Identification of a Novel PP2A Regulator, WNK1, as Critical For Uterine Function

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Abstract

WNK1 is critical for uterine function as a mediator of stromal cell decidualization \textit{in vitro}. Here, we employed a mouse model with conditional WNK1 ablation from the female reproductive tract to define its \textit{in vivo} role in uterine biology. Loss of WNK1 altered uterine morphology, causing endometrial epithelial hyperplasia, adenomyosis and a delay in embryo implantation, ultimately resulting in compromised fertility. Mechanistic investigations through transcriptomic and proteomic approaches uncovered the regulatory role of WNK1 in controlling the PP2A-AKT-FOXO1 signaling axis. We show that WNK1 interacts directly with PPP2R1A, which is crucial for PP2A phosphatase activity. PP2A phosphatase in turn dephosphorylates AKT, thereby reducing its inhibitory effect on FOXO1. This permits the nuclear entry of FOXO1 to transcriptionally regulate implantation-associated genes. Our findings revealed a novel function of WNK1 in regulating AKT-FOXO1 post-translational modification, and demonstrated that this signaling pathway is critical in normal uterine physiology and pregnancy.
Introduction

The ability of the uterus to support and maintain the development of an embryo is critically dependent upon the process of implantation, the aberrant occurrence of which causes a ripple effect leading to pregnancy complications and miscarriage\textsuperscript{1}. Embryo implantation occurs during the “window of receptivity” and requires a fully prepared and receptive uterus. Investigation of uterine function mediators during receptivity identified with no lysine kinase (K) 1 (WNK1) as potentially functional in the uterus acting downstream of EGFR, whose inhibition impaired decidualization\textsuperscript{2}. Functional analysis of human endometrial stromal cells \textit{in vitro} demonstrated that WNK1 is needed for proliferation, migration and differentiation\textsuperscript{3}. Collectively, these findings indicate a previously unrecognized function of WNK1 in the female reproductive tract and led us to hypothesize that WNK1 is a mediator of uterine function.

WNK1 belongs to a family of serine/threonine protein kinases\textsuperscript{4, 5}, with its name derived from the unusual placement of the catalytic lysine in subdomain I\textsuperscript{6}. To date, Wnk1’s function is the most extensively explored in the kidney and the nervous system due to the link between its mutation and familial hypertension and autonomic neuropathy\textsuperscript{7-9}. In the renal system, WNK1 controls ion homeostasis through diverse mechanisms including activation of the SGK1/epithelial sodium channel pathway\textsuperscript{10}, regulating the potassium channel Kir1.1 cell surface localization\textsuperscript{11}, as well as controlling the activity of Na-K-Cl cotransporter through phosphorylating OSR1 (OXSR1) and SPAK\textsuperscript{12, 13}. Interestingly, WNK1’s regulatory function on OSR1 is critical for cardiovascular development, thereby contributing to embryonic lethality when WNK1 is ablated from the endothelium\textsuperscript{14, 15}. These findings suggest that although WNK1 exhibits organ-specific
physiological functions, the underlying cellular components regulated by WNK1 may share
similarity between the different tissues.

Despite its ubiquitous expression pattern, WNK1’s role in organs other than those described
above remain unexplored. Given the role of WNK1 in regulating uterine stromal cell biology in
vitro, we hypothesized that WNK1 is essential in regulating uterine functions. To test this idea, we
established a conditional uterine WNK1 ablated mouse model, which resulted in severely
compromised fertility. We demonstrated that WNK1 is critical in maintaining uterine morphology,
regulating epithelial proliferation and permitting appropriate embryo implantation. Additionally, we
defined the molecular mechanism through both transcriptomic and proteomic approaches to show
that the WNK1 signaling cascade begins by its direct interaction with the scaffold subunit of the
phosphatase complex PP2A, PPP2R1A. This interaction stabilized PP2A which prevented
unregulated AKT phosphorylation, ultimately alleviating AKT’s inhibitory effect on FOXO1 nuclear
localization, an indispensable mediator of implantation\textsuperscript{16}. 
RESULTS

WNK1 is expressed in the uterus during early pregnancy in both humans and mice

WNK1 expression was examined by immunohistochemistry in human endometrium during the proliferative and mid-secretory phases as well as in the peri-implantation uterus of mice. In humans, WNK1 is expressed in both the epithelial and stromal cells during the proliferative and mid-secretory phases (Fig. 1.A). Similarly in mice, WNK1 is expressed during and after implantation on gestation days (GDs) 4.5 and 5.5 (Fig. 1.B). These findings support the in vivo involvement of WNK1 in regulating functions of the female reproductive tract.

