The planar cell polarity gene \textit{Vangl2} is required for mammalian kidney-branching morphogenesis and glomerular maturation

Laura L. Yates\textsuperscript{1,†}, Jenny Papakrivopoulou\textsuperscript{2,3,†}, David A. Long\textsuperscript{2}, Paraskevi Goggolidou\textsuperscript{1}, John O. Connolly\textsuperscript{3}, Adrian S. Woolf\textsuperscript{4} and Charlotte H. Dean\textsuperscript{1,∗}

\textsuperscript{1}Mammalian Genetics Unit, Medical Research Council, Harwell, Oxfordshire OX11 0RD, UK, \textsuperscript{2}Nephro-Urology Unit, UCL Institute of Child Health, London, UK, \textsuperscript{3}UCL Centre for Nephrology, Royal Free and University College Medical School, London, UK and \textsuperscript{4}University of Manchester and Royal Manchester Children’s Hospital, Manchester, UK

Received July 28, 2010; Revised and Accepted September 8, 2010

The planar cell polarity (PCP) pathway, incorporating non-canonical Wnt signalling, controls embryonic convergent (CE) extension, polarized cell division and ciliary orientation. It also limits diameters of differentiating renal tubules, with mutation of certain components of the pathway causing cystic kidneys. Mutations in mouse \textit{Vangl} genes encoding core PCP proteins cause neural tube defects (NTDs) and \textit{Vangl2} mutations also impair branching of embryonic mouse lung airways. Embryonic metanephric kidneys also undergo branching morphogenesis and \textit{Vangl2} is known to be expressed in ureteric bud/collecting duct and metanephric mesenchymal/nephron lineages. These observations led us to investigate metanephroi in \textit{Vangl2} mutant mice, \textit{Loop-tail} (\textit{Lp}). Although ureteric bud formation is normal in \textit{Vangl2\textsuperscript{Lp/Lp}} embryos, subsequent \textit{in vivo} and \textit{in vitro} branching morphogenesis is impaired. Null mutant kidneys are short, consistent with a CE defect. Differentiating glomerular epithelia express several PCP genes (\textit{Vangl1/2}, \textit{Celsr1}, \textit{Scrib}, \textit{Mpk1/2} and \textit{Fat4}) and glomeruli in \textit{Vangl2\textsuperscript{Lp/Lp}} fetuses are smaller and contain less prominent capillary loops than wild-type littermates. Furthermore, \textit{Vangl2\textsuperscript{Lp/+}} kidneys had modest reduction in glomerular numbers postnatally. \textit{Vangl2\textsuperscript{Lp/Lp}} metanephiroi contained occasional dilated tubules but no overt cystic phenotype. These data show for the first time that a PCP gene is required for normal morphogenesis of both the ureteric bud and metanephric mesenchyme-derived structures. It has long been recognized that certain individuals with NTDs are born with malformed kidneys, and recent studies have discovered \textit{VANGL} mutations in some NTD patients. On the basis of our mutant mouse study, we suggest that PCP pathway mutations should be sought when NTD and renal malformation co-exist.

INTRODUCTION

The planar cell polarity (PCP) pathway, incorporating non-canonical Wnt signalling, directs embryonic convergent (CE) extension, polarized cell division and cell differentiation and also orients cilia (3–6). \textit{Vangl} (\textit{Van Gough}/\textit{Strabismus}) genes code for core PCP proteins. The transmembrane protein \textit{Vangl2} is expressed in the embryonic neural tube (7) and the mouse \textit{Loop-tail} (\textit{Lp}) allele comprises a \textit{Vangl2} mutation which encodes an unstable and mislocalized, and thus non-functional, protein product (8,9). The mutation is ‘semi-dominant’ with heterozygous mice being overtly healthy but characterized by their malformed tails; \textit{Vangl2\textsuperscript{Lp/Lp}}, however, causes a severe neural tube defect (NTD) called craniorachischisis (10). In this context, the normal function of \textit{Vangl2} is to facilitate, via Rho kinase (ROCK) and cytoskeletal rearrangement, cell-autonomous CE in axial mesoderm and neuroepithelium before neurulation (11). \textit{Vangl1\textsuperscript{Lp}}/\textit{Vangl2\textsuperscript{Lp}} compound heterozygous mice have severe NTDs (12) and subsets of individuals born with NTDs have been

†To whom correspondence should be addressed at: Tel: +44 1235841100; Fax: +44 1235841282; Email: c.dean@har.mrc.ac.uk

© The Author 2010. Published by Oxford University Press.
This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (http://creativecommons.org/licenses/by-nc/2.5), which permits unrestricted non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited.
reported to have either VANGL1 or VANGL2 variants (13–16) which may lead to dysfunctional gene products and thus contribute to the pathogenesis of neural defects (17). Genetic experiments in mice and zebrafish have shown that Vangl2 functions together with centrosomal/basal body-associated proteins, mutated in the Bardet–Beidl (18) and oral facial digital (19) syndromes, to drive embryonic CE. Furthermore, Vangl2 directs the orientation of primary cilia, thus allowing them to function optimally (5,6,20).

