhibition of flow in theory should randomize LR asymmetry, and it is unknown why inv/inv mutants instead display situs inversions. To resolve this conundrum, it is crucial to validate in independent models that LR asymmetry and vectorial fluid flow in vivo are linked to the planar orientation of cilia.

Besides perturbing LR asymmetry, mutations in inversin and other ciliary proteins give rise to polycystic kidney disease (PKD), and eventually to renal failure (Benzing and Walz, 2006; Fischer et al., 2006). The hallmark of this diverse group of genetic disorders is a progressive disruption of renal tubular morphology, preceded by defects in apicobasal protein sorting and misoriented divisions of renal epithelial cells (Benzing and Walz, 2006; Fischer et al., 2006; Germino, 2005; Wilson, 2004). In the kidney, cilia act as mechanosensors that stimulate Ca2+ channels in response to urinary flow (Prætorius and Spring, 2003). In addition, kidney cilia harbor the atypical cadherin Fat4, a conserved regulator of planar cell polarity (PCP) that is essential to orient renal cell divisions and suppress cyst formation (Saburi et al., 2008). These findings directly link cilia to PCP. However, the mechanisms by which cilia maintain normal polarity and the tubular architecture of renal epithelial cells remain poorly understood.

Several studies suggest that cilia and associated basal bodies mediate PCP at the expense of canonical Wnt signaling (reviewed by Gerdes et al., 2009), even though loss of cilia in mice lacking intraflagellar transport proteins (IFT) other than Ift88 (Jones et al., 2008) does not generally perturb classic readouts of PCP (reviewed by Eggenschwiler and Anderson, 2007). In the canonical signaling branch, Wnt proteins bind receptor complexes of frizzled and Lrp5 or Lrp6 that are endocytosed and recruit cytoplasmic dishevelled (Dvl1) to block a β-catenin destruction complex composed of axin, Apc and Gsk3β. Alternatively, to activate the PCP branch, complexes of Wnts and frizzled retain Dvl at the plasma membrane and reorganize the actin cytoskeleton by stimulating the small GTPases RhoA or Rac (reviewed by Kikuchi et al., 2009). The propagation of PCP between cells in vertebrates relies on the core PCP proteins frizzled (Fz3, Fz6 and Fz7), dishevelled (Dvl1, Dvl2 and Dvl3), prickle (Pk1 and Pk2), Van Gogh-like (Vangl1 and Vangl2), Celsr1, diversin and possibly inversin (reviewed by Simons and Mlodzik, 2008). A role for inversin is likely because...
depletion of inversin in *Xenopus* stabilizes cytoplasmic Dvl1 and thereby hyperactivates β-catenin while inhibiting polarized convergence extension movements during gastrulation (Simons et al., 2005). Similarly, the inversin-related protein diversin, an ortholog of the core PCP component Diego (Feiguin et al., 2001), promotes PCP signaling of Dvl1 during gastrulation, at least in part, by activating the β-catenin destruction complex (Moeller et al., 2006; Schwarz-Romond et al., 2002). When delivered ectopically to zebrafish pronephric duct, diversin can substitute for inversin to suppress renal cyst formation (Simons et al., 2005). Together, these observations led to the notion that both diversin and inversin function as a molecular switch between PCP and canonical Dvl signaling. β-catenin is also stabilized at the expense of PCP signaling during zebrafish gastrulation upon suppression of the ciliogenic kinesin Kif3a, or after depleting the basal body proteins Bbs1, Bbs4 or Mksks (Bbs6) (Gerdes et al., 2007). Similarly, mutations in ciliary (Kif3a, Ift88) or basal body proteins (Otfd) enhance Wnt/β-catenin signaling in the mouse (Corbit et al., 2008). However, the molecular machinery that links cilia to PCP remains ill-defined.