WNK1 ablation altered uterine morphology and microenvironment

To examine WNK1’s function in the female reproductive tract, we established a mouse model with WNK1 ablation in progesterone receptor (PGR) expressing cells, which comprised all compartments of the uterus. The conditional Wnk1 allele mice (Wnk1^ff) were crossed to the PGR^Cre mice, and confirmed that in the PGR^Cre^x^Wnk1^ff offspring (Wnk1^dd), Cre mediated excision led to the removal of exon 2 (Fig. S1. A and S1.B). Analysis of protein expression in uterine tissue via western blot confirmed reduced WNK1 protein levels in the Wnk1d/d mice (Fig. S1.C), indicating that exon 2 excision led to reduced total protein levels.

We employed a tissue clearing technique to visualize uterine morphology in 3D during the window of receptivity (Fig. 2.A and Videos 1 and 2). WNK1 deficiency substantially impacted uterine morphology, exhibiting increased number of, as well as altered structure of the endometrial glands (Fig. 2.A, Videos 1 and 2). Among the abnormalities observed in the Wnk1^dd uteri was the failure
of gland reorientation surrounding the embryo\textsuperscript{19, 20}. Examination of uterine cross sections from older mice (26 and 50 weeks) further demonstrated invasion of glands into the myometrium, suggesting that WNK1 ablation caused adenomyosis (Fig. 2.B). This was supported by the elevated expression of \textit{Moesin} (\textit{Msn}) in the Wnk1\textsuperscript{d/d} uteri, a biomarker for adenomyosis in humans (Fig. 2.C)\textsuperscript{21}. Quantification of gland number and \textit{Foxa2} gene expression showed significant elevation in the Wnk1\textsuperscript{d/d} uteri (Fig. 2.D and E), confirming the substantial increase in glandular tissues. To examine whether the increased glands were a result of increased proliferation in the uterus, we examined the expression of two mitotic markers - cyclin D1 (CCND1) and phosphorylated histone H3 (H3S10p). Elevated levels of both proteins in the glandular epithelial cells of the Wnk1\textsuperscript{d/d} uteri was observed (Fig. 2.F). In addition to the increased CCND1 and H3S10p in the glandular epithelium, higher expression of both proteins was also observed in the luminal epithelium of Wnk1\textsuperscript{d/d} uteri demonstrating that WNK1 ablation induced epithelial hyperplasia was not restricted to the glands. Moreover, we observed increased extracellular matrix deposition especially surrounding the glands in the Wnk1\textsuperscript{d/d} uteri, as shown by Masson’s trichrome staining (Fig. 2.G). These results suggest that the adenomyotic phenotype could be associated with increased epithelial proliferation as well as excessive extracellular matrix deposition\textsuperscript{22, 23}.

\textit{Uterine WNK1 ablation compromised fertility in mice through impaired implantation}

A six month breeding trial was conducted to determine the impact of WNK1 ablation on female fertility. Of the 8 control mice, 7 were able to complete the breeding trial, with 1 found dead midtrial. Necropsy showed neither pregnancy nor abnormality associated with the reproductive tract in this one mouse, indicating the cause of death was not related to abnormal uterine function. The 7 mice produced 31 litters totaling 245 pups, which was equivalent to 4.4 litters and 35 pups per mouse during the 6 months (Fig. 3.A and B). In contrast, only 4 of the 8 Wnk1\textsuperscript{d/d} mice initiated in the trial were able to complete the trial. This was due to 4 females succumbing to complications
during pregnancy or delivery. Of the 4 females that completed the trial, 10 litters and 18 pups were produced, which averaged to 2.5 litters and 4.5 pups per mouse (Fig. 3.A and B). The average litter size was also significantly smaller, averaging 3.3 pups per litter in the Wnk1\(^{dd}\) mice, compared to 7.7 in the control Wnk1\(^{ff}\) mice (Fig. 3.C). While the Wnk1\(^{ff}\) mice bred consistently, producing the last litters in the twentieth week of the trial, most of the Wnk1\(^{dd}\) mice stopped breeding after 3 litters less than 15 weeks into the trial, with only one mouse producing after the twentieth week (Fig. 3.D). Taken together, these results illustrated compromised ability to support pregnancy and progressive sterility with uterine loss of WNK1.