The embryonic precursor of the metanephros, which differentiates into the mature mammalian kidney, initiates around the fifth week of human gestation and embryonic day 11 (E11) in mice (21,22). The first phase of its morphogenesis is dominated by epithelial branching, starting with the outgrowth of the ureteric bud from the mesonephric duct, followed by arborization of the bud into the collecting ducts which drain forming urine into the renal pelvis from where it enters the ureter (21). Near the branching tips, mesenchymic mesenchyme converts into another epithelium, the nephron vesicle, which subsequently segments, elongates and differentiates into components including the glomerular podocytes, cells involved in ultrafiltration of blood in the first step of urine production (23,24), and the proximal tubules featuring actin-rich apical brush borders (25) which modify the ultrafilterate by reabsorption of specific molecules.

The embryonic lung is another organ which forms by branching morphogenesis, and we recently reported that Vangl2−/− mice have defective embryonic lung airway branching (26). Lung branching was also perturbed in Crash mice which have mutation of cadherin EGF LAG seven pass G-type receptor 1/Flamingo (Celsr1), another gene encoding a PCP protein (26). Intriguingly, in murine developing kidneys, Vangl2 protein is known to be expressed in ureteric bud epithelia (7) and in fetal podocytes (27). Although the exact pattern of branching is different in kidneys and lungs, the above observations led us to hypothesize that Vangl2 may have roles in kidney-branching morphogenesis and perhaps glomerular maturation. On the basis of the results of our current Vangl2 mutant mouse study, which indeed show that this PCP component is required for normal kidney morphogenesis, we suggest that VANGL mutations should especially be sought in humans who are found to have both an NTD and a malformed renal tract (28–30).

**RESULTS**

**Branching morphogenesis is disrupted in Vangl2LP kidneys in vivo and in vitro**

To determine whether kidney development was affected by Vangl2 mutation, we first examined the anatomical initiation of the metanephros. In transverse sections of both wild-type and Vangl2LP LP E11.5 embryos, we observed paired metanephric rudiments, each with a ureteric bud surrounded by mesenchymal cells (Fig. 1A and B). By E13.5, the normal ureteric bud has branched several times (21,22). Whole-mount immunostaining for cytokeratin was undertaken (Fig. 1C and D) to visualize both branching tips and collecting duct stalks (31). E13.5 wild-type organs contained significantly more ureteric bud branch tips than either heterozygous or homozygous Vangl2 mutant E13.5 metanephroi (Fig. 1E). None of the several tens of mutant embryos examined between E11 and late gestation had other ureteric defects such as duplication or dilatation (data not shown). We noted that both at E13.5 and in later gestation (E18.5) Vangl2LP/LP kidneys appeared smaller compared with wild-type littermates, especially in their antero-posterior axis (Fig. 1C, D and F). In fact, homozygous mutant E18.5 kidneys had a significantly (P < 0.01) lower length/width ratio (mean ± SEM; 1.48 ± 0.02, n = 3) than wild-type organs (1.74 ± 0.02, n = 3). Vangl2LP/LP embryos are known to have a shortened rostrocaudal axis, and this has been attributed to a primary CE defect (11).

Thus, a trivial explanation for the small mutant metanephroi would be that, in vivo, an abnormally shaped embryonic body might have physically constrained their morphogenesis. We therefore isolated late E11.5 metanephroi and followed their progress in serum-free organ culture (32). After 2 days, wild-type organs had undergone marked growth (Fig. 1G) whereas, in comparison, the progress of Vangl2LP/LP organs appeared retarded to variable degrees (Fig. 1H and I). At this stage, wild-type kidneys had significantly more ureteric bud branch tips versus Vangl2LP/LP organs (Fig. 1J). In other experiments, E12 organs, in which the bud had branched once, were cultured for 5 days and then co-immunostained for cytokeratin, to highlight the ureteric bud lineage, and Wilms tumour 1 (WT-1), to highlight glomerular podocytes (33) (Fig. 1K and L). At the end of the observation period, Vangl2LP/LP organs always appeared smaller than wild-type metanephroi (Fig. 1K and L). Mutant kidneys in vitro contained branch tips which frequently appeared distorted and wider than normal (compare Fig. 1 inserts in C and D and also frames K and L); we have previously noted a similar phenotype in the lungs of Vangl2LP/LP mice (26). Cultured organs from both genotypes contained clusters of WT-1-expressing cells, indicating that lack of Vangl2 function did not block formation of glomeruli in vitro (Fig. 1K and L). The detailed effects of Vangl2 mutation on glomerular maturation in vivo are explored below.

The late-gestation Vangl2LP/LP kidney has poor cortico-medullary definition, a hypoplastic medulla and mildly dysmorphic tubules