Apart from a potential role in promoting PCP, cilia themselves are also subject to regulation by PCP. Evidence that cilia respond to PCP comes from multiciliated *Xenopus* epithelial cells, in which docking of the basal bodies at the apical plasma membrane is directed by Dvl2 (Park et al., 2008), Vangl2 (Mitchell et al., 2009) and by the PCP pathway. The planar positioning of cilia and fluid flow in a manner reminiscent of that observed in *inv/inv* embryos, thus establishes LR asymmetry. Possibly, this is due to the hyperactivation of β-catenin while inhibiting polarized extension microscopy. For scanning electron microscopy, embryos were dissected 8.0 days post coitum in Hanks balanced salt solution, fixed at room temperature for 2 hours in 5% glutaraldehyde for 2 hours. For transmission electron microscopy, embryos were fixed for 2 hours in 0.1 M cacodylate buffer (pH 7.4) containing 4% paraformaldehyde and 0.1% trimethyl methylene blue. For transmission electron microscopy, embryos were stained with 2% osmium tetroxide, dehydrated in acetone and critical point dried. The samples were then coated with ~20 nm of platinum and examined using a transmission electron microscope.
(Schweickert et al., 2007). For numerical analysis, the positioning of cilia was counted in three or four embryos in which the ventral nodes were sufficiently shallow to clearly see more than 75% of the anchoring points.

**Video microscopy and image analysis**
The flow of fluorescent beads (1000-fold dilution of FluoSpheres carboxylate-modified microspheres, 0.2 μm, Invitrogen) in the cavity of the PNC/ventral node was recorded for durations of 2 to 10 seconds as described (Schweickert et al., 2007) at 20 to 50 frames per second (fps) and at 20X magnification on an Axioplan 2 imaging microscope equipped with an AxioCam HSi video camera. Flow at the GRP in dorsal explants of *Xenopus* embryos was recorded as described (Schweickert et al., 2007), except that bead solution was diluted 1:2500. Cilia movements in the marine PNC/ventral node were recorded at 63-fold magnification for 2 seconds at 100 fps. To visualize trajectories, 50 frames were analyzed using ImageJ software in combination with the MTrackJ plugin (http://www.imagescience.org/meijering/software/mtrackJ/).

**Cell culture and transfection**
HEK293T, COS1 and MDCK cells were cultured in DMEM (Sigma) supplemented with 10% fetal bovine serum (FBS, Sigma), glutamine 1% (Invitrogen) and gentamycin 1% (Invitrogen). Polarized MDCK cells were obtained after 10 days at confluence. HEK293T and MDCK cells were transfected using calcium phosphate. COS1 cells were transfected using DEAE-dextran. For stable transfection, HA-tagged BicC, ΔSAM and ΔKH constructs were subcloned into a pEF-IRESpac plasmid (Hobbs et al., 1998), and transfected into MDCK cells. Transfected cells were selected in concentrations of up to 2.5 μg/ml puromycin and expanded for 3 weeks.

**Luciferase assays using the Tcf/Lef-sensitive reporter TOPFLASH**
HEK293T cells were plated into 96-well dishes in triplicate at a density of 7.5 x 10^3 cells/well. After incubation for 24 hours, cells were transfected with TOPFLASH (0.1 μg/well) (Korinek et al., 1997) together with an expression plasmid using Lipofectamine 2000, lacZ and β-galactosidase activity. Western blot and indirect immunofluorescence analysis

**RESULTS**

**Heterotaxia and randomization of asymmetric Nodal signaling in Bicc1-/+ mice**

Using whole-mount in situ hybridization, we confirmed that mouse *Bicc1* is induced throughout the PNC/ventral node between embryonic day (E) 7.5 and E8.5 (Wessely et al., 2001) (see Fig. S2 in the supplementary material). To determine the function of *Bicc1* during development, we generated a targeted allele. *Bicc1* resides on mouse chromosome 10 B5.2 and gives rise to two splice variants A and B, differing by the presence or absence of exon 21. The sequence of the longer isoform is altered in the affected allele from mouse chromosome 10 B5.2 and gives rise to two splice variants A and B, differing by the presence or absence of exon 21. The sequence of the longer isoform is altered in the affected allele from mouse chromosome 10 B5.2 and gives rise to two splice variants A and B, differing by the presence or absence of exon 21. 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the supplementary material). The deletion introduced a frameshift after K131 in exon 12, resulting in premature termination of translation. Analysis by RT-PCR and sequencing confirmed that the mutant mRNA encodes a truncated protein that comprises KH1 but terminates after K131 (see Fig. S1C in the supplementary material).