We next examined whether the subfertile phenotype was associated with an implantation defect in the Wnk1\(^{dd}\) mice. Dams were euthanized on GD 4.5, and embryo implantation was visualized by Evan's blue dye staining. As expected, 84.2% of the control mice successfully permitted embryo implantion on GD 4.5 while the remaining 15.8% had no embryos present in the uterus as indicated by uterine flushing (Fig. 3.E, top panel and Fig. 3.F). On the other hand, only 29.4% of the mated Wnk1\(^{dd}\) mice were able to form implantation sites on GD 4.5; however, 52.9% of the mice harboured fertilized embryos inside the uterus as identified by uterine flushing (Fig. 3.E, top panel and Fig. 3.F). Examination of the uterus on GD 5.5 showed 76.9% of the Wnk1\(^{dd}\) mice with implantation sites which is, at this time point, comparable to their Wnk1\(^{ff}\) control littermates (Fig. 3.E, bottom panel). Histological examination showed that the control Wnk1\(^{ff}\) mice had already degraded the epithelium, enabling the embryo to invade the underlying stroma (Fig. 3.G, left), while Wnk1\(^{dd}\) uteri had intact maternal epithelium at this time point with the embryo trapped inside the luminal space (Fig. 3.G). These findings demonstrate that the majority of the implantation events in the Wnk1\(^{dd}\) mice were delayed.
In addition to the uterine expression, PGR is also expressed in the ovaries and pituitary, hence WNK1 was also ablated from those tissues. We next questioned whether the implantation phenotype could be attributed to a dysfunction in the hypothalamic-pituitary-ovarian axis. Ovarian function was evaluated by assaying ovulation and ovarian steroid hormone levels. The mice were subjected to superovulatory regimen of donadotropins, followed to mating to wildtype male mice, and euthanized on GD 1.5 to simultaneously monitor fertilization of the oocytes. We found no significant decrease in the number of 2-cell embryos produced by the Wnk1^{dd} dams indicating that ovulation and fertilization was not affected (Fig. S2.A). Additionally, serum estradiol (E_2) and progesterone (P_4) levels were similar between the Wnk1^{ff} and Wnk1^{dd} mice on GD 4.5 (Fig. S2.B and C), demonstrating that the ovaries were able to produce and maintain hormone levels. These results indicate that the main contributing factor for the delayed implantation was not a malfunction of the hypothalamic-pituitary-ovarian axis.

Prerequisites for uterine receptivity are the production of leukemia inhibitory factor (LIF) from the uterine glands and the cessation of epithelial proliferation prior to implantation (GD 3.5)\textsuperscript{24,25}; as well as suppression of epithelial PGR expression and the nuclear localization of FOXO1 during implantation (GD 4.5)\textsuperscript{16,26}. Hence, we examined the uterus to see whether the delayed implantation was associated with impairment of the those parameters. Lif gene expression on GD 3.5 was similarly induced in the Wnk1^{ff} and Wnk1^{dd} mice, which decreased to basal level on GD 4.5 in the Wnk1^{ff} mice (Fig. 3.H). The Wnk1^{dd} mice showed significantly higher Lif levels on GD 4.5 (Fig. 3.H), possibly resulting from the increased glandular tissue. The proliferative markers Ki-67, PGR and FOXO1 were all deregulated in the Wnk1^{dd} mice on GD 4.5. High levels of Ki-67 were observed in the luminal epithelium of Wnk1^{dd} uteri (Fig. 3.I), indicating failure to impede epithelial proliferation. Increased PGR coupled to decreased nuclear FOXO1 and increased cytoplasmic FOXO1 were simultaneously observed in the luminal epithelium, indicating aberrant
regulation of this signaling pathway. Moreover, the underlying stromal cells also exhibited reduced nuclear FOXO1, suggesting that the cellular impact of WNK1 on FOXO1 localization may be similar in both the epithelial and stromal compartments. Our findings illustrated that several crucial implantation-associated molecular events were deregulated in the Wnk1<sup>dd</sup> mice.

**Delayed implantation negatively impacted pregnancy and embryo development in the Wnk1<sup>dd</sup> mice**

Interestingly, of the 29.4% mated Wnk1<sup>dd</sup> mice that were able to permit embryo implantation on time (GD 4.5), the number of implantation sites were similar to their Wnk1<sup>ff</sup> control littermates (Fig. 4.A). However, the number of implantation sites present on GD 5.5 was significantly lower in the Wnk1<sup>dd</sup> mice (Fig. 4.B). This finding indicated that the delay in implantation is associated with reduced number of implantation sites. Additionally, spacing between the implantation sites in the Wnk1<sup>dd</sup> mice were irregular whereas the implantation sites observed in the Wnk1<sup>ff</sup> mice were evenly distributed (Fig. 4.C). Interestingly, for the Wnk1<sup>dd</sup> mice that were able to implant promptly, implantation spacing was more evenly distributed (Fig. 3.F), suggesting that the delay impacted both implantation numbers and spacing. Examination of the uterus and embryo during mid-pregnancy (GD 8.5) further demonstrated that the Wnk1<sup>dd</sup> mice carried either resorbed embryos (Fig. 4.D, middle panel, blue arrows) or abnormally formed decidual balls (Fig. 4.D.right panel), compared to the normally sized decidual balls observed in the control mice (Fig. 4.D, left panel). Moreover, we also observed multiple embryos within one decidual zone (Fig. 4.D, red arrows), possibly from the cluttered/delayed implantation. Morphology was evaluated by examining cross sections through the center of the decidual ball, which showed that the Wnk1<sup>ff</sup> mice have vascularized and initiated placentation (Fig. 4.E, black arrows and dashed lines, respectively), both of which were lacking in the Wnk1<sup>dd</sup> uteri. Ultrasound scans demonstrated decreased gestation sac size (Fig. 4.F and G) and decreased embryo size at both GD 8.5 and GD 10.5 (Fig.
4. F and H. By GD 12.5, embryo resorption was frequently observed in the Wnk1^{d/d} mice (Fig. 4 F, bottom panel). Collectively, these findings demonstrate that uterine WNK1 ablation led to abnormal implantation and negatively impacted embryo development, resulting in the compromised pregnancy outcome and subfertility.