At E14.5, homozygous mutant metanephroi appeared smaller than those of wild-types (Fig. 2A–C). Inside E14.5 Vangl2LP/LP and Vangl2LP LP organs, as in wild-type littermate metanephroi, condensing renal mesenchyme was visualized around ureteric bud branch tips, and adjacent nephron vesicles were apparent (Fig. 2D–F). Despite the smaller size of E14.5 organs, both proliferation and apoptosis measured in situ across the whole metanephros was similar in wild-type and Vangl2LP/LP organs (Fig. 2G–P). By E18.5, the wild-type metanephros had undergone further growth and acquired its characteristic ‘kidney bean shape’, and its centre was occupied by a slit-like renal pelvis, the cavity which receives urine from the collecting duct system (Fig. 3A). Overall histology of E18.5 Vangl2LP/LP kidneys (Fig. 3B) appeared similar to their wild-type counterparts, whereas Vangl2LP/LP E18.5 kidneys lacked the normal curved profile and contained a renal pelvis which resembled a hole rather than the normal
elongated cavity (compare the outline of the organ in Fig. 3C with A). At this stage, the wild-type kidney contained a well-demarcated cortex and medulla (Fig. 3D). The outer cortex comprised ureteric bud branch tips whereas the cortex below this contained S-shaped bodies, which are immature nephrons beginning to segment into glomerular and proximal tubule epithelia (Figs. 4A–D), as well as proximal tubules and glomeruli (Figs 3D, G, 4E and F). The demarcation between the cortex and medulla was unclear in Vangl2Lp/Lp organs (Fig. 3F), with heterozygous kidneys (Fig. 3E) having an intermediate appearance between null mutant and wild-type littermates. Whereas proximal tubules and glomeruli appeared grossly normal in Vangl2Lp/+ organs (compare Fig. 3H with G), homozygous mutant kidneys contained subsets of glomeruli which were small or distorted (a feature examined in detail below) and some tubules contained attenuated brush borders as visualized by periodic-acid-Schiff (PAS) staining (Fig. 3I). The brush border of a differentiated proximal tubule cell comprises microvilli inserting into the apical zone of the epithelium, and both zones are rich in actin (25). Using phalloidin staining to visualize the f-actin cytoskeleton, some proximal tubule profiles in E18.5 Vangl2Lp/Lp kidneys had an irregular and discontinuous actin pattern versus proximal tubules in wild-type littermates (compare Fig. 3K with J). Although some tubules in fetal null mutant kidneys appeared dilated (Fig. 3I), an overt polycystic phenotype was not seen.

Several PCP molecules are expressed in the nephrogenic cortex and in differentiating podocytes

Vangl2 protein has previously been immunodetected in developing murine kidneys in ureteric bud branch tips, collecting ducts, nephron vesicles and forming glomeruli (S-shaped bodies and podocyte epithelia) (7,27). Because the results of

Figure 1. Early branching morphogenesis is disrupted in Vangl2Lp kidneys in vivo and in vitro. Haematoxylin- and eosin-stained transverse sections through whole E11.5 embryos showed similar appearances in both wild-type (A) and Vangl2Lp/+ (B) littermate embryos (n = 3 each genotype). Both had paired metanephric rudiments in which the ureteric bud (UB) penetrated into metanephric mesenchyme (MM). E13.5 whole z-stack images of wild-type (C) and Vangl2Lp/+ (D) kidneys immunostained for cytokeratin (green signal). Inserts in (C) and (D) show representative zoomed-in images of typical ureteric bud tips; Vangl2Lp/+ bud tips are frequently shorter and broader than in wild-type. (E) There was a significant (P < 0.01) reduction in numbers of ureteric bud branch tips in Vangl2Lp heterozygote (n = 12) and homozygote (n = 5) versus wild-type (n = 4) metanephroi. At E18.5, Vangl2Lp kidneys (F, right) appeared smaller and abnormally shaped in comparison to wild-type littermates (F, left). After 2 days in culture, E11.5 mutant explants (H and I) appeared variably growth-retarded versus wild-type organs (G). Furthermore, these cultured Vangl2Lp/+ metanephroi (n = 10) had significantly (P = 0.009) fewer bud tips in comparison to wild-type littermates (n = 7) (J). (K and L) To assess potential further differentiation in vitro, E12.0 metanephroi were cultured for 5 days and then immunostained for both cytokeratin (red) to highlight the arborizing ureteric tree, and WT1 (green) to highlight podocytes within glomeruli. After 5 days, wild-type explants (K; n = 3) appeared consistently larger than the Vangl2Lp/+ organ (L; n = 3). The ureteric tree in mutants appeared distorted with wider branch tips. Balls of WT-1-expressing cells, primitive glomeruli, formed in both genotypes. Scale bars: (A and B) 50 μM, (C and D) 125 μM, (F) 62.5 μM, (G–I) 63 μM and (K and L) 125 μM.
those studies are clear, we did not attempt to replicate them here. Instead, to gain a greater understanding of PCP pathway components in kidney development, we examined the distribution of Scribble 1 (Scrib), a protein mutated in the Circletail mouse (34), and Celsr1, a molecule known to be critical for lung branching (26). In wild-type E18.5 kidneys, which contain a spectrum of differentiating structures, Scrib was immunodetected (Fig. 4A, C and E) most prominently in the outer (nephrogenic) cortex where ureteric bud branch tips, mesenchymal condensates, vesicles and S-shaped bodies expressed the protein. Maturing podocytes were also positive. Celsr1 (Fig. 4B, D and F) was expressed in the cortex of the E18.5 kidney where it was most prominent in more mature components such as collecting duct stalks and S-shaped bodies (Fig. 4D) rather than bud tips and vesicles. Celsr1 was also detected in podocytes and adjacent proximal
tubules (Fig. 4F). No significant signal was obtained when primary antibodies were omitted (Fig. 4G and H). Previous studies have established the specificity of these antibodies by attenuation and/or mislocalization of the Vangl2 protein in Lp/+ embryos and the Celsr1 protein in Crsh/Crsh embryos (26,35,36). Finally, we used an established in vitro cell culture system (37) to seek the expression of a panel of PCP genes in both undifferentiated and differentiated mouse podocytes (Fig. 5). The in vitro differentiation is characterized by a change from proliferating cells with few processes, to quiescent