Although adult \textit{Bicc1}\textsuperscript{+/-} heterozygotes were healthy and fertile, only 50% of the homozygotes developed to term (see Table S1 in the supplementary material) and thereafter died within 2-15 days, apparently as a result of renal failure (data not shown). \textit{Bicc1}\textsuperscript{--} newborns frequently displayed complete situs inversions (53%, \(n=18/34\)) or situs ambiguus (6%, \(n=2/34\); see Fig. 1A,B). During embryonic stages E13.5 to E15.5, we also detected ventricular septal heart defects (\(n=7/13\), Fig. 1B), and situs ambiguus was more frequent (41%, \(n=11/27\); Table 1), which is likely to account for the embryonic lethality. Besides confirming a role for BicC in kidney morphogenesis, targeted inactivation thus reveals a new function in LR axis formation.

The LR axis is patterned during early somite stages by the TGF\(\beta\) family member Nodal and its feedback inhibitors Lefty1 and Lefty2 (Shiratori and Hamada, 2006). Nodal signaling is confined to the left side by a leftward fluid flow that is propelled by motile cilia in the posterior notochord (PNC), also known as ventral node (Blum et al., 2007; Hirokawa et al., 2006). Inhibition of ectopic Nodal signaling on the right side is reliant on axial midline tissues (Shiratori and Hamada, 2006). To test whether BicC is essential for midline formation, we monitored the expression of \textit{Shh}, \textit{Brachyury} and \textit{Foxa2} mRNAs in axial mesoderm and the ventral neural tube. All of these markers were expressed normally in \textit{Bicc1}\textsuperscript{--} embryos (Fig. 1C; data not shown). Nevertheless, only eight out of 47 (17%) mutants expressed \textit{Nodal} and its target genes \textit{Lefty1}, \textit{Lefty2} and \textit{Pitx2} asymmetrically on the left side. In the remaining \textit{Bicc1}\textsuperscript{--} embryos, expression in lateral plate was bilateral (15/47), inverted (10/47) or absent (14/47; see Fig. 1D,E). Interestingly, all embryos
lacking Lefty2 mRNA in lateral plate mesoderm (n=7/17) instead showed ectopic expression of Lefty1 and/or Lefty2 behind the PNC. These results show that loss of BicC randomizes the sidedness of Nodal signaling without disrupting midline formation.

**BicC is necessary to correctly orient cilia and to generate nodal flow**

Next, we investigated whether laterality defects arose owing to perturbations of cilia morphogenesis or flow. Scanning electron and video microscopy analysis revealed no overt abnormalities in cilia length or motility (Fig. 2A-J, see also Movies 1, 2 in the supplementary material). However, although 82% (n=205/250) of cilia emanated from the posterior hemisphere in control PNC cells (Fig. 2D), this number was reduced to 38% (n=94/246) in BicC mutants (Fig. 2H). Thus, the majority of cilia in the mutants failed to become positioned correctly. Movies of beating cilia confirmed that the rotational axes of control cilia were tilted towards the posterior pole, whereas this polarity was perturbed in mutants (Fig. 2I,J). To assess the ability of cilia to generate flow, we recorded the movement of fluorescent beads in cultured PNC explants. Compared with the leftward flow in control embryos, flow in BicC mutants was less directed (see Movie 3 in the supplementary material) or, in a rare case (1/31), was even diverted to the right (see Fig. S3 in the supplementary material). Image analysis confirmed that trajectories in wild-type embryos were directed strictly to the left (Fig. 2K). By contrast, in BicC mutants, particles meandered considerably.
Individual trajectories pointed to the left, right, or were circling (Fig. 2L). The frequency distribution of trajectory angles revealed leftward movement of about 50% of control beads (Fig. 2M), whereas beads in BicC mutant embryos were transported in all directions (Fig. 2N). As a measure of flow quality, we have used $\rho$ to describe the scattering of trajectory directions. The maximum value $\rho=1$ indicates that all trajectories point in the same direction, whereas a $\rho$-value of 0 indicates uniform distribution in all directions. Control flow such as the one displayed in Fig. 2K reached a $\rho$-value of 0.71, whereas the BicC mutant flow was characterized by a $\rho$-value of 0.31 (Fig. 2L,N). Together, these results demonstrate that BicC is necessary to align the tilt of PNC cilia with the anteroposterior body axis as a prerequisite of leftward flow and LR axis formation.  