Loss of uterine WNK1 altered the transcriptome and kinase phosphorylation network during the receptive phase

To fully characterize the molecular mechanisms underlying the loss of WNK1 induced-implantation defect, we next examined global gene expression profile by RNA sequencing (RNA-seq) in the uterus during receptivity. To ensure that the analysis was conducted only on the maternal uterine tissues and not the embryos, we used vasectomized wild-type males to induce pseudopregnancy in the Wnk1^{f/f} and Wnk1^{d/d} mice, which was confirmed by serum progesterone levels on pseudopregnancy day (PPD) 4.5 (Table S1). In total, there were 14,423 and 14,337 genes expressed in the Wnk1^{f/f} and Wnk1^{d/d} uterus, respectively; of which 14,024 were expressed in both. The transcriptomes were subjected to principle component analysis (PCA) as a measure of quality control, which segregated according to genotype indicating that the samples were well-characterized by genotype (Fig. S3). Using a defining threshold of q-value under 0.05 for significance and fold change (FC) over 1.5 as differential expression, we identified 1,727 significantly and differentially expressed genes (DEGs) in the Wnk1^{d/d} uterus during receptivity (Table S2). We then conducted detailed analyses to characterize the molecular alterations associated with uterine WNK1 ablation using the DAVID Bioinformatic Database and Ingenuity Pathway Analysis (IPA). The top biological processes associated with the DEGs were adhesion, cell movement and locomotion, inflammation and blood vessel development (Table S3). Many important molecular functions associated with implantation were also deregulated in the Wnk1^{d/d} uteri, such as cell proliferation and apoptosis, Notch signaling, cell differentiation, epithelial to
mesenchymal transition (EMT), cytokine production, and response to estrogen. Prediction of upstream regulator activity further showed altered activity for many important receptivity mediators, including the suppression of JAG, HEY2, PTEN and SERPINE1 (Fig. 5.A). On the other hand, TGFB1, ERBB2, AKT, estrogen, ERK, MUC1 and KLF5 were predicted to show increased activity (Fig. 5.A, for the complete list see Table S4).

We have earlier shown that there was a decrease in nuclear FOXO1 during the window of implantation in the Wnk1<sup>d/d</sup> mice (Fig. 3.I). Since FOXO1 is a transcription factor that is critical for implantation, its nuclear exclusion would likely results in its reduced activity in transcribing genes. Therefore, we compared the DEGs to known FOXO1-controlled endometrial genes<sup>16</sup>, and found that of the 631 FOXO1-regulated endometrial genes, 313 showed altered expression in Wnk1<sup>d/d</sup> uterus (Fig. 5.B). Detailed comparison further illustrated that 90% of these common genes were deregulated in the same direction under WNK1 and FOXO1 deficient conditions (Fig. 5.C). These findings suggest that the impaired implantation observed in the Wnk1<sup>d/d</sup> mice was mediated partially through insufficient FOXO1 signaling during receptivity.