Figure 3. Histology of E18.5 kidneys. E18.5 (A–K) kidneys from wild-type (A, D, G and J), Vangl2Lp/+ (B, E and H) and Vangl2Lp/Lp (C, F, I and K) mice. At E18.5, homozygous (C) kidneys appeared smaller than wild-type littermate kidneys (A). Vangl2Lp/Lp kidneys are visibly misshaped, with a hypoplastic medulla (compare F with D) and the renal pelvis (*) lacks the normal slit-like shape seen in wild-type and Vangl2Lp/+ kidneys (compare C with A and B). Occasional enlarged and distorted proximal tubules with attenuated brush borders (pink) were observed in Vangl2Lp/Lp mice (I), but there was no overt cystic phenotype. Phalloidin staining highlights disruption to the organization of the actin cytoskeleton in Vangl2Lp/Lp proximal tubules; particularly at the apical surface (K), in comparison to the ordered cytoskeletal arrangement in Vangl2Lp/+ (J). Images are representative of at least three animals in each category. c, cortex; g, glomerulus; m, medulla; pt, proximal tubule. Scale bars: (A–C) = 200 μm, (D–I) = 25 μm and (J and K) = 5 μm.
cells with up to several actin-rich processes which grow from the cell body (Fig. 5A and B). In both conditions (Fig. 5C and D), podocytes expressed Vangl2 and well as Vangl1, Celsr1, Fat4, mouse prickle1 and mouse prickle2 (mpk1 and mpk2; 38,39).

E18.5 Vangl2<sup>Lp/Lp</sup> kidneys show abnormalities of glomerular maturation

The above observations prompted us to look closely at forming glomeruli in E18.5 kidneys. We conducted a careful analysis of glomerular morphology undertaken by an observer, blinded as to the genotypes of the organs, who assigned at least several tens of glomeruli in each kidney into one of four categories, according to the extent of maturation or morphological disruption, with ‘a’ being the most mature and ‘d’ being the most abnormal (Fig. 6A). In wild-type kidneys, over 95% of glomerular cross-sections displayed two or more capillary lumens (a marker of maturity), and most had more than two such lumens (Fig. 6B). By contrast, in late-gestation Vangl2<sup>Lp/Lp</sup> kidneys, <90% of all glomeruli contained two or more capillary lumens and, more strikingly, only half of these had more than two lumens. Moreover, in the null mutant kidneys, a small subset of glomeruli were malformed, with a distorted glomerular tuft (subtype d); such profiles were never observed in several hundreds of glomeruli scored in wild-type kidneys. The average diameter of Vangl2<sup>Lp/Lp</sup> glomerular tufts (i.e. the central ball of cells without the surrounding Bowman’s space and outer parietal epithelium) was significantly less than in wild-type kidneys (Fig. 6C). For Vangl2<sup>Lp/+</sup> kidneys, values were intermediate between wild-types and null mutants (Fig. 6B and C). We also sought proliferating cells in glomerular tufts by an immunohistochemical method (Fig. 6D) and found that the proportion of positive nuclei was significantly reduced in Vangl2<sup>Lp/Lp</sup> glomeruli versus either wild-type or Vangl2<sup>Lp/+</sup> kidneys (Fig. 6E). When comparing the most mature-looking glomeruli (i.e. subtype ‘a.’) in the three genotypes, immunostaining patterns for both synaptopodin, a molecule which regulates podocyte actin dynamics (40), and nephrin, a component of the junctions between podocyte foot processes (24),

Figure 4. PCP pathway components are expressed in E18.5 kidney. Immunostaining (brown) for Scrib (A, C and E) and Celsr1 (B, D and F) in E18.5 sections (note there is no counterstain in these sections). Scrib expression was prominent in the outer (nephrogenic) cortex (c in A) where it was found in diverse structures such as ureteric bud tips, mesenchymal condensates and vesicles (C). Scrib was also expressed in maturing podocytes located in the periphery of the glomerular tuft (E). Celsr1 expression was noted in collecting duct stalks and S-shaped bodies just below the nephrogenic zone (B and D). Expression of Celsr1 was also observed in proximal tubule and podocyte epithelia (F). Where primary antibodies were omitted, no significant staining was observed (G and H). Dashed line in (G) marks edge of kidney; below line is tissue. c, cortex; g, glomerulus; m, medulla; pt, proximal tubule. Scale bars: (A and B) 100 μM, (C and D), (G) 25 μM and (E, F and H) 12.5 μM.
appeared similar (Fig. 6D). As assessed by phalloidin staining, the crenellated pattern of f-actin evident in wild-type glomerular tufts, probably representing actin in both the basal zone of podocytes and in closely adjacent endothelia in forming capillary tufts (41), appeared less prominent in \( \text{Vangl2}^{Lp/Lp} \) glomeruli (Fig. 6F).