A role for BicC in generating cilia-driven flow is conserved in Xenopus

To quantitate flow dynamics, we depleted xBicC in Xenopus embryos, which can be analyzed in large numbers using advanced automated software tools (Schweickert et al., 2007). During flow stages (stages 16-19), xBicC was expressed in the gastrocoel roof plate (GRP), the floor plate, and the epithelial lining of the circumblastoporal collar (Fig. 3A,B), consistent with a conserved role for xBicC in regulating flow. To deplete xBicC in the GRP and floor plate, the morpholino oligonucleotides xBicC MO1 and MO2 (Tran et al., 2007) were injected into the marginal zone of four-cell embryos (Blum et al., 2009). If xBicC MO1/2 was injected on both sides (Fig. 3C,C') or unilaterally (Fig. 3D,D'),...
neural tube closure between stages 12-24 was delayed. Therefore, to analyze the left-sided marker genes *Xnr1* and *Pitx2*, injected embryos were cultured until stages 22-34. Embryos injected with DsRed mRNA or a control morpholino oligonucleotide (Co-MO) revealed normal left-sided gene expression patterns (Fig. 3E; see also Table S2 in the supplementary material). By contrast, injection of xBicC MO1/2 dose-dependently perturbed gene expression patterns in 20-60% of cases (Fig. 3E; see also Table S2 and Fig. 4 in the supplementary material). To elucidate the cause of LR asymmetry defects, xBicC morphants and controls were analyzed by scanning electron microscopy at stage 17/18. In representative control embryos, these stages were characterized by about 75% (71/94) of cilia being tilted towards the posterior pole in the flow-relevant areas (dotted lines) of wild-type GRPs (Schweickert et al., 2007) (Fig. 3F,G). By contrast, in xBicC morphants, posteriorly polarized cilia were reduced in number to less than 50% (40/83), with the percentage of misaligned cilia rising accordingly (Fig. 3J,K). These results suggest a conserved role for xBicC in aligning the planar polarity of ciliated cells with the anteroposterior body axis.

To monitor flow, dorsal explants of control- and MO-injected embryos were analyzed by adding fluorescent beads (see Movie 4 in the supplementary material). For the evaluation of flow, trajectories were selected that were directed and that exceeded the empirically selected threshold of maximum Brownian movement. Explants were grouped into four categories on the basis of flow phenotype. Category I represented robust leftward flow (ρ=0.66-1); category II (ρ=0.33-0.65) and category III (ρ<0.33) mildly and severely affected flow, respectively (see Fig. S5 in the supplementary material). If less than 25 particles per movie displayed velocities of >2.5 μm/sec (i.e. above threshold), embryos were classified as category IV (no flow). Trajectories of individual beads were generated by automated computation of gradient time trails (GTTs) (Schweickert et al., 2007). In representative control specimens (category I), a robust leftward flow was evident by the straight direction of trajectories towards the left (Fig. 3H). By contrast, beads in the GRP of representative xBicC morphants (category III) were frequently trapped by non-polarized cilia, resulting in non-directional, meandering trails (Fig. 3L, see also Movie 4 in the supplementary material). The frequency distribution of trajectory angles above the GRP, and the ρ values for flow in explants from xBicC morphants clearly differed from that in controls (Fig. 3L,M,N). Furthermore, the effects of xBicC MO1/2 were dose dependent, and a clear correlation between LR marker gene expression patterns and flow categories was obvious (compare Fig. 3E with 3N), confirming that altered gene expression resulted from aberrant flow.

