Planar cell polarity signaling in the uterus directs appropriate positioning of the crypt for embryo implantation

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Blastocyst implantation is a complex process requiring coordination of a dynamic sequence of embryo-uterine interactions. Blood vessels enter the uterus from the mesometrium, demarcating the uterus into mesometrial (M) and antimesometrial (AM) domains. Implantation occurs along the uterine longitudinal axis within specialized implantation chambers (crypts) that originate within the evaginations directed from the primary lumen toward the AM domain. The morphological orientation of crypts in rodent uteri was recognized more than a century ago, but the mechanism remained unknown. Here we provide evidence that planar cell polarity (PCP) signaling orchestrates directed epithelial evaginations to form crypts for implantation in mice. Uterine deletion of Vang-like protein 2, but not Vang-like protein 1, conferred aberrant PCP signaling, misdirected epithelial evaginations, defective crypt formation, and blastocyst attachment, leading to severely compromised pregnancy outcomes. The study reveals a previously unrecognized role for PCP in executing spatial cues for crypt formation and implantation. Because PCP is an evolutionarily conserved phenomenon, our study is likely to inspire implantation studies of this signaling pathway in humans and other species.

PCP | Vangl2 | embryo implantation | uterus | crypt

E ffective reciprocal cross-talk between the receptive uterus and a competent blastocyst is critical for successful implantation (1). In nearly all mammals studied, the uterus differentiates into a receptive state timed appropriately with the migration of blastocysts into the uterus to initiate bidirectional communication for implantation. The receptive state is transient; in mice the uterus becomes receptive on day 4 of pregnancy (day 1 =vaginal plug) and refractory to implantation by the afternoon of day 5 (2). The coordinated actions of progesterone and estrogen regulate the proliferation and/or differentiation of major uterine cell types in a spatiotemporal manner to establish this brief window of receptivity for implantation. These changes involve an interplay of ovarian hormones, transcription factors, growth factors, morphogens, cytokines, and other signaling molecules. Aberrations or defects in any of these pathways may result in implantation failure or defective implantation, which propagates adverse ripple effects through the remaining course of gestation, compromising pregnancy outcomes (2-5).

Blood vessels enter the uterus from the mesometrium and delineate the uterus into mesometrial (M) and antimesometrial (AM) domains. In mice, blastocyst attachment occurs within specialized crypts (implantation chambers), which originate as epithelial evaginations from the main lumen at orderly spaced intervals at the AM domain. The formation of these directed evaginations is intiated on the morning of day 3, and they elongate by the afternoon of day 4 for blastocyst attachment and subsequent crypt formation in the evening along the Wnt5a gradient (6). Blastocyst attachment coincides with increased endometrial vascular permeability at the site of the attachment, which can be monitored by i.v. injection of a blue dye; distinct blue bands indicate sites of blastocyst attachment in the uterus (3, 7). On day 5, the blue bands are distinct, and the underlying stroma surrounding the implantation chamber begins to differentiate into decidual cells (decidualization) to create a favorable environment to support embryonic growth (2).

The unique architectural organization of crypts for blastocyst homing and attachment was first recognized in rodents more than a century ago and later by other investigators (8, 9). However, the mechanism by which epithelial evaginations are appropriately directed to form crypts at the AM domain for embryo homing and implantation remained unknown.

The regular spacing of luminal epithelial (LE) evaginations in preparation for implantation is reminiscent of the spatial patterning and directed morphogenetic movements resulting from planar cell polarity (PCP) signaling during embryogenesis (10, 11). PCP is essential for establishing spatial cues during organogenesis in multicellular tissues, directing actin-dependent morphogenetic cell movements to polarize structures in a wide range of settings,

Significance

Blastocyst implantation is a complex process that coordinates reciprocal embryo–uterine interactions. The uterus is demarcated by mesometrial (A)–antimesometrial (AM) domains; implantation occurs at regularly spaced intervals along the longitudinal axis of the AM pole, within specialized implantation chambers (crypts). The organized crypt formation in the rodent uterus was first observed in 1901, but the mechanism driving this phenomenon remained unknown. We found that planar cell polarity (PCP) signaling coordinates luminal epithelial evaginations toward the AM domain to form crypts for embryo homing and implantation in mice. Disruption of PCP signaling by inactivation of Vang-like protein 2, but not Vang-like protein 1, in the uterus disturbs the regular formation of epithelial evaginations and crypts, disrupting implantation and decidualization and severely compromising pregnancy outcomes.

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including *Drosophila* wing hair (12) and mammalian neural development (13). Disturbances in PCP signaling produce developmental anomalies in neural tube closure, left–right asymmetry, and hair cell orientation in the inner ear (14–20). In humans, mutations or polymorphisms of PCP components are associated with an array of developmental defects including spina bifida, cardiac outflow malformations, and cystic renal disease (21–24).

Vertebrate core PCP components are highly conserved and include noncanonical Wnt family members, their Frizzled receptors, coreceptors (Ror1/Ror2), and membrane-bound signaling intermediaries (Celsr1, Vangl, Scribble, Fat/Dachous) as well as intracellular protein mediators (Dvl, Prickle). These mediators are intracellularly sorted and reside asymmetrically along the apicobasal axis of the cell, initiating spatial organization through cellcell contacts (19).

Here we show that PCP components display unique uterine expression before and during embryo implantation and, moreover, that mice with uterine-specific deletion of Vangl2 (Vangl2^{d/d} mice) show abnormal uterine PCP activity with changes in the spatial distribution of other PCP components. These changes lead to aberrant LE evaginations with random distribution of glands within both the M and AM domains, resulting in severe subfertility. Interestingly, conditional deletion of both Vangl1 and Vangl2 (Vangl1^{d/d} Vangl2^{d/d}) in the uterus mostly recapitulated the infertility phenotype seen in Vangl2^{d/d} females, suggesting that Vangl2 is the predominant isoform in this role; Vangl1^{d/d} females showed normal fertility. We also found that disrupted PCP signaling is associated with derailed Noggin-Bmp2 interaction. Finally, we observed that PCP signaling pathways converge on Dvl2 signaling, and aberrant PCP signaling conferred defective decidualization in the underlying stroma surrounding the implantation chamber, suggesting PCP's role in the initiation of this process. Taken together, these findings provide clear in vivo and in vitro evidence that PCP signaling is critical for the formation of the directed epithelial evaginations leading to the formation of the crypts required for optimal blastocyst homing, implantation, and decidualization in the mouse uterus.