**Gene expression in E18.5 kidneys**

To assess whether the various morphological aberrations described above in E18.5 \( \text{Vangl2}^{Lp/Lp} \) kidneys corresponded to transcriptional changes, levels of transcripts were measured by quantitative PCR (32) in RNA extracted from whole organs. The genes analysed (Supplementary Material, Table S1) included those coding for: (i) transcription factors, growth factors/receptors and adhesion molecules implicated in metanephric-branching morphogenesis and nephron formation; (ii) proteins characteristically expressed by specific differentiated epithelia within collecting ducts, glomeruli, proximal tubules and loops of Henle; and (iii) ciliary/basal body proteins, mutations of which cause polycystic kidney disorders. Comparing levels of these mRNAs factored for levels of housekeeping transcripts between wild-type and \( \text{Vangl2}^{Lp/Lp} \) organs, none were significantly different (i.e. \( P < 0.05 \); data not shown). This analysis did, however, suggest that levels of four genes (\( \text{Shh} \), \( \text{Patched} \) 1, \( \text{Fgf7} \) and \( \text{Ift88} \)) tended to differ between the two genotypes. We therefore re-examined levels of these transcripts using an independent set of qPCR assays on cDNA from additional embryos. The results of these assays (Supplementary Material, Fig. S1) failed to confirm the previously observed trends.

**Postnatal \( \text{Vangl2}^{Lp/+} \) kidneys have a modest reduction in glomerular numbers**

As depicted in Figure 7 and data not shown, \( \text{Vangl2}^{Lp/+} \) kidneys of 1-month-old mice were found to have a modest but significant deficit in glomerular numbers, although immunostaining for synaptopodin and nephrin was normal, and there was no sign of glomerulosclerosis (scarring) as assessed by PAS staining (Fig. 7). Tubules showed no cyst formation. Finally, we found no difference in urinary albumin (Fig. 7H) or retinol-binding protein excretion (Fig. 7I) between 12-week-old wild-type and \( \text{Vangl2}^{Lp/+} \) mice. An increased level of albumin would be expected if there existed significant impairment of the glomerular permeability barrier (i.e. a podocyte function). An increased level of retinol binding protein would be expected if proximal tubule uptake of small molecules had been impaired.

**DISCUSSION**

**PCP genes and kidney development**

In the fetal kidney, the PCP pathway is considered to set a limit on the diameter of differentiating tubules because \( \text{Wnt9b} \) mutation causes cystic kidneys, perturbing the normal CE-like rearrangement of existing epithelial cells during tubule elongation and narrowing (42). Later in mouse kidney maturation, tubule diameter is limited by a predominance of polarized cell divisions aligned along the longitudinal axis of the tubule (43). This, in turn, requires the function of primary cilia (43), organelles which protrude from the apical epithelial surface and which transduce signals triggered by tubular fluid flow and perhaps also by yet-to-be defined components in the fluid (44). Oriented cell divisions and tubule elongation are disrupted by mutation of \( \text{Fat4} \), a PCP gene, the protein product of which is also detected in primary cilia (44,45). Intriguingly, the cystic kidney phenotype observed in \( \text{Fat4} \) mutants is accentuated when mice are additionally haplosufficient for \( \text{Vangl2} \) (45). Another link between renal cysts, PCP and ciliary sensing is provided by \( \text{Invs} \), a protein mediating a switch from canonical to non-canonical Wnt signalling (46). Expression of \( \text{Invs} \) is upregulated by fluid flow, and \( \text{Invs} \) mutations lead to cystic kidneys (46).

In our current study, we failed to find a cystic phenotype either in \( \text{Vangl2}^{Lp/Lp} \) and \( \text{Vangl2}^{Lp/+} \) fetal kidneys or in
1-month-old Vangl2 heterozygous kidneys. It remains possible, however, that the absence of Vangl2 might lead to cysts in the weeks after birth and that Vangl2 haploinsufficiency might cause kidney cysts later in adulthood. As assessed by PCR of late-gestation kidneys, the Vangl2<sup>Lp/Lp</sup> mutation did not significantly alter the expression of numerous transcripts (e.g. Bls1/4, Itf88, Invs, Mks1, Ofd1 and Pkd1/2), which encode ciliary proteins implicated in genetic cystic kidney diseases. Furthermore, we found that tubules in Vangl2<sup>Lp/Lp</sup> fetal kidneys did contain primary cilia (data not shown), as assessed by immunostaining for acetylated α-tubulin and IFT88 (19,47), although their orientation or numbers were not formally assessed.
Vangl2 is required for kidney-branching morphogenesis

Our data show that Vangl2 mutation strikingly impairs the arborization of the ureteric tree and that it is also associated with impaired glomerular maturation. This is the first demonstration that a core PCP component is required for normal development of both ureteric bud and metanephric mesenchyme-derived structures. The fact that developing epithelia in both lineages express Vangl2 protein is consistent with intrinsic functions in ureteric bud branches and nascent glomeruli (7, 27). Compared with time-matched wild-type organs, the growth restriction observed in explanted Vangl2 null mutant E11.5 rudiments appeared more prominent than that seen in either E12.0 explants or Vangl2 null mutant metanephroi in vivo. One possible explanation is that the serum-free tissue culture conditions used for such experiments may be more ‘stressful’ for younger rudiments and that any additional insult (in this case the Vangl2 mutation) results in a more severe in vitro phenotype.