The SAM domain recruits BicC to the periphery of P-bodies and downregulates BicC protein levels in polarized MDCK cells

Homologs of BicC in *Drosophila* and nematodes are implicated in regulating the localization or translation of target mRNAs (Chicoine et al., 2007; Eckmann et al., 2002; Mahone et al., 1995). To determine how BicC might regulate target genes, we visualized the subcellular localization of tagged BicC and truncated mutant forms lacking the KH- or SAM domains. Immunostaining of transfected HEK293T cells and confocal imaging detected BicC in discrete cytoplasmic foci that do not overlap with molecular markers of the ER, Golgi, endosomes or lysosomes (see Fig. S6 in the supplementary material). BicC formed similar structures in transfected COS1 cells (Fig. 4A).

Reconstruction of the 3D surface indicated that BicC foci formed tube- and vesicle-like structures around or adjacent to P-bodies that were marked by a GFP fusion of the decapping enzyme Dcp1a (Cougot et al., 2004) (Fig. 4A). In representative cells (n=10), 51±26% of the Dcp1a-GFP spots were at least partially coated by BicC-HA. A similar distribution was observed for mutant BicC lacking the KH domains (AKH, Fig. 4B), whereas deletion of the SAM domain gave rise to a diffuse cytoplasmic staining (Fig. 4C). Similarly in MDCK kidney epithelial cells, BicC and AKH localized in cytoplasmic foci, whereas ΔSAM staining was diffuse (Fig. 4D-G; data not shown). When MDCK cells expressing BicC were differentiated into fully polarized, ciliated epithelial cells (Fig. 4F-G, arrow), full-length BicC failed to accumulate, indicating that the combination of the KH and SAM domains negatively regulates BicC protein translation or stability under the conditions examined (Fig. 4H). Altogether, these experiments show that BicC is recruited to the periphery of P-bodies in a SAM domain-dependent manner.
### BicC inhibits Dvl signaling in the canonical Wnt pathway

Dvl, a positive regulator of the canonical Wnt pathway has been shown to multimerize in similar structures. (Bilic et al., 2007; Schwarz-Romond et al., 2007a; Schwarz-Romond et al., 2005). Prompted by our observations, we assessed whether Dvl and BicC would colocalize in cytoplasmic punctae. Consistent with previous reports, FlagDvl2 was detected in discrete puncta that arise by multimerization via the DIX domain (Schwarz-Romond et al., 2007a; Schwarz-Romond et al., 2007b). Interestingly, in Dvl2-expressing cells, P-bodies formed irregularly shaped clusters and, among 28±11 Dcp1a-GFP surfaces examined per cell (n=9), 7±3 (38±20%) abutted FlagDvl2 puncta (Fig. 5A). In cells co-expressing FlagDvl2 and BicC-HA, P-bodies remained dispersed, and, of those occupied by FlagDvl2 (40±9% of n=451 in seven cells), 60±7% cuddled to the BicC-HA surface (Fig. 5B-D). Furthermore, 35±6% of the FlagDvl2 puncta far from P-bodies (n=516/695 in seven cells) were at the BicC-HA surface (Fig. 5C,D, red arrowheads). To determine whether BicC directly interacts with Dvl2, transfected COS1 cells were analyzed by co-immunoprecipitation. Neither BicC nor ΔKH pulled down FlagDvl2, and their overexpression did not deplete FlagDvl2 (see Fig. S7 in the supplementary material). However, BicC dose-dependently inhibited FlagDvl2-mediated induction of TOPFLASH, a reporter of canonical Wnt signaling (Fig. 5E). This inhibitory effect was reduced upon deletion of the SAM domain, indicating that concentration in cytoplasmic foci is important. By contrast, the activity of ΔKH was similar to that of wild-type BicC. Furthermore, BicC failed to diminish induction of TOPFLASH by LiCl (Fig. 5F), indicating that BicC blocks signaling of Dvl upstream of Gsk3β. Taken together, these results suggest that BicC, via its SAM domain and independently of its KH domains, is concentrated in cytoplasmic platforms that inhibit canonical Dvl2 signaling.