Results

Uterine Vangl2 Deletion Causes Defects in Epithelial Evaginations and Crypt Formation and Severely Compromises Pregnancy Outcome. Vangl2, a core PCP component, works in collaboration with Vangl1, Celsr1, Scribble (Scrib), Dvl2, Wnt5a, and ROR to execute PCP signaling (16, 25–27). Immunofluorescence (IF) staining using a highly specific Vangl2 antibody (28) reveals that Vangl2 is distinctly localized in luminal and glandular epithelial cells on days 4 and 5 (the days of uterine receptivity and implantation, respectively) and at relatively lower levels in surrounding endothelial cells and stromal cells adjacent to the epithelium (Fig. 1A and Fig. S1). We chose to use conditional uterine deletion of Vangl2, because looptail mice homozygous for Vangl2 mutation have structural defects in the uterus, with imbibition of fluid into the lumen and defects in the vaginal opening, precluding studies of PCP on implantation (29). In contrast, mice with conditional uterine deletion of Vangl2 (Vangl2^{d/d}) by Cre expression under the control of progesterone receptor (Pgr-Cre) have normal uterine structures and morphology; uterine deletion of Vangl2 was highly efficient (Fig. S1). However, Vangl2^{d/d} females show severe subfertility: Approximately 46% of plugpositive females fail to produce any litters after multiple mating, and the remaining 54% of plug-positive *Vangl2*^{d/d} females produce significantly smaller litters (Fig. 1B). These results show that Vangl2 is critical for normal fertility. A careful examination of fertility phenotypes showed that these females exhibited pregnancy stagespecific effects. Vangl2^{d/d} females had numerous unevenly spaced resorption sites on days 12 and 14 of pregnancy (Fig. 1C and Fig. S2 A and B) preceded by retarded placental development with a predominance of trophoblast giant cells (TGC) and reduced expression of a TGC marker gene Mash2 (Fig. S2C).

On day 8 of pregnancy, *Vangl2*^{d/d} females showed an atypical morphology of the implantation sites: In comparison with the

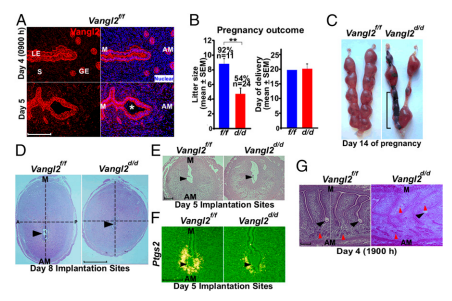


Fig. 1. Uterine inactivation of Vangl2 results in compromised pregnancy outcome. (A) IF localization of Vangl2 in uteri on days 4 and 5 of pregnancy. GE, glandular epithelium; LE, luminal epithelium; S, stroma; *, embryo. (Scale bar, 100 μm.) (B) Litter sizes and day of delivery in littermate $Vangl2^{bf}$ and $Vangl2^{bf}$ females. Percentages and numbers above the bars indicate the number of litters born compared with the total number (n) of plug-positive females in each genotype. Data are shown as mean ± SEM; **P < 0.01. (C) Day 14 implantation sites in $Vangl2^{bf}$ and $Vangl2^{bf}$ demales. The bracket indicates adjacent resorption sites. (D) Histology of day 8 implantation sites in $Vangl2^{bf}$ and $Vangl2^{bf}$ dimice. Note the elliptical vs. round shape of $Vangl2^{bf}$ and $Vangl2^{bf}$ and $Vangl2^{bf}$ and $Vangl2^{bf}$ day 5 implantation sites in $Vangl2^{bf}$ and $Vangl2^{bf}$ and $Vangl2^{bf}$ dimplantation sites in $Vangl2^{bf}$ and $Vangl2^{bf}$ an

normally elliptical implantation sites of $Vangl2^{f/f}$ females, which align with the embryonic axis, $Vangl2^{d/d}$ implantation sites were more spherical. The ratio of length along the M-AM axis respective to the anterior-posterior axis was 24% less in Vangl2^{d/d} females (Fig. 1D), indicating abnormal implantation chamber development and pregnancy progression in these mice. These morphological changes suggested that adverse pregnancy effects in Vangl2^{d/d} mice originated earlier. When examining the status of implantation sites on day 5 by blue dye injection (7), we often found very weak blue bands, indicating inferior blastocyst attachment in $Vangl2^{d/d}$ uteri (Fig. 1E and Fig. S2D). This finding was reinforced by the observation of aberrant cell-specific expression of Ptgs2 (Cox2), a marker for successful implantation, on day 5 (Fig. 1F) (30).

To explore further LE evagination before blastocyst attachment on the evening of day 4, we examined the histology of uterine longitudinal sections. Normally, epithelial evaginations for crypt formation occur at regular intervals aligned with glands and are invariably directed toward the AM domain (Fig. 1G). We found that in Vangl2^{d/d} females evaginations lack direction and are situated at both the M and AM domains, resulting in dysfunctional formation of crypts and defective attachment of the embryos positioned within these crypts on the evening of day 4 (Fig. 1G). Interestingly, this phenotype of severe subfertility in $Vangl2^{d/d}$ females was not caused by altered uterine receptivity, as evaluated by the expression of *Ihh*

and Msx1, markers of uterine receptivity, on day 4 (Fig. S2E) (3) or by changes in epithelial cell proliferation (Fig. S2F). In addition, ovarian hormone levels, ovarian Vangl2 expression, and uterine expression of progesterone receptor (PR) and estrogen receptor (ER) showed no apparent changes in Vangl2^{f/f} or Vangl2^{d/d} mice (Fig. S3). Taken together, these results show a previously unrecognized role of uterine Vangl2 in the orientation of epithelial evaginations and crypt formation, because loss of this signaling in Vangl2^{d/d} mice led to aberrant implantation and adverse pregnancy outcome.

PCP Activity Directs LE Evaginations to Form Crypts at the AM Domain.

Vangl2 executes PCP signaling with Celsr1, Scrib, and Vangl1 as well as with intracellular protein mediators such as Dvl (28, 31-33). We focused on these PCP components because Vangl2 is known to interact directly with Scrib and Celsr1; we also confirmed the direct interaction of Scrib with Vangl2 (Fig. S44). Spatiotemporal localization of PCP components shows that Vangl2 and Scrib are first substantially expressed in the LE with localization to the apical surface on day 3 of pregnancy (Fig. S4B) and are further amplified on day 4 before implantation (Fig. 1A), suggesting progesterone regulation of Vangl2 expression. This expression is similar to Wnt5a expression (6). To examine the spatial distribution of PCP proteins as a functional output of PCP