Our immunohistochemistry studies which localized the PCP proteins Scrib and Celsr1 in diverse structures in the still-developing outer cortex of the late-gestation kidney concur with patterns of transcripts as reported in the Genito-Urinary Development Molecular Anatomy Project database [GUDMAP (48)]. We suggest that the branching impairment found in both Vangl2 heterozygous and homozygous metanephroi is an intrinsic defect operating within ureteric bud epithelia. In a quantitative PCR survey, we found no significant differences in levels of transcripts for growth factor signalling and HoxB7/Cre lineage-specific Cre-LoxP mRNA in Vangl2 null mutant E11.5 kidneys and the defects in lung branching reported by Yates et al. (26) in Vangl2 mutant mice. This study showed that branching defects were due to impaired tissue morphogenesis as a result of cytoskeletal disruption. In fact, the lung-branching defect in these mutants could not be rescued in vitro upon the addition of Fgf10, a growth factor that normally induces lung branching (26).

More likely, Vangl2 mediates branching in both the lung and kidney in a cell-autonomous manner acting, as it does in neural tube closure (11), through ROCK-mediated cytoskeletal actions. ROCK indirectly inhibits actin filament depolymerization (49), and when perturbed, over time, actin monomers that are required to continue actin polymerization for migration and cell movement become limited. Indeed, the branching defects observed in Vangl2 embryos showed that branching morphogenesis in the metanephros is sensitive to ROCK activity (50–52). While we did not detect any marked alteration in pattern of f-actin in Vangl2 mutant ureteric bud branch tips (data not shown), it was notable that the apical f-actin pattern appeared irregular in mutant proximal-like tubules. Future studies should use conditional deletion Cre-LoxP strategies, for example using the ureteric lineage-specific HoxB7/Cre line (53), to confirm the hypothesis that the kidney-branching defect is caused by deficiency of Vangl2 within arborizing epithelia.

Despite the above insights into tubule diameter regulation, the roles, if any, of the PCP pathway on other metanephric processes, such as initiation of the kidney, ureter formation, branching morphogenesis and glomerular morphogenesis, have been unclear. We found that, at least in the C3H/HeH background studied, Vangl2 deficiency did not prevent ureteric bud formation nor aggregation of intermediate mesodermal cells, which together comprise the metanephric rudiment. Nor did Vangl2 deficiency lead to ureteric duplication or ureter dilatation, the latter structure derived from the distal part of the ureteric bud.

Figure 7. Histology of adult Vangl2+/+ kidneys. (A and B) PAS staining of 4-week-old Vangl2+/+ mice kidneys show a normal appearance with no scarring. Expression patterns of synaptopodin (green in C and D) and nephrin (green in E and F) were similar in wild-type and heterozygous glomeruli. (G) Reduced glomerular numbers per kidney in Vangl2+/+ mice (n = 4) versus wild-type littermates (n = 4). No significant differences in the ratio of urinary albumin/creatinine (H) or retinol-binding protein (RBP)/creatinine (I) were found between Vangl2+/- and Vangl2+/+ mice. Scale bars = 15 μm.
The retardation of metanephric-branching morphogenesis which we measured in the first days of Vangl2Lp/Lp metanephric development likely explains why the late-gestation null-mutant kidney has a poorly developed medulla and renal pelvis because bud-derived collecting ducts normally populate the medulla and contribute to the shape of the pelvis. As assessed by qPCR, there were no differences between null mutant and wild-type late-gestation kidneys in expression of differentiation markers of specific tubule epithelia (e.g. Ggt1 for proximal tubules, Umod for loops of Henle and Aqp2 for collecting ducts). These observations are consistent with the idea that Vangl2 is not required for terminal differentiation of tubules but instead modulates their morphogenesis.

**Humans with NTDs sometimes also have kidney malformations**

Historically, individuals born with NTDs often died from renal failure, which was perceived as arising from excessive blood pressure on the kidneys from impaired urine flow combined with bacterial infection ascending from urine pooled in poor-emptying urinary bladders. Some individuals with NTDs, however, are born with kidney malformations such as agenesis (unilateral absent kidney), gross defects in renal shape (horse-shoe kidney) and ureteric duplication (duplex kidneys) (28–30). These historical clinical observations are consistent with the idea that both the renal malformation and NTD arise from common, perhaps genetic or environmental, causes. On the basis of our observations in mice, we suggest that the subset of individuals with NTDs who also have kidney malformations may be found to have VANGL or other PCP gene mutations.

**Vangl2 is required for kidney glomerular maturation**

We also found that the tufts of glomeruli in Vangl2Lp/Lp late-gestation kidneys were smaller than their wild-type littermates. This was associated with a decrease in percentage of proliferating cells within the tufts. Glomerular tufts initially comprise only differentiating podocytes and they then become invaded by blood capillary loops and pericyte-like mesangial cells (23,54). Our methodology could not distinguish whether the proliferation defect in Vangl2Lp/Lp tufts affected a specific type of cell although proliferation did appear to be reduced in the periphery of the tuft (compare null mutant with wild-type in Fig. 6D) where podocytes are found. The cause of the proliferation defect is uncertain, but we note that Vangl2 has been implicated in the expansion of muscle (4) and neural (3) stem cells. In the latter example (3), late-gestation Vangl2Lp/Lp brains showed a significant reduction in dividing progenitors even though earlier in gestation, exactly as found in E14.5 Vangl2 mutant metanephroi, the numbers of proliferating cells were the same as in controls. Thus, we speculate that Vangl2 may regulate the numbers of glomerular precursor cells within the primitive nephron. Consistent with this interpretation of the Vangl2Lp/Lp glomerular tuft size in relation to podocyte proliferation, we found that isolated differentiating podocytes express an array of PCP transcripts, including Vangl2.