As WNK1 is a kinase, we next examined the alterations in the kinase phosphorylation network to understand the impact of WNK1 ablation. To this end, we employed an image-based phosphokinase array to simultaneously evaluate the phosphorylation status of multiple kinases in the uterus during receptivity (Fig. 5.D). Loss of WNK1 altered the phosphorylation of various kinases, including TOR (mTOR), SRC, PRAS40 (AKT1S1), JNK, AMPKα1 (PRKAA1), GSK-3α/β (GSK3A and GSK3B) and AKT (Fig. 5.E, all kinases with > 2 foldchange in phosphorylation are shown in Fig. S4.A). The phosphorylation of AKT, GSK-3α/β and PRAS40 were independently validated via western blotting (Fig. S4.B) and all showed elevated phosphorylation in Wnk1<sup>d/d</sup> uteri.
during receptivity. Interestingly, AKT was identified as an activated upstream regulator by IPA, and the phosphokinase array demonstrated its elevated phosphorylation in the Wnk1<sup>dd</sup> uteri during receptivity (Fig. 5.D, E and Fig. S4.B). Furthermore, elevated AKT phosphorylation on GD 4.5 was confirmed in both the epithelium and the stroma (Fig. 5.F). Indeed, we found that in the control mice, phosphorylation of AKT was actively suppressed as the mice transitioned into the receptive phase from GD 3.5 to PPD 4.5, however, the Wnk1<sup>dd</sup> mice maintained high AKT phosphorylation both prior to and during receptivity (Fig. 5.G). As AKT directly phosphorylates FOXO1 leading to its nuclear exclusion<sup>27, 28</sup>, and decreased nuclear FOXO1 was observed in the Wnk1<sup>dd</sup> uteri, we next examined FOXO1 phosphorylation. Western blot analysis showed a robust induction in FOXO1 phosphorylation during receptivity in the Wnk1<sup>dd</sup> mice, while total FOXO1 level remained constant (Fig. 5.H). These findings suggest that the elevated FOXO1 phosphorylation, and hence nuclear exclusion may be attributed to increased AKT phosphorylation.

**WNK1 ablation led to FOXO1 nuclear exclusion via AKT phosphorylation, which is associated with decreased PP2A expression and activity**

Having demonstrated that the loss of WNK1 led to increased phosphorylation of AKT and FOXO1 in mouse uteri, we next examined whether this regulatory axis was similarly maintained in human endometrial HEC1A (epithelial) and THESC (stromal) cells. Using small interfering RNA against WNK1 (siWNK1), WNK1 protein expression was inhibited, which robustly induced AKT and FOXO1 phosphorylation in both cell lines (Fig. 6.A). In order to test whether AKT facilitated FOXO1 nuclear exclusion downstream of WNK1, we next treated these cells with an AKT inhibitor, GDC0941, and examined whether it could rescue WNK1 ablation-induced phosphorylation and nuclear exclusion of FOXO1. FOXO1 localization clearly decreased in the nucleus of both cells after transfection with siWNK1 (Fig. 6.B, panels 1 VS 2, and 4 VS 5).
However, when the siWNK1 transfected cells were treated with GDC0941, nuclear FOXO1 was readily restored (Fig. 6.B, panels 3 and 6). This suggested that WNK1 inhibition-induced nuclear exclusion of FOXO1 is mediated through AKT. This is further supported by the findings that AKT inhibition rescued WNK1 knock-down induced FOXO1 phosphorylation (Fig. 6.C). Interestingly, GDC0941 treatment reduced the phosphorylation of AKT and FOXO1 to a level that is lower than seen in the siCTRL transfected, untreated cells (considered basal level). As GDC0941 inhibits AKT through its upstream regulator PI3K, it is likely that PI3K lies upstream of WNK1 in regulating AKT. Indeed, none of the PI3K family members were impacted by WNK1 inhibition, including p110-α, p110-β, p110-γ, Tyr458 phosphorylated p85 and Tyr199 phosphorylated p55 (Fig. 6.C). Similarly in mice, the expression of these proteins in the uterus were comparable between the Wnk1\textsuperscript{fl} and Wnk1\textsuperscript{dd} mice during receptivity (Fig. 6.D).

A search of the upstream regulators predicted by IPA identified candidate AKT regulators with altered activities in the Wnk1\textsuperscript{dd} uteri, including PTEN, PPP2CA, and sirolimus (rapamycin, Table S4). PTEN and PPP2CA are both phosphatases that regulate AKT phosphorylation, and both displayed repressed activities in the Wnk1\textsuperscript{dd} mice during receptivity (Z-scores of -2.079 and -1.195, respectively, Table S4). Sirolimus, on the other hand, is a drug targeting the kinase mTOR, which was strongly inhibited (Z-score of -2.95, Table S4). We found that mTOR phosphorylation and PP2A subunits A and C were altered in the Wnk1\textsuperscript{dd} mice, while PTEN level was not significantly different (Fig. 6.E). This finding suggested that increased AKT phosphorylation in the Wnk1\textsuperscript{dd} mice may be mediated through elevated mTOR or repressed PP2A activity. As mTOR is both a regulator and a substrate of AKT\textsuperscript{30, 31}, we examined whether WNK1 ablation-induced AKT phosphorylation is mediated through mTOR. We inhibited mTOR activity using rapamycin and examined AKT/FOXO1 phosphorylation as well as FOXO1 localization as a readout of AKT activity. As shown in Fig. S5.A, rapamycin treatment did not reverse the nuclear exclusion of
FOXO1 induced by WNK1 inhibition. Additionally, AKT and FOXO1 phosphorylation was not rescued by rapamycin treatment (Fig. S5.B). Similar results were observed in HEC1A cells where WNK1 and mTOR double knock-down failed to rescue AKT and FOXO1 phosphorylation (Fig. S5.C). Thus, mTOR is likely not the WNK1 mediator controlling AKT activity, and its elevated phosphorylation is a result of elevated AKT activity, rather than its cause.