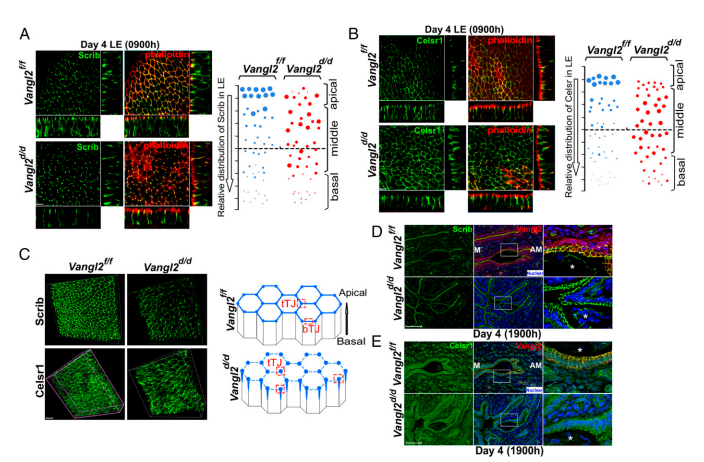


Fig. 2. Aberrant distribution of PCP proteins in Vangl2^{d/d} luminal epithelia. (A, Left) 3D reconstruction of Scrib localization by IF in ex vivo explants of Vangl2^{t/f} and Vangl2^{d/d} uterine LE on day 4 showing that colocalization of Scrib and phalloidin was lost in Vangl2^{d/d} LE cells. (Scale bar, 10 μm.) (Right) Relative apicobasal distribution of Scrib in Vangl2^{f/f} and Vangl2^{d/d} LE. (B, Left) 3D reconstruction of Celsr1 localization on the morning of day 4. (Scale bar, 10 μm.) (Right) Relative apicobasal distribution of Celsr1 in Vangl2^{(ff} and Vangl2^{(dd} LE. (C, Left) Merged 3D images of IF of Scrib and Celsr1 at the apical surface of $Vangl2^{fif}$ and $Vangl2^{d/d}$ LE cells on the morning of day 4. (Right) Changes in Scrib and Celsr1 accumulation at the apical surface of $Vangl2^{fif}$ Vang/2^{d/d} LE cells. Note that in floxed LE, Scrib and Celsr1 localize to bicellular (bTJ) and tricellular (tTJ) tight junctions, but they are assembled primarily in tricellular junctions in $Vang|2^{d/d}$ LE. (Scale bar, 10 μ m.) (D) IF of Scrib in the longitudinal uterine sections of $Vang|2^{d/d}$ and $Vang|2^{d/d}$ mice on the evening of day 4. (Scale bar, 100 μ m.) (E) IF of Celsr1 in the longitudinal uterine sections of $Vang|2^{d/f}$ and $Vang|2^{d/d}$ mice on the evening of day 4. (Scale bar, 100 μ m.) Asterisks in D and E indicate the location of embryos.

activity, we performed 3D reconstruction of IF images of Vangl2, Scrib, and Celsr1 in isolated, intact LE. The results show that Scrib and Celsr1 are colocalized with Vangl2 at the apical surface in the epithelium on the morning of day 4 when LE evagination is in progress (Fig. 2 *A* and *B*). In contrast, Scrib and Ceslr1 are not appropriately localized in *Vangl2*^{d/d} LE (Fig. 2 *A* and *B* and Movies S1–S4). We also found that these PCP components accumulate less at bicellular junctions and more at tricellular junctions in *Vangl2*^{d/d} LE, with diffuse distribution of phalloidin staining (Fig. 2*C* and Movies S5–S8). These results prompted us to examine the status of Scrib and Celsr1 in the uterus on the evening of day 4 before blastocyst attachment. We found that the localization of these PCP components is strikingly altered in irregularly oriented evaginations and crypts in *Vangl2*^{d/d} mice (Fig. 2 *D* and *E*).

Celsr1 is clearly displayed in the crypt epithelial cells of $Vangl2^{t/f}$ females on day 5, but localization is patchy in $Vangl2^{d/d}$ mice (Fig. 3A). However, Scrib becomes undetectable in the $Vangl2^{t/f}$ crypt epithelium, appearing instead in the basal lamina and underlying stroma, but is retained in the epithelium in $Vangl2^{d/d}$ mice (Fig. 3B). In contrast, the abnormal spatial distribution of Scrib is associated with increased protein levels and phosphorylation, as shown by gel shift assays (Fig. 3C). There is evidence that Scrib phosphorylation in transfected cells interferes with its apical localization (34). Collectively, PCP activity is consistently aberrant in $Vangl2^{d/d}$ uteri around the time of implantation, manifesting poorly coordinated epithelial evaginations and aberrant crypt formation and implantation. These results suggest that disorganization of apical Scrib and Ceslr1 in the absence of Vangl2 compromises the apical actin-myosin cytoskeleton and directional

cues between cells required for the execution of proper LE evaginations toward the AM domain.

We speculated that LE tight junctions were also altered in these mice, and thus we assessed the distribution of the tight junction protein ZO-1. Although it is expressed uniformly in the $Vangl2^{f/f}$ LE on day 4, its distribution is heterogeneous with variable intensity and distribution in $Vangl2^{d/d}$ LE (Fig. 3D). In the same context, the tight junction protein Claudin 1 is expressed at the bottom region of the crypt in $Vangl2^{f/f}$ uteri but is expressed diffusely throughout the crypt epithelium around the blastocyst in $Vangl2^{d/d}$ mice on day 5 (Fig. 3E). These changes, along with changes in PCP components, are reflected in inefficient removal of the LE barrier surrounding the embryo in $Vangl2^{d/d}$ mice on day 6, as indicated by the mislocalization of E-cadherin and β -catenin (Fig. 3F).

Vangl2 Is the Principal Isoform for Executing PCP Signaling in the Uterus. We speculated that Vangl1, a Vangl family member that interacts via the same PDZ domain as Vangl2, compensates for the loss of Vangl2 to retain partial fertility in some Vangl2^{d/d} females. This assumption is supported by the observation of elevated levels of Vangl1 by quantitative RT-PCR (qRT-PCR) in Vangl2^{d/d} uteri on day 4 of pregnancy (Fig. 4A). Because we could not identify any authentic commercial Vangl1 antibody, we used RNAscope in situ hybridization to localize cell-specific expression of Vangl1 transcripts and found that the distribution of Vangl1 expression in the uterus is similar to that of Vangl2 (Fig. 4B). To provide genetic evidence, we generated mice with uterine-specific deletion of Vangl1 (Vangl1^{d/d}), and determined that deletion was proficient (Fig. 4C). We then examined the fertility phenotypes

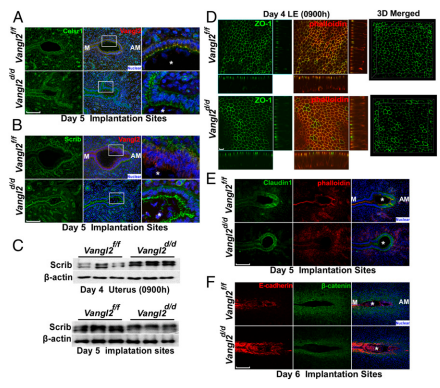


Fig. 3. Interactions among PCP components in the absence of Vangl2. (A) Celsr1 colocalizes with Vangl2 at the apical surface in $Vangl2^{fif}$ implantation sites. In $Vangl2^{did}$ implantation sites Celsr1 localization was aberrant. (Scale bar, 100 μm.) (B) IF localization of Scrib and Vangl2 in day 5 implantation sites. Although Scrib was displayed at the apical side of the crypt epithelium in the $Vangl2^{did}$ mice, it was absent in $Vangl2^{fif}$ females. (Scale bar, 100 μm.) (C) Western blotting for Scrib in day 4 uteri and day 5 implantation sites of $Vangl2^{did}$ mice showed higher levels of Scrib and its phosphorylation in $Vangl2^{did}$ uteri. n = 3 females per genotype. (D, Left) The tight junction marker ZO-1 lost its uniform distribution at the apical surface of $Vangl2^{did}$ LE. (Right) 3D reconstruction of all layers. (Scale bar, 10 μm.) (F) IF of Claudin1 and phalloidin in day 5 implantation sites of $Vangl2^{did}$ mice. (Scale bar, 100 μm.) (F) Poorly developed primary decidual zone with epithelial retention of E-cadherin and limited β-catern with inefficient invasion of the trophoblast through the crypt epithelium in $Vangl2^{did}$ mice. (Scale bar, 200 μm.) Asterisks in A, B, E, and F indicate the location of embryos.