Podocytes are extremely plastic cells and as such they are able to alter their shape via molecular regulation of their intracellular cytoskeleton. Cytoskeletal disruption caused by injury or genetic disorders of glomerular function leads to misshaping or absence of foot processes (24,55). Unless this process can be reversed, it results in leakage of protein into the glomerular filtrate, glomerular scarring and eventually, chronic renal failure. Therefore, an understanding of the molecular mechanisms regulating podocyte architecture is likely to prove valuable for treatment of glomerular dysfunction. Synaptopodin plays an essential role in the regulation of podocyte actin dynamics and RhoA is a target of the molecule (40). However, we noted no overt abnormality in the pattern of synaptopodin protein in Vangl2 mutant glomeruli. In a recent cell culture study, Babayeva et al. (27) reported that siRNA depletion of podocyte Vangl2 reduced the number of cell projections, decreased actin stress fibres and impaired cell motility. How this result relates to the glomerular phenotype we observed in Vangl2Lp/Lp kidneys is not clear although it provides proof of principle that Vangl2 plays an active role in podocyte biology. Ichimura et al. (56) recently reported that immature podocytes express primary cilia in vivo but that these organelles are lost with full differentiation. In future, it would be interesting to determine whether the Vangl2 protein can be found associated with these structures.

We note that endothelia in developing glomeruli have been reported to express Vangl2 transcripts (48) and, if capillary migration were to be compromised, this may help explain why Vangl2Lp/Lp tufts have a deficit in capillary lumens. The apparent attenuation of f-actin we observed in mutant fetal glomeruli may represent a combined aberration in the cytoskeleton of both podocytes and closely adjacent endothelia (41). According to our current data and those from GUDMAP (48), podocytes also express Vangl1 transcripts. Given the fact that, in other tissues, there is genetic interaction between Vangl1 and Vangl2 (12), we predict that double mutants would have a much more severe glomerular phenotype than that displayed by Vangl2 mutants in the current study. Finally, we note that upregulation of canonical Wnt/β-catenin signalling has been implicated in the pathophysiology of acquired proteinuric glomerular diseases (57). In future, a definitive exploration of the roles of Wnt canonical and non-canonical/PCP pathways in glomeruli genes will require their selective deletion in these cells at different stages of glomerular differentiation using Cre-LoxP technology (58).

**Vangl2 and nephron numbers**

In mice, new nephrons continue to be generated during the first week after birth (22,59). Non-viability of homozygous Vangl2 mice due to severe NTDs precluded analyses of their kidneys beyond the fetal period. Instead, we analysed kidneys of Vangl2Lp/Lp mice 4 weeks after birth. These animals are overtly healthy although they have a deformity consistent with their name, Looptail. Our data show that haploinsufficiency of Vangl2 results in a modestly decreased nephron number. Our method for quantifying glomerular numbers relied on acid-mediated dissociation of the kidney (60). While it is possible this method may be less accurate...
than stereology (61,62), any directional bias would be equal in wild-type and \textit{Vangl2}Lp/+ samples because the latter displayed no tissue abnormalities such as cyst formation which might affect the efficiency of dissociation. Because we demonstrated that \textit{Vangl2}Lp/+ as well as homozygous \textit{Vangl2} mice have a branching defect, the reduction in final glomerular numbers may be caused by a relative lack of branch tips available to induce mesenchymal cells to differentiate into nephrons.

It is striking that human populations contain otherwise healthy individuals with a very wide range of glomerular numbers (~1–2 million) per kidney (62). On the basis of the results of animal experiments (60,61) and human studies (63), it is likely that both environmental influences (e.g. the diet a mother eats when pregnant) and genetic factors (e.g. polymorphisms of genes expressed in the developing kidney) together determine nephron number in an individual. Moreover, people with essential hypertension probably have a congenital nephron deficit (62) and animals born with too few nephrons are prone to arterial hypertension later in life (61). Further studies will be required to determine whether\textit{Looptail} mice are prone to hypertension or susceptible to glomerular scarring as they age. However, we suggest that mutation or variants of \textit{VANGL} and other PCP genes should be considered as possible causes of the variable glomerular numbers observed in human kidneys.

**MATERIALS AND METHODS**

All reagents were obtained from Sigma Chemical (Poole, UK) unless otherwise stated.

**Histology and immunostaining**

The \textit{Looptail} mice (Jackson Laboratories, Bar Harbour, USA) contain a point mutation which causes a serine to asparagine transition at position 464 resulting in loss of \textit{Vangl2} function (10,64). Mice were maintained on a C3H/HeH background. \textit{Vangl2}2+/− homozygotes and heterozygotes were identified by craniorachischisis and looped-tail phenotypes, respectively, or genotyped using a pyrosequencing assay to amplify the mutation directly (forward: 5′-Biotin GTCTGGCCGCTTTCAAGAGGA; reverse: NNNGGCCAAACAGTGGAACCTTGG; sequence: CAGTTGACCTTGGTA); wild-type littermates were used as controls. Kidneys were fixed in 4% paraformaldehyde, dehydrated, wax-embedded and sectioned at 5 µm. For some antibodies, 10 µm cryosections were used. In some experiments, the following stains were used: PAS to visualize all basement membranes and also proximal tubule brush borders; haematoxylin to visualize nuclei; and eosin to define cytoplasm. Immunohistochemistry was performed for the following antibodies to: Celsr1 (gift from C. Formstone, King’s College London), nephrin (Progen Biotechnik, Heidelberg, Germany), proliferating cell nuclear antigen (PCNA; BD Biosciences, Oxford, UK), phospho-histone H3 (pH3; Millipore, Dundee, UK), Cleaved caspase-3 (Cell Signalling Technology, Danvers, MA, USA), rhodamine-conjugated phalloidin (Invitrogen, Paisley, UK), Scrib (Santa Cruz Biotechnology, USA), synaptopodin (Acris, Herford, Germany) and WT-1 (Acris). Bound antibodies were detected with either Alexa Fluor 488 or 594 secondary antibodies or appropriate horse-radish-conjugated antibodies followed by diamobenzidine detection. Cell nuclei were visualized by staining with 4′,6-diamidino-2-phenylindole (DAPI, HCl). As negative controls, primary antibodies were omitted.