To monitor the influence of BicC on the canonical Wnt pathway in vivo, Bicc1 mutants were crossed with transgenic BAT-gal reporter mice, which express a lacZ reporter of β-catenin/TCF signaling (Maretto et al., 2003). Whole-mount staining of BAT-gal embryos at E7.5-E8.0 showed that lacZ expression is reduced in the PNC compared with the adjacent primitive streak region in Bicc1 wild-type embryos, and that this local down-modulation of BAT-gal is impaired in the PNC of Bicc1−/− embryos (n=6/7; Fig. 6A), indicating that BicC attenuates canonical Wnt activity in the posterior notochord.

### DISCUSSION

This study reveals a conserved role for BicC in directing the planar orientation of cilia and leftward fluid flow during LR axis formation that has not been described in bpk and jcpk mice. Thus, our new Bicc1 allele unequivocally confirms for the first time the prediction of the flow hypothesis that misorientation of PNC cilia should randomize LR asymmetry. Furthermore, we have shown that the SAM domain concentrates BicC in cytoplasmic tube- and vesicle-like structures harboring P-bodies and cytoplasmic Dvl2, and that BicC inhibits Dvl2 signaling via the canonical β-catenin/TCF
Role of BicC during LR axis formation

Based on these results, we propose that BicC links cilia orientation to PCP signals, possibly by counteracting Dvl-induced activation of β-catenin.

In hpk mice, a point mutation disrupting one of the two splice variants of BicC1 causes ARPKD, whereas Jcpk mice carry a dominant mutant allele associated with an ADPKD-like phenotype (Cogswell et al., 2003). Our targeted mutation leads to a randomization of visceral tissue positioning, combined with a cystic phenotype in newborn kidney and pancreas that will be described elsewhere (I.G., unpublished observation). Transcription of this new allele might give rise to a truncated N-terminal peptide of 131 amino acids comprising the first KH domain. However, LR patterning and kidney defects are not observed in heterozygotes, suggesting that a gain-of-function is unlikely to account for LR defects. Corroborating this conclusion, morpholino-mediated knockdown confirmed that a homolog of BicC is also required for LR axis formation in *Xenopus*.

LR asymmetry is specified by Nodal activity, which is symmetrically induced in the PNC by the Notch pathway (Brenman et al., 2002; Krebs et al., 2003; Rayat et al., 2003), but is then biased by leftward flow to signal preferentially on the left side (Nonaka et al., 2002). Flow generation is thought to depend on the posterior tilting of node cilia (Cartwright et al., 2004; Nonaka et al., 2005; Okada et al., 2005). Consistent with this model, inhibition of flow in Bicc1 mutants was accompanied by a decrease in the number of correctly oriented cilia from 80% to 38%. A similar cilia misorientation phenotype has only been described in inv/inv mice (Okada et al., 2005). Even though cilia fail to align parallel to the anteroposterior axis, they remain fully motile and thus counteract the leftward fluid flow in both inv/inv and Bicc1−/− embryos. However, the outcome differs between these mutants. Poor flow in BicC1 mutants leads to situs randomization. By contrast, cilia malpositioning in inv/inv embryos is associated with situs inversions (Yokoyama et al., 1993). Although depletion of inversin can also randomize heart positioning in zebrafish, other aspects of LR asymmetry or cilia positioning have not been analyzed in that model (Otto et al., 2003). The present observations thus indicate that the inv/inv mutation probably has some additional function besides perturbing the planar orientation of cilia.

In zebrafish pronephros, depletion of inversin leads to cystic growth (Otto et al., 2003), which can be suppressed by injecting diversin, an ortholog of the core PCP protein Diego (Simons et al., 2005). Inversin selectively sequesters the cytoplasmic, but not the membrane-bound pool of Dvl1, suggesting that it promotes PCP by inhibiting the activity of Dvl in the Wnt/β-catenin pathway. Consistent with this model, hyperactivation of β-catenin signaling is sufficient to trigger cystic growth in the kidney of transgenic mice (Qian et al., 2005; Saadi-Kheddouci et al., 2001). However, whether an imbalance between PCP and Wnt/β-catenin signaling in inv/inv mutants also accounts for the misorientation of node cilia during LR axis formation is not known.