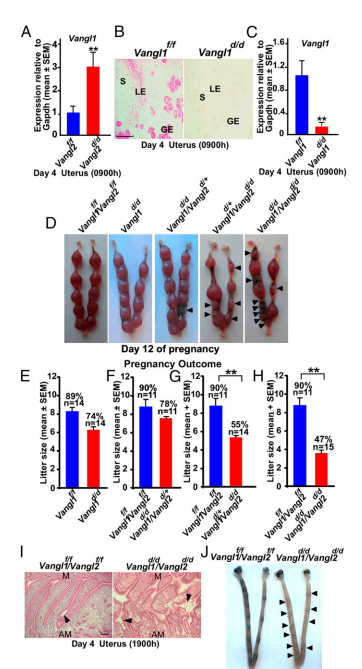


Fig. 4. Fertility phenotypes in Vangl1^{d/d} mice and Vangl1^{d/d}Vangl2^{d/d} mice (A) Results of Vangl1 qPCR in day 4 uteri of Vangl2^{d/d} mice. n = 3; data are shown as mean \pm SEM; **P < 0.01. (B) In situ hybridization of Vangl1 in Vangl1^{ttt} and Vangl1^{d/d} uteri on the morning of day 4 showing specific localization of Vangl1 transcripts. Red granular speckles indicate transcripts. (Scale bar, 200 µm.) (C) qPCR results of Vangl1 in day 4 uteri of Vangl1^{flf} and Vangl1^{dld} mice. Data are shown as mean \pm SEM; **P < 0.01. (D) Implantation sites in Vangl1^{f/f}Vangl2^{f/f} Vangl1^{d/d}, Vangl1^{d/d}Vangl2^{d/+}, Vangl1^{d/+}Vangl2^{d/d}, and Vangl1^{d/d}Vangl2^{d/d} mice on day 12. Arrowheads indicate degenerating embryos. (E) Litter sizes in littermate Vanql1^{f/f} and Vanql1^{d/d} females. Percentages and numbers above the bars indicate the number of litters born compared with the total number of plugpositive females (n) in each genotype. (F-H) Litter sizes in Vangl1^{f/f} Vangl2^{f/f} and Vangl1^{d/d}Vangl2^{d/+}females (F), Vangl1^{f/f}Vangl2^{f/f} females and Vangl1^{d/+}Vangl2^{d/c} females (G), and Vangl1ft Vangl2ftf and Vangl1dd females (H). (I) Histology of longitudinal sections of day 4 uteri from Vangl1^{f/f} Vangl2^{f/f} females and Vangl1^{d/d}Vangl2^{d/d} females. Arrowheads indicate the location of embryos. (Scale bar, 500 μm.) (J) Implantation sites in Vangl1^{f/f}Vangl2^{f/f} and Vangl1^{d/d}Vangl2^{d/d} mice on day 5. Arrowheads indicate very weak bands in Vangl1^{d/d}Vangl2^{d/d} mice.

in different crossings of mice with uterine deletions of Vangl1 and/or Vangl2. The fertility phenotypes are normal in $Vangl1^{d/d}$ or $Vangl1^{d/d}/Vangl2^{d/+}$ females (Fig. 4 D–F), whereas the fertility phenotype in $Vangl1^{d/+}/Vangl2^{d/d}$ females closely resembles that of $Vangl2^{d/d}$ mice (Fig. 4 D, G, and H). However, the infertility rate is somewhat higher in $Vangl1^{d/d}/Vangl2^{d/d}$ females; 53% of plugpositive females were infertile, and the remaining 47% of plugpositive females generated smaller litters than $Vangl2^{d/d}$ females (Fig. 4H). Poor morphological coordination of LE evaginations for crypt formation is also apparent in day 4 uteri of $Vangl1^{d/d}/Vangl2^{d/d}$ females (Fig. 4I), with weak blue bands (implantation) on day 5 (Fig. 4I). This result is different from studies in other systems in which deletion of both isoforms led to more severe phenotypes than the deletion of Vangl2 alone (27). The results suggest that Vangl2 is the predominant isoform critical for epithelial evaginations and crypt formation for implantation.

PCP Signaling Is Disrupted Under Aberrant Uterine Wnt5a-ROR Signaling. The failure of organized epithelial evaginations and crypt formation in Vangl2^{d/d} females is reminiscent of the subfertility phenotypes observed in females with aberrant uterine Wnt5a-ROR signaling (6). Thus, we sought to examine the localization and activity of the major PCP components under altered Wnt5a-ROR signaling. To do so, we generated mice with uterine overexpression of Wnt5a (Wnt5a^{GOF}), uterine inactivation of Wnt5a (Wnt5a^d/dd), and double deletion of Ror1 and Ror2 (Ror1^{d/d}/Ror2^{d/d}) using the Pgr-Cre driver (6). Single deletion of Ror1 or Ror2 showed no observable reproductive phenotypes (6). We found that the intensity of Vangl2 signaling correlates with the degree of Wnt5a-ROR signaling: In Wnt5a^{GOF} LE, Vangl2 localization is distinct in both the apical and basal sides of the LE, but the signal intensity is lower in *Wnt5a*^{d/d} and *Ror1*^{d/d}/*Ror2*^{d/d} females on the morning of day 4 (Fig. 5A). Similarly, signal intensities of Celsr1 and Scrib are aberrantly low in the crypt epithelia of Wnt5a^{d/d} and Ror1^{d/d}/Ror2^{d/d} uteri, but the intensity is appreciably higher in Wnt5a^{GOF} crypt epithelium on the evening of day 4 and the morning of day 5 (Figs. S5 and S6). This low signaling intensity was preceded by aberrant apical distribution of Vangl2, Scrib, and ZO-1 in the LE of Wnt5a^{d/d}, Wnt5a^{GOF}, and Ror1^{d/d}/Ror2^{d/d} uteri on the morning of day 4 (Fig. 5 B and C). These findings show defective PCP activity in uteri with inappropriate Wnt5a-ROR signaling, providing evidence that Wnt5a-ROR signaling intersects PCP signaling in implantation.