**Explant culture and whole-mount metanephros staining**

Freshly isolated metanephiroi were collected from \textit{Vangl2}Lp homozygotes and wild-type littermates. Some metanephiroi were used immediately for staining but others were cultured for up to 5 days as described (32). Metanephiroi were fixed and incubated with anti-pan-cytokeratin (Sigma) and/or anti-WT1, followed by appropriate Alexa Fluor 488 or 594 secondary antibodies. Explants were imaged using confocal microscopy and the number of tubules in each explant counted.

**Urine analysis**

Twelve-week-old male \textit{Vangl2}2+/− heterozygotes and wild-type littermates were individually housed in metabolic cages for 24 h for urine collection. Urine concentration of albumin (Exocell Incorporation, Philadelphia, PA, USA) and retinal-binding protein-4 (Alpco Diagnostics, Salem, OR, USA) were assessed using commercially available kits according to the manufacturer’s protocols. All data were normalized to urinary creatinine concentrations, measured by an Olympus Au400 autoanalyser.

**Glomerular morphology and counts**

Glomerular morphology was assessed to the following categories: a, normal; b, only one or two capillary loops; c, no capillary loops; d, abnormal. Glomeruli from E18.5 kidney sections from wild-type, \textit{Vangl2}2+/− and \textit{Vangl2}Lp/Lp mice were photographed and assigned to each category by an assessor unaware of the genotypes. At least 100 glomeruli in each of the three kidneys from each of the three genotypes were assessed. In individual kidneys, care was taken to use sections at least 100 µm from each other to avoid counting the same glomeruli more than once. For each kidney, the proportion of glomeruli in each category was then calculated. Results are expressed as a percentage of total glomeruli in each genotype. The same images were used to calculate glomerular tuft diameters (which was taken as the maximum diameter of each tuft). To measure the number of glomeruli in each kidney, organs were removed from 4-week-old heterozygote \textit{Vangl2}2+/− and wild-type male mice and placed in 0.1 M hydrochloric acid for 30 min at 37°C. The acid was removed and replaced with PBS and the tissue homogenized. Twenty imicrolitres of this was pipetted into a haemacytometer and using a 10× objective lens, the number of glomeruli in the aliquot counted. This was repeated in triplicate for each sample and the three results averaged to determine the number of glomeruli in each sample and therefore each kidney (65).
Podocyte cell culture

Mouse podocytes (gift from Peter Mundel, University of Miami, USA) (37) were cultured on tissue culture plastic coated with 1% Matrigel substrate. For ongoing proliferation, cells were cultured at 33°C in a 5% CO₂ incubator in RPMI (Invitrogen) supplemented with 10% FCS (Invitrogen), antibiotics and interferon-γ (10 U/ml). To induce differentiation, cells were thermoshifted to 37°C and cultured in RPMI in the absence of interferon-γ. Cells were allowed to differentiate under these conditions for 7 days when they formed extensive processes similar to their in vivo phenotype.

Quantitative RT–PCR

One mg of RNA was isolated from proliferating and differentiating podocytes using TRI Reagent and cDNA prepared for PCR of the PCP genes Celsr1, Fat4, Mpk1, Mpk2, Vangl1 and Vangl2 using previously described methods (66). To assess differences in gene expression, quantitative real-time RT–PCR was performed (66) on samples from three independent experiments. All measurements were performed in duplicate and HPRT was used as a housekeeping gene to allow quantification. Alterations in gene expression in differentiated cells which was given a standardized value of 1. Negative controls of reactions without cDNA template were included. Primer details are available on request.

Statistical methods

Data were analysed using either unpaired two-tailed t-tests, or when more than two groups were analysed by one-way ANOVA with least square difference correction. Significance was accepted at P < 0.05; error bars in all data represent standard error mean.

SUPPLEMENTARY MATERIAL

Supplementary Material is available at HMG online.

ACKNOWLEDGEMENTS

We thank Caroline Formstone for providing Celsr1 antibody, Peter Mundel for the mouse podocyte cell line and Lauren Chessum for technical assistance.

Conflict of Interest statement. None declared.

FUNDING

This work was supported by the Medical Research Council, the NIHR Manchester Biomedical Research Centre, St Peter's Trust for Kidney Bladder & Prostate Research, the Wellcome Trust, a Walport/NIHR clinical lecturership to J.P. and a Kidney Research UK Non-Clinical Senior Fellowship to D.A.L. Funding to pay the Open Access Charge was provided by Medical Research Council.

REFERENCES