WNK1 regulates AKT phosphorylation through direct interaction with PPP2R1A

WNK1 ablation led to impaired implantation arising from dysregulated AKT-FOXO1 signaling with a concomitant repression of PP2A activity and reduced PP2A subunits A and C expression. We explored the possible regulatory link between WNK1 and PP2A/AKT using a non-biased WNK1 immunoprecipitation-mass spectrometry (IP-MS) approach to identify WNK1 binding partners. Successful WNK1 IP was confirmed by examining the lysate for WNK1 expression after immunoprecipitation using a a rabbit IgG (negative control) or WNK1 targeting antibody from HEC1A cells (Fig. S6), and the peptides identified by mass-spectrometry are listed in Table S5. Amongst those were peptides belonging to WNK1 itself, as well as a known WNK1 substrate, oxidative stress responsive kinase 1 (OXSR1/OSR1)\(^1\), confirming the validity of the pull-down results (Table S5).

Putative WNK1 binding proteins identified in this experiment include Wnt regulators (OFD1 and CCDC88C), chromosome modulating and DNA repair proteins (SMCA1, KIF11, FANCI, RAD50 and SLC25A5), proteins associated with the endoplasmic reticulum and ribosomal functions (UGGT1, SEC23A, HYOU1, EMC1, AIFM1, HM13, SCFD1) as well as the mitochondria (AIFM1, SLC25A5). Of particular interest were the components of protein phosphatase complexes PP2A (PPP2R1A) and PP6 (PPP6R3), as both regulate AKT phosphorylation. Since the enzymatic
activity of PP2A was predicted by IPA as repressed in the Wnk1\(^{dd}\) mice during receptivity (PPP2CA, Table S4), we postulated that these observations were associated with the direct interaction of WNK1 and PPP2R1A, the alpha isoform of the scaffold subunit A of PP2A. In order to confirm the interaction of WNK1 and PPP2R1A, a YFP-tagged WNK1 (c4161, Fig. S7) was expressed in HEC1A cells, then immunoprecipitated using a YFP nanobody, followed by detection for PPP2R1A in the pulldown. We first confirmed that c4161 transfection induced exogenous WNK1 expression when compared to the control cells transfected with YFP only expressing construct (cYFP, Fig. 7.A). WNK1 was subsequently detected in the lysate immunoprecipitated for YFP (Fig. 7.B, upper panel), which co-immunoprecipitated with PPP2R1A (Fig. 7.B, middle panel).

Having confirmed the interaction of WNK1 and PPP2R1A, we next explored the biological implications of this interaction. The PP2A phosphatase complex is comprised of the scaffold subunit A with 2 isoforms, the regulatory subunit B with 13 isoforms and the enzymatic subunit C with 2 isoforms. As shown earlier, uterine WNK1 ablation led to decreased protein levels of subunits A and C (Fig. 6.E), yet RNA-seq showed no alteration in transcription of the 4 genes encoding these 2 subunits in the Wnk1\(^{dd}\) mice. It has been reported that the stability of the PP2A subunits depends on their association with each other\(^{32}\). Hence, reduced subunit levels could be an indication that the complexing capacity of the subunits were reduced in the absence of WNK1, leading to their degradation. We therefore postulated that the WNK1-PPP2R1A interaction is necessary for the association of the PP2A subunits. To test this idea, we examined the levels of PPP2R1A, total PP2A subunit A and total PP2A subunit C in WNK1 knock-down HEC1A cells, and accordingly found their reduced levels upon WNK1 inhibition (Fig. 7.C). Lastly, to validate that PP2A mediates AKT/FOXO1 signaling, we inhibited PPP2R1A expression in HEC1A cells using siRNA, and examined the components of the PP2A-AKT-FOXO1 signaling axis. As
expected, PPP2R1A knock-down caused a reduction in total subunits A and C of PP2A (Fig. 7.D). Interestingly, AKT phosphorylation was selectively induced on threonine 308, but not serine 473 after PPP2R1A knock-down (Fig. 7.D). This nonetheless, translated to elevated FOXO1 phosphorylation, indicating that loss of PP2A activity-induced AKT phosphorylation on this residue alone is sufficient to alter FOXO1 phosphorylation (Fig. 7.D). These findings confirmed that in endometrial cells, WNK1 controls FOXO1 phosphorylation through directly interacting with PPP2R1A, which leads to stabilization of the PP2A complex and induces AKT dephosphorylation (Fig. 7.E). In the Wnk1\textsuperscript{ld} uterus, the loss of WNK1 destabilizes and reduces PP2A activity, and this in turn, causes an elevation of AKT and subsequently FOXO1 phosphorylation, rendering FOXO1 cytoplasmic. As such, implantation-mediating genes reliant on FOXO1 transcriptional regulation becomes deregulated, ultimately resulting in impaired implantation in these mice.
Discussion