To assess whether BicC regulates Wnt signaling, we explored interactions between BicC and Dvl. We have shown that BicC downregulates expression of the BAT-gal reporter transgene in the PNC/ventral node, and that it can inhibit β-catenin/TCF signaling induced by Dvl2 in TOPFLASH reporter assays. By contrast, BicC failed to inhibit the induction of TOPFLASH by LiCl. Immunostaining showed that BicC forms cytoplasmic platforms, which accommodate foci of Dvl2 previously associated with PCP signaling, whereas the canonical Wnt pathway relies on a cytoplasmic pool of Dvl aggregating with axin during or after endocytosis of Lrp6 (Bilic et al., 2007; Kikuchi et al., 2009; Wallingford and Habas, 2005). Although the activities of these distinct pools of Dvl antagonize each other (for a review, see Wallingford and Habas, 2005), the mechanisms...
regulating pathway selection are poorly understood. Recent studies in zebrafish revealed that Dvl interacts with seahorse, a novel regulator of LR asymmetry and renal morphogenesis also known as leucine rich repeat containing 6 (Lrrc6), and that seahorse reduces the induction of Wnt/β-catenin target genes during gastrulation (Kishimoto et al., 2008). In addition, a genome-wide screen in Drosophila cells recently showed that PCP signaling depends on an association of Dvl with negatively charged phospholipids in the plasma membrane, and that this interaction is facilitated by a reduction of the intracellular pH by the sodium proton exchanger Nhe2 (Simons et al., 2009). It might be interesting, therefore, to assess in future studies whether canonical Wnt signals antagonize the expression or activity of Nhe2.

Here, we have directly demonstrated for the first time that BicC, through its SAM domain, forms cytoplasmic structures that can accommodate P-bodies and Dvl2. P-bodies are key regulators of mRNA surveillance, degradation, translational repression and RNA-mediated gene silencing (Cougou et al., 2004; Eulalio et al., 2007), but to our knowledge, they are not implicated in transmitting canonical Wnt signals. Our finding that the SAM domain promotes both BicC localization and the inhibition of Dvl2 indicates that P-bodies probably must at least communicate with BicC. In one possible scenario, BicC could mediate translational silencing of a Gsk3 inhibitor X by P-bodies (Fig. 6B, left box). The recent findings that β-catenin can be released from Gsk3β by the RNA helicase activity of p68 (Yang et al., 2006), and that p68 activates the let-7 miRNA precursor are consistent with such a model (Salzman et al., 2007). Alternatively, or in addition, BicC might promote PCP by blocking a direct inhibition of Gsk3β by Dvl2, the mechanism of which is currently unknown.

In Drosophila oocytes, BicC confines Oskar expression to the posterior pole to promote anterior cell fates (Mahone et al., 1995; Saffman et al., 1998). Posterior localization of oskar mRNA also depends on the Dcp1a homolog Dep1 (Lin et al., 2006), which has been detected in cytoplasmic foci of Drosophila oocytes and nurse cells together with Maternal Expression at 31B (Me31B) (Lin et al., 2006). Me31B, the homolog of the yeast decapping activator Dihlp, is localized to cytoplasmic foci by Drosophila BicC, apparently through a direct interaction (Chicoine et al., 2007; Kugler et al., 2009). Drosophila BicC has also been shown to repress translation of its own mRNA by recruiting the NOT3/5 component of the CCR4 deadenylase complex (Chicoine et al., 2007). CCR4 (also known as Twin) shortens the poly A tail length of oskar mRNA (Benoit et al., 2005), and in mammalian cells it stimulates the assembly of P-bodies (Andrei et al., 2005; Cougou et al., 2004; Zheng et al., 2008). These observations are consistent with our model that BicC regulates translational silencing of target mRNAs in P-bodies. However, we do not exclude that besides inhibiting cytoplasmic Dvl, BicC promotes PCP through additional mechanisms (Fig. 6B, right box). It will be interesting to determine in future studies the function of the novel P-body microenvironment defined by BicC, and whether BicC is needed to correctly canalize a response to Wnt signals at the level of RNA silencing.

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Competing interests statement

The authors declare that they have no competing financial interests.

Supplementary material

Supplementary material for this article is available at http://dev.biologists.org/cgi/content/full/136/17/3019/DC1

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Role of BicC during LR axis formation

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