Aberrant PCP Signaling Derails Noggin-Bmp2 Interaction. The complex interactions between the embryo and uterus during implantation show common features of epithelial-mesenchymal interactions underlying developmental processes including limb, tooth, hair follicle, lung, and kidney development. Bone morphogenetic proteins (BMP) and their antagonists play a critical role in these processes (35–37). Noggin (Nog) is an antagonist to Bmp2 (38). We hypothesized that PCP signaling influences Nog-Bmp2 interaction to execute blastocyst attachment within the crypt. Using in situ hybridization, we examined the expression of Bmp2 and Nog. In $Vangl2^{f/f}$ females Nog is expressed in the subepithelial stroma, but its expression is rapidly down-regulated, followed by Bmp2 expression in the stroma at the site of embryo implantation before and after blastocyst attachment on the evening of day 4 and the morning of day 5. This pattern is disrupted in $Vangl2^{d/d}$ females during the same time periods (Fig. 6 A and B). In pseudopregnant $Vangl2^{f/f}$ or $Vangl2^{d/d}$ females, generated

In pseudopregnant *Vangl2*^{1/1} or *Vangl2*^{2/1} females, generated by mating with vasectomized males, *Nog* expression is present in the subepithelial stroma along the entire length of the epithelial evagination, with no sign of *Bmp2* expression at the expected time of implantation. However, *Nog* expression is more robust in deleted uteri with aberrant epithelial evaginations (Fig. 6C). These results suggest that PCP signaling in the uterus is essential for implantation-competent blastocysts to confer down-regulation of *Nog* with induction of *Bmp2* to initiate the attachment reaction and subsequent decidualization at the implantation site.

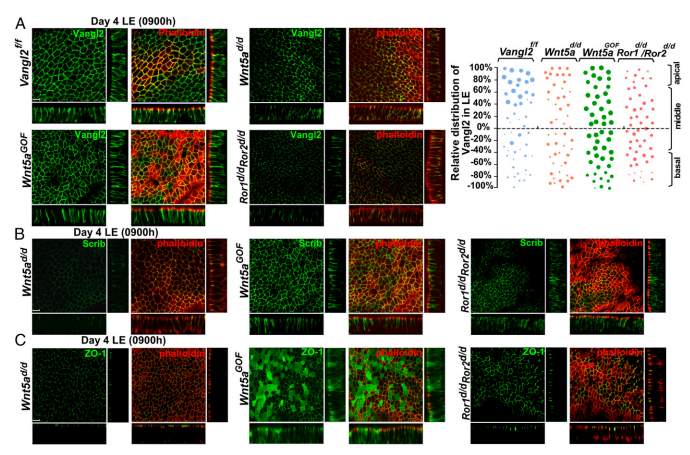


Fig. 5. Aberrant Wnt5a–ROR signaling dysregulates the localization of PCP components and tight junctions in LE. (*A, Left*) 3D reconstruction of Vangl2 IF showing aberrant expression patterns in *Wnt5a*^{d/d}, *Wnt5a*^{GOF}, and *Ror1*^{d/d}/*Ror2*^{d/d} LE cells on the morning of day 4. (*Right*) Relative apicobasal distribution of Vangl2 in *Vangl2*^{f/f} mice which was altered in *Wnt5a*^{d/d}, *Wnt5a*^{GOF}, and *Ror1*^{d/d}/*Ror2*^{d/d} mice. (*B*) Comparison of aberrant apicobasal expression of Scrib in LE of *Wnt5a*^{d/d}, *Wnt5a*^{GOF}, and *Ror1*^{d/d}/*Ror2*^{d/d} mice. (*C*) Aberrant apicobasal expression of ZO-1 in LE of *Wnt5a*^{d/d}, *Wnt5a*^{GOF}, and *Ror1*^{d/d}/*Ror2*^{d/d} mice. (Scale bars, 10 μm.)

Bmp2 signaling was also shown to be critical for the initiation and progression of decidualization (38, 39). BMPs execute their effects by phosphorylating Smad proteins. We noted dampened BMP signaling, as indicated by reduced intensity of IF localization of pSmad1/5/8 in the primary decidual zone (pdz) on day 5 in *Vangl2*^{d/d} females (Fig. 6D).

Dvl2 Signaling Is Dysregulated Under Abnormal PCP Signaling. Because PCP signaling can converge on the Dvl pathway, we explored the status of Dvl2 in the uterus under aberrant Wnt5a-ROR-Vangl2 signaling. Dvl2 is a scaffolding protein that interacts directly with Vangl2 and plays an important role downstream of PCP signaling (31). Western blotting results show that the levels of phosphorylated Dvl2 (pDvl2, active) are lower in Vangl2^{d/d} uteri on the evening of day 4 with no apparent changes at the implantation site on day 5 (Fig. 7 A and B), although Dvl2 localization patterns in WT and $Vangl2^{d/d}$ uteri show little change on the evening of day 4 (Fig. S7A). We then asked if disturbances in Wnt5a–ROR signaling alter the Dvl2 phosphorylation status at the implantation sites. Indeed, levels of pDvl2 are very low to undetectable in Wnt5a^{d/d} and Ror1^{d/d} Ror2^{d/d} uteri compared with those in WT uteri on the evening of day 4 and in day 5 implantation sites (Fig. 7 C and E–G). In contrast, $Wnt5a^{\rm GOF}$ mice show increased uterine pDvl2 levels on these days (Fig. 7 C and D). Phosphorylation was confirmed by treatment with calf intestinal alkaline phosphatase (CIAP) that abolished the phosphorylated band (Fig. 7H). Collectively, these results suggest that Wnt5a-ROR-PCP signaling converges on the downstream mediator Dvl2 in the uterus at the time of

implantation. Studies in different systems showed that Wnt5a-ROR and Vangl2 signaling converge on Dvl2 activity for cell shape and cell-directed movements (10).

Aberrant PCP Signaling Resulted in Defective Decidualization. In mice, differentiating stromal cells surrounding the blastocyst initially form the pdz on day 5. This differentiation involves the transformation of stromal cells into epithelioid cells, which are densely packed to form an avascular zone around the implantation chamber. By day 6, the pdz is well formed, and a secondary decidual zone is formed at the periphery of the pdz. After day 8, placental and embryonic growth gradually reduce the decidual thickness. One proposed function of the pdz is to form a selective epithelioid barrier surrounding the embryo, safeguarding the embryo against harmful agents that may infiltrate from maternal circulation.

Because aberrant PCP activity interferes with BMP signaling, and because Bmp2 can influence Dvl2 distribution (40), we asked if Dvl2 activity is also associated with the initiation of decidualization (pdz formation). We examined Dvl2 localization by IF and found that Dvl2 accumulates in puncta within decidualizing stromal cells in the nascent pdz. Puncta density and Dvl2 IF intensity are markedly up-regulated in *Wnt5a* GOF implantation sites on day 5, whereas *Wnt5a* or *Ror1/Ror2* pdz sites have significantly fewer puncta than those in WT mice (Fig. 7I). To explore the relationship between Dvl2 and Vangl2 in decidualization, IF of Dvl2 in stromal cells differentiating to decidual cells was examined. We found that Dvl2 is localized to distinct puncta in stromal cells differentiating to pdz at the implantation sites on day 5 in