Reproductive biology has relied profoundly on transcriptomic analyses to identify novel players that may serve crucial functions in the regulation of fertility. While this approach has uncovered many key components in the reproductive tract, it is unable to detect alterations at the proteomic level, such as post translational modifications (PTMs). In many cases, the PTMs control protein activity and stability, and hence are the actual determinants of functional output. Using a proteomic approach, we identified WNK1 as a potential regulator of uterine biology with previously unreported reproductive functions. In this study, we examined the in vivo function of WNK1 using a whole-uterus WNK1 knock-out mouse model. We demonstrate that loss of WNK1 led to hyperplasia, adenomyosis and impaired implantation, which could all contribute to compromised fertility. Importantly, we demonstrate for the first time that WNK1 is a direct regulator of the PP2A-AKT-FOXO1 signaling pathway. As FOXO1 is critical for embryo implantation in mice\textsuperscript{16}, and the subfertile phenotype of Wnk1\textsuperscript{d/d} mice is associated with implantation impairments, it is likely that aberrant FOXO1 localization is at least partially responsible for the subfertility. It is worth noting that the Wnk1\textsuperscript{d/d} mice did not fully recapitulate the uterine FOXO1 knock-out phenotype, possibly due to decreased FOXO1 signaling, rather than complete inhibition.

Using a combination of in vivo and in vitro studies, we demonstrate that FOXO1 deregulation resulted from elevated AKT phosphorylation. This in turn was attributed to the reduced levels of PP2A subunits A and C, and hence reduced PP2A phosphatase activity. Interestingly, the RNA-seq results did not show altered transcription of Ppp2ca, Ppp2cb, Ppp2r1a and Ppp2r1b, which encode the different isoforms of subunits A and C. This suggests that the altered protein level was mediated through post-transcriptional or post-translational measures. As the half-life of the PP2A subunits is dependent on their ability to complex with each other\textsuperscript{32}, a possibility is that WNK1 facilitates the binding of the subunits, resulting in PP2A complex stabilization. Therefore,
in the absence of WNK1, the decreased coupling of the subunits to each other led to shorter half-lives and decreased protein levels. It is worth noting that the antibody against PP2A subunit B used in this study targets only one of the thirteen isoforms (PPP2R2A). While this particular isoform was unaffected by loss of WNK1, it is possible that other subunit B isoforms may be affected. Mechanistically, WNK1 directly interacts with PPP2R1A and hence this interaction may be crucial for PP2A complex formation. Future studies will aim to understand the biochemical nature of the PPP2R1A-WNK1 interaction, for example, whether WNK1 phosphorylates PPP2R1A, and if so, whether the phosphorylation interferes with PP2A subunits coupling. As WNK1, PP2A, AKT and FOXO1 are all ubiquitously expressed with crucial functions, this signaling cascade is likely to have broad-spectrum functions in other tissues.

Whilst we were able to demonstrate the dysregulation of PP2A-AKT-FOXO1 signaling pathway in the Wnk1<sup>dd</sup> mice and the associated implantation defect, it is likely that the effects of WNK1 ablation on uterine biology is not limited to implantation. Given that 30% of the Wnk1<sup>dd</sup> mice were able to implant promptly with normal numbers of embryos, we rarely observed normal sized litters from these mice. This suggests that even after a seemingly normal implantation, there are likely other impairments in subsequent pregnancy development accounting for the compromised fertility. Indeed, gene expression analysis indicated altered signaling in many receptivity regulators including suppressed Notch signaling (HEY2 and JAG2)<sup>33</sup>, elevated ERK<sup>34</sup> and MUC signaling<sup>35</sup>, which are all important decidualization mediators. Additionally, WNK1 is a major ion channel regulator in renal cells<sup>36</sup>, and ion channels are also significant players in endometrial functions<sup>37</sup>. For example, the cAMP-activated anion channel cystic fibrosis transmembrane conductance regulator (CFTR) and epithelial sodium channel (ENaC) are two ion channels responsible for regulating uterine fluid flow<sup>38, 39</sup>, and both are WNK1-regulated in other systems<sup>10, 40</sup>. Interestingly, there was a 2.7-fold increase in Cftr expression in the Wnk1<sup>dd</sup> uteri during
implantation (Table S2), suggesting that WNK1 may regulate the expression of this gene’s RNA in addition to its reported regulatory function at the post-translational level. Moreover, WNK1 is expressed in the placenta, indicating that WNK1 may also have placental functions and hence a role in supporting embryo development during mid-pregnancy\textsuperscript{41}. These findings offer additional mechanisms through which WNK1 ablation could potentially impair fertility in mice, and will be addressed in future studies to explore WNK1’s alternative uterine functions.