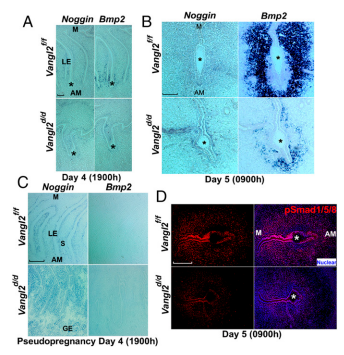


Fig. 6. Uterine deletion of Vangl2 derails Noggin and Bmp2 expression. (A) In situ hybridization of Nog and Bmp2 in $Vangl2^{dif}$ and $Vangl2^{did}$ uteri on the evening of day 4 showing their aberrant expression in $Vangl2^{did}$ uteri. (Scale bar, 200 μ m.) (B) In situ hybridization of Nog and Bmp2 in $Vangl2^{dif}$ and $Vangl2^{dif}$ implantation sites on day 5 showing the disappearance of Nog with robust Bmp2 expression in $Vangl2^{dif}$ females; this pattern was disrupted in $Vangl2^{did}$ implantation sites. (Scale bar, 100 μ m.) (C) In situ hybridization of Nog and $Vangl2^{did}$ implantation sites. (Scale bar, 100 μ m.) (C) In situ hybridization of $Vangl2^{did}$ implantation sites. (Scale bar, 100 μ m.) (C) In situ hybridization of $Vangl2^{did}$ in the subepithelial stroma along the entire length of the LE evagination in $Vangl2^{dif}$ and $Vangl2^{did}$ uteri, with more robust expression in the latter. $Vangl2^{did}$ uteri, with more robust expression in the latter. $Vangl2^{did}$ uteri, $Vangl2^{did}$ uteri. (Scale bar, 200 $Vangl2^{did}$ uteri, $Vangl2^{did}$ uteri. (Scale bar, 200 $Vangl2^{did}$ uteri.) (E, luminal epithelium; S, stroma. (D) If of pSmad1/5/8 in $Vangl2^{dif}$ and $Vangl2^{did}$ uteri. Asterisks indicate the location of the embryo.

Vangl2^{ff} females but is mostly absent from Vangl2^{dd} implantation sites (Fig. 7J). A greater abundance of puncta in the pdz cells is noted with the formation of the definitive pdz on day 6 in WT mice but not in conditionally deleted mice. These puncta are not evident when an antibody (105B), reported to detect the non-phosphorylated form of Dvl2 (41), is used (Fig. S7 B and C), suggesting that these puncta are positive for phosphorylated Dvl2.

Because puncta were abundant in pdz cells in vivo, we asked whether puncta are formed in stromal cells induced to decidualization in vitro. Primary stromal cells isolated from WT day 4 uteri were induced to decidualize in vitro (42). The decidual cells show higher pDvl2 levels than nondecidualized stromal cells in vitro (Fig. S7D), but they fail to show puncta formation. However, results with 10B5 Dvl2 antibody show lower levels in decidual cells in culture than in nondecidualized stromal cells (Fig. S7E). Although the definitive role of puncta is currently unknown (43–46), their formation in the uterus is clearly associated with the formation of the pdz. It is possible that puncta formation is relevant to the context of tissue patterning in vivo. Overall, these results suggest that PCP signaling is also involved in patterning stromal cells toward an epithelioid cell type during the pdz formation.

Discussion

Uterine receptivity is achieved primarily by the ovarian steroid hormones that provide an environment conducive to the complex molecular and physical interactions between the blastocyst trophectoderm and the uterine epithelium to achieve implantation.

This study reveals a previously unrecognized but essential step in the receptive uterus before the attachment reaction: the morphological, cellular, and molecular changes that contribute to the formation of directed, regularly spaced LE evaginations and the formation of implantation chambers (crypts) at the AM domain. Previous studies had shown that aberrant uterine Wnt5a-ROR signaling caused defective implantation (6), but the mechanism by which Wnt5a-ROR signaling coordinates crypt formation remained unknown. Because Wnt5a forms a signaling gradient along the LE evaginations toward the AM domain (6), we speculated that directed cell movement of the epithelial evaginations through the stromal bed is guided by PCP signaling. Indeed, the present findings show that noncanonical Wnt5a-ROR signaling intersects with PCP signaling to initiate epithelial evaginations from the main LE toward the AM domain. This process is evident from our studies showing that uterine inactivation of Vangl2 (Vangl2^{d/d}) disrupts directed epithelial evaginations. The crucial role of PCP in LE evaginations was reinforced further by aberrant spatial distribution and expression of the PCP components Scrib, Celsr1, and Dvl2 in $Vangl2^{d/d}$ uteri and by the poor, asymmetric sorting of PCP components in LE cells in the absence of Vangl2. Aberrant uterine PCP signaling with misdirected evaginations results in poor fertility, with intrauterine demise of embryos and reduced litter size.

This study uniquely identifies a physiologic role for PCP signaling in female fertility. This pathway has several roles during development, and we show here its requirement in maternal uterine tissues before and during implantation (Fig. S8). Mouse blastocysts do not express *Pgr-Cre* (3), eliminating the role of embryonic PCP in this event. Because this pathway is likely evolutionarily conserved across flies, worms, and mammals, the discovery of its role in the mouse uterus in providing directional cues for epithelial evagination and crypt formation at the AM domain is of paramount importance in implantation. Although the Vangl1 and Vangl2 proteins belong to the same family, and both manifest protein–protein interactions via the PDZ domain, Vangl2 appears to be the dominant isoform in the uterus that influences the pregnancy events, because Vangl2 deficiency shows a severe adverse fertility phenotype, whereas fertility is normal in *Vangl1*-deleted uteri.

A previous study used unauthenticated Vangl1 and Vangl2 antibodies to show cell-specific localization and used intraluminal injections of Vangl1 and Vangl2 antisense oligonucleotides in an unspecified volume of water to show their effects on implantation in mice (47). The authors concluded that uterine Vangl1 and Vangl2 are required for implantation, although both embryos and uterine cells were exposed to antisense oligos. Because the uterine luminal volume is merely 150-200 nL per horn around the time of uterine receptivity and implantation with the closure of LE, intraluminal infusion of solution is likely to distort epithelial morphology, cell polarity, and function (48). Furthermore, the intraluminal infusion exposes both the embryo and uterus to antisense oligonucleotides, making it difficult to delineate the roles of Vangl from the uterus versus Vangl from the embryo. Therefore, methodological and experimental design limitations raise concerns over these results, which are contrary to our findings derived using conditional knockout mouse models and authenticated antibodies with mechanistic information on uterine PCP signaling in pregnancy events.

Despite PCP's numerous functions in developmental processes, such as embryogenesis and organogenesis, its function in implantation was unknown. We now show that PCP signaling is a potential mechanism underlying this process. In most mammals, the uterus houses and supports a growing embryo throughout pregnancy until parturition; embryo implantation is the seed of embryogenesis and is a crucial, beginning step in development. In this context, *Vangl2* transcripts in developing neonatal uteri from conditionally deleted *Wnt7a* mice were reduced showing poor adenogenesis (49).

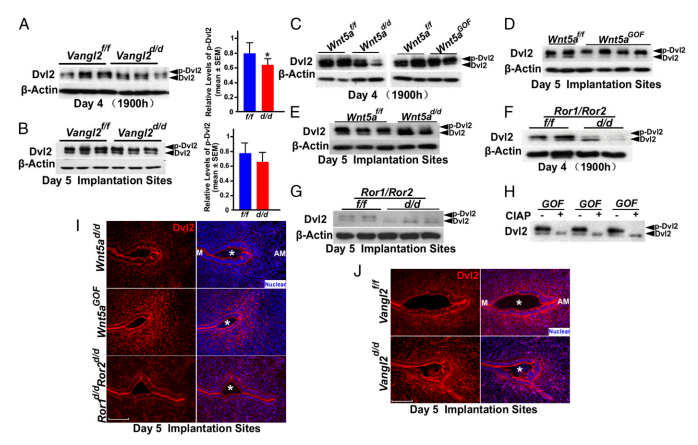


Fig. 7. Wnt5a–ROR–Vangl2 signaling pathways influence Dvl2 levels and activity during implantation. (A and B, Left) Western blotting results of Dvl2 in day 4 uteri (A) and day 5 implantation sites (B) of $Vangl2^{b/d}$ and $Vangl2^{b/d}$ mice. (Right) Relative levels of pDvl2. Data are shown as mean \pm SEM; *P < 0.05. (C) Results of Western blotting of Dvl2 in day 4 uteri of $Wnt5a^{d/d}$ and $Wnt5a^{GOF}$ mice. (D and E) Western blotting for Dvl2 in day 5 implantation sites of $Wnt5a^{GOF}$ (E) mice. (E) mice. (E) and E) Western blotting for Dvl2 in day 5 implantation sites (E) of E0 and E1 western blotting for Dvl2 in day 4 uteri (E1) and day 5 implantation sites (E2) of E3 implantation sites (E3) of E4 uterine samples leads to undetectable levels of phosphorylated Dvl2. (E4) Dvl2 localization in E4 in E5 implantation sites. (E6 calle bar, 100 E7) units in E8 implantation sites. (E8 bar, 100 E9) Asterisks indicate the location of embryos.

Bmp2 signaling is localized primarily to stromal cells surrounding the attaching blastocyst and increases further during decidualization (38). Our observation of reduced expression of pSmad in the stroma in Vangl2^{d/d} uteri on day 5, at the time of implantation and decidualization, suggests that this process is compromised in the uterus deficient in PCP signaling. This observation implies that epithelial-mesenchymal interactions in the context of implantation and decidualization involve PCP-BMP signaling. Previous work showing the role of BMP in embryo spacing (38) suggests that BMP signaling depends on Vangl2 signaling in regular spacing of the LE evaginations and crypt formation for implantation and then decidualization. Our observation that Nog disappears with the induction of Bmp2 in the uterus before and after the initiation of implantation suggests that the Noggin-BMP2 interaction is tightly coupled with blastocyst attachment. The disruption of this interaction in the Vangl2^{d/d} uterus points toward an intimate association of Noggin-BMP interactions with PCP activity. Indeed, there is evidence that Nog expression is tightly coupled with hair follicle development, in which PCP signaling is also a dominant feature (50). The definitive mechanism by which this Nog-Bmp2 engagement is realized remains to be discovered.

Although epithelial-mesenchymal interactions are critical for the execution of many uterine events, it is not yet understood how stromal cells help LE evaginations form implantation chambers and then initiate decidualization following implantation. The distinct architectural arrangement of differentiating stromal cells to epithelioidlike cells with the acquisition of epithelial markers such as ZO-1 surrounding the implanting blastocyst to form the pdz is the hallmark of the decidualization process in rodents (2, 51). Subepithelial stromal cells surrounding the blastocyst express Vangl2, raising the possibility that PCP activity patterns stromal cells to acquire epitheliumlike characteristics at the site of implantation to initiate the pdz formation. The appearance of puncta as shown by pDvl2 localization in the pdz cells in vivo on day 5 is interesting and suggests that pDvl2 has a role in puncta formation that could serve as a marker for transitioning stromal cells to epithelioid cells in vivo. The absence of puncta in cells decidualized in vitro suggests that these decidual cells have lost the characteristics of pdz cells. The role of Dvl2 puncta has yet to be elucidated despite its identification in several systems (46, 52–54).

Our findings have uncovered a previously unidentified but critical role of PCP signaling to direct epithelial evaginations at regular intervals toward the AM domain for crypt formation and successful implantation. This finding is particularly interesting because severe subfertility in mice with aberrant PCP signaling is independent of altered uterine receptivity and steroid hormonal changes, providing evidence that PCP orchestrates LE evaginations to form implantation chambers for blastocyst homing and implantation. This study provides evidence for a physiological role for PCP signaling in the dynamically active adult mouse uterus. Whether this signaling is active in the human endometrium or in mammals other than rodents during implantation is not known. There is indication of undulation of the epithelium at the site of

the embryo implantation in human (Carnegie collections, plate 41) (55). Determining whether this undulation represents epithelial evagination is hampered by the availability of very few samples with limited details on early stages of pregnancy. Furthermore, research to identify very early stages of implantation in humans has been limited by ethical restrictions, and rightfully so. Therefore, the presence of epithelial evagination/undulation before blastocyst attachment is yet to be experimentally documented. Nonetheless, the human endometrium undergoes profound epithelial cell-surface changes with the formation of pinopode (uterodome) and with changes in surface microvilli and cilia during the endometrial receptive phase for implantation (56, 57). It is possible that PCP activity participates in these processes (58, 59). In addition, the formation of epithelial plaque at the site of embryo implantation in humans and, more noticeably, in subhuman primates (macaque) (60) suggests the possible participation of PCP signaling. Thus, the knowledge derived from this study in mice will encourage basic scientists and clinical investigators to probe for PCP signaling in implantation events in humans and other species. This signaling is particularly relevant to investigations of the cause of abnormal implantation in multiple-gestation pregnancies in humans and placenta previa, a source of significant mortality and/or morbidity of the mother and fetus (61, 62). The proteome atlas showing the expression of major PCP components including Vangl2, Vangl1, Scrib, ROR, and Wnt5a in the human endometrium (63) suggests the involvement of PCP signaling in human implantation. Taken together, our research addresses a mechanism behind a long-standing observation in rodent implantation and can spur further studies on PCP signaling in the human uterus and other mammals to improve our understanding of how embryos are directed to select a uterine site for implantation.

Materials and Methods

Mice. $Pgr^{Cre/+}$, $Wnt5a^{d/d}$, $Wnt5a^{GOF}$, and $Ror1^{d/d}Ror2^{d/d}$ mouse lines were generated as previously described (6, 11, 64, 65). $Vangl1^{d/d}$, $Vangl2^{d/d}$, and $Vangl1^{d/d}/Vangl2^{d/d}$ mice were generated by mating floxed females with $Pgr^{Cre/+}$ males (3). Vangl2 floxed mice were generated in Y.Y.'s laboratory (NIH), and Vangl1 floxed mice were purchased from Jackson Laboratories. All mice used in this study were housed under a constant 12-h/12-h light/dark cycle in the Cincinnati Children's Animal Care Facility according to NIH and institutional guidelines for the use of laboratory animals. All protocols were approved by the Cincinnati Children's Animal Care and Use Committee. Mice were provided with autoclaved Laboratory Rodent Diet 5010 (Purina) and UV light-sterilized reverse osmosis/deionized constant-circulation water ad libitum.

Analysis of Pregnancy Events. Female mice were mated with littermate floxed males to induce pregnancy. Ovulation, fertilization, preimplantation embryo development, and implantation were assessed as previously described (3). The morning of finding the vaginal plug was considered day 1 of pregnancy. To examine uterine receptivity and implantation, pregnant dams were killed on day 4 (0900–1000 h), and day 5 (0900 h), respectively. Implantation sites were visualized by i.v. injection of a Chicago blue dye solution before mice were killed, and the number of implantation sites demarcated by distinct blue bands was recorded. The number of blue bands on day 5 in control and experimental animals was compared as an index of ovulation, fertilization, and oviductal embryo transport and development. For litter size analysis, pregnant females were monitored from days 17–21 by observing mice daily for parturition and litter size.

Immunohistochemistry and Histology. Immunohistochemistry of ER (sc-542; Santa Cruz Biotechnology) and PR (8757; Cell Signaling Technology) was performed in paraffin-embedded sections using specific antibodies. Tissue sections from control and experimental groups were processed on the same slide. A Histostain-Plus (diaminobenzidine) kit (859243; Thermo Scientific) was used to visualize specific antigens. Paraffin sections (6 μm) were deparaffinized, hydrated, and stained with H&E for light microscopy analysis (3).

IF and Confocal Microscopy. IF for Vangl2 (custom generated in the J.-P.B. laboratory, INSERM), ZO-1 (33-9100; Thermo Scientific), rabbit E-cadherin (3195s; Cell Signaling Technology), goat E-cadherin (sc-31020; Santa Cruz Biotechnology),

β-catenin (sc-1496; Santa Cruz Biotechnology), Claudin 1 (71-7800; Thermo Scientific) and phalloidin (43166A; Thermo Scientific), Scrib (H300; Santa Cruz Biotechnology), Celsr1 (custom generated in the Fuchs laboratory, Rockefeller University, New York), phospho-Smad1/Smad5/Smad8 antibody (generated by Dan Vasiliauskas, Susan Morton, Tom Jessell, and Ed Laufer, Columbia University, New York), Dvl2 (bs-9059R; Bioss Antibodies), Ki67 (RM-9106-S; Thermo Scientific), and 10B5 (sc-8026; Santa Cruz) was performed using secondary antibodies conjugated with Cy-2, Cy-3, or Cy-5 (Jackson ImmunoResearch). Nuclear staining was performed using Hoechst 33342. Tissue sections from control and experimental groups were processed onto the same slide.

Whole-Mount LE Recovery and Immunostaining. Uteri were cut into 2- to 3-mm pieces and were incubated in HBSS containing 120 mg/mL dispase and 125 mg/mL pancreatin. The pieces were incubated at 37 °C for 1 h. The intact LE then was gently squeezed out of the uterus by a small pair of forceps. The isolated LE was fixed in 4% (wt/vol) paraformaldehyde (PFA) for 10 min at room temperature, washed gently three times in PBS, and then subjected to IF as described above. The z axis and x, y axes were set up for IF scanning by a Nikon A1 invert confocal microscope, and Imaris software was used to analyze the images. At least two independent samples were analyzed.

RNA Isolation and qPCR. RT-PCR was performed as described (3), using the following primers: 5'-CCAAACAGCCTTACCAT-3' and 5'-TCCTTAAGCCCATT-GGTCAC-3' for *Vangl2* (product size 196 bp); 5'-TGGAGGGTAATGTCACAGCA-3' and 5'-AAACCGTTCTTTGCCCTTCT-3' for *Vangl1* (product size 200 bp); and 5'-TCCATGACAACTTTGGCATTG-3' and 5'-CAGTCTTCTGGGTGGCAGTGA-3' for mouse *Gapdh* (product size, 72 bp). *Gapdh* served as an internal control.

In Situ Hybridization. PFA-fixed frozen sections from control and experimental groups were processed onto the same slide and hybridized with ³⁵S-labeled cRNA probes (3, 30) or were hybridized with digoxigenin (DIG)-labeled cRNA probes. DIG-labeled probes were generated according to the manufacturer's protocol (Roche). RNAscope in situ hybridization was performed as described (66). Briefly, frozen tissue sections were fixed in 4% PFA/PBS and then were incubated with hydrogen peroxide and Protease IV according to the manufacturer's protocol (Advanced Cell Diagnostics, ACD). The slides then hybridized with the *Vangl1* probe generated by ACD at 40 °C for 2 h. After hybridization, slides were subjected to signal amplification using an HD 2.0 Detection Kit (ΔCD).

Immunoblotting. Antibodies used included Vangl2 (J.-P.B.), and Actin (sc-1615; Santa Cruz Biotechnology). Bands were visualized using an ECL kit (RPN2108; GE Healthcare). Actin served as a loading control. For the gel-shift assay, proteins were separated in 10% Bis-acrylamide gel for 4 h at 70 V (for Vangl2) or 7% Bis-acrylamide gel for 4 h at 70 V (for Dvl2). For phosphatase treatment, $Wnt5a^{\rm GOF}$ implantation sites were homogenized in RIPA buffer as described above, and an aliquot of each sample was treated with calf intestine alkaline phosphatase (CIAP), 1 unit/µg protein, for 40 min at 37 °C. Vangl2 and Dvl2 proteins were examined by immunoblotting.

Primary Stromal Cell Culture. Uteri from day 4 pseudopregnant WT mice were split open longitudinally and cut into small pieces (2–3 mm long). Tissue pieces were incubated with pancreatin and dispase for 1 h on ice, followed by 30 min at room temperature and 10 min at 37 °C. Fragments of LE sheets were liberated by pipetting the digests several times. The remaining tissue fragments were incubated in type II collagenase to free stromal cells. Stromal cells were suspended in DMEM:F12 Nutrient Mixture (DMEM:F12) colntaining 10% heat-inactivated charcoal-stripped FCS, 50 units/mL penicillins 50 μ g/mL streptomycin, and 1.25 μ g/mL fungizone. Cell suspensions were filtered through a 70- μ m nylon mesh to remove glands and clumps of epithelial cells. Stromal cells were plated in 3.5-mm Petri dishes and allowed to attach for 48 h at 37 °C in 5% CO₂ in air. Protein lysates were collected 24 h later in RIPA buffer for immunoblotting.

In Vitro Decidualization. Stromal cells from day 4 of pregnancy were collected by enzymatic digestion as described previously (67). The cells were cultured overnight before the initiation of decidualization by treatment with estradiol-17 β (10 nM) and progesterone (1 μ M) in phenol-red free DMEM/F12 medium supplemented with charcoal-stripped 1% FBS (wt/vol).

Measurement of Serum Estradiol-17β and Progesterone Levels. Sera were collected on days 4, 8, and 12 of pregnancy, and hormone levels were measured by enzyme immunoassay kits (Cayman) (4).

Statistics. Statistical analyses were performed using a two-tailed Student's t test. A value of P < 0.05 was considered statistically significant.

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