As aforementioned, uterine WNK1 ablation exhibited pleiotropic effects, one which was particularly apparent was the epithelial abnormality in the Wnk1\textsuperscript{d/d} uteri. We observed epithelial hyperplasia which manifested through abnormal gland growth and associated with increased numbers of CCND1 and H3S10P positive cells. This was accompanied by adenomyosis which is characterized by extension of the glands into the myometrium. Adenomyosis is known to reduce implantation\textsuperscript{42}, which is associated with dysregulated adhesion molecules expression, including ITGB3 and SPP1\textsuperscript{43}. Interestingly, we found elevated Spp1 expression in the Wnk1\textsuperscript{d/d} uteri during implantation, revealing another possible mechanism of impaired implantation in the Wnk1\textsuperscript{d/d} mice. In addition to implantation, adenomyosis negatively impact pregnancy in several other ways including abnormal muscle contractility, inflammation and metabolism\textsuperscript{42, 44, 45}. Interestingly, the ERM protein family member moesin (encoded by the MSN gene) has been correlated with adenomyosis in humans\textsuperscript{21}, and we accordingly found higher Msn expression in the Wnk1\textsuperscript{d/d} mice. This finding suggests that WNK1 ablation-induced adenomyosis may share molecular similarities to this disease in humans. We propose that the uterine WNK1 ablation mouse model could be employed to study this disease.
Although neither epithelial hyperplasia nor adenomyosis are cancerous, both are progressive conditions which may lead to malignant transformation\textsuperscript{46, 47}. Functional interpretation of the transcriptome reiterated this, where many cancer development and progression associated signaling pathways were altered in the Wnk1\textsuperscript{d/d} uteri, including elevated TGFB, AKT and estrogen\textsuperscript{47-50}. Interestingly, a recurrent mutation of \textit{Ppp2r1a} is associated with serous endometrial carcinoma\textsuperscript{51, 52}, and this mutation has been found to impact oncogenic signaling through a dominant negative effect\textsuperscript{53}. It is possible that WNK1 could act to protect cells against cancer progression through stabilizing PP2A subunits and hence activity. We speculate that WNK1 signaling in the uterus also acts to maintain tissue homeostasis, such as preventing unregulated epithelial proliferation. As such, uterine loss of WNK1 displayed epithelial hyperplasia and may promote cancer development with aging.

In summary, this study is a first to explore the reproductive function of WNK1 \textit{in vivo}. We demonstrate that WNK1 is critical in maintaining normal uterine morphology, mediating epithelial homeostasis and implantation. Future studies will further explore the molecular actions of this complex protein in female reproductive tract, as well as the significance in human fertility and pathological outcomes when WNK1 is deregulated.
Online Methods

Ethics statement

This study was conducted according to the federal regulations regarding the use of human subjects. Procedures were approved by the following ethics committee: Institutional Review Board/Committee-A (IRB) of Greenville Health System under IRB file #Pro0000093 and Pro00013885 and the University of Chapel Hill at North Carolina IRB under file #: 05-1757. Written, informed consents were obtained from all patients prior to participation.

All animal studies were conducted in accordance with the Guide for the Care and Use of Laboratory Animals, as published by the National Institute of Health. Animal protocols were approved by the Animal Care and Use Committee (ACUC) of National Institute of Environmental Health Sciences (protocol numbers 2015-0012 and 2015-0023). The mice were housed with a maximum of 5 per cage with a 12-hour light and dark cycle, and fed irradiated Teklad global soy protein-free extruded rodent diet (Harlem Laboratories, Inc., Indianapolis, IN) and fresh water ad libitum. Euthanization was carried out by carbon dioxide inhalation followed by cervical dislocation.

Generation of transgenic mice

The Wnk1<sup>f/f</sup> mice were a kind gift from Dr CL Huang (University of Iowa Healthcare), the generation of which was described in a previous publication<sup>14</sup>. Briefly, the Wnk1 floxed allele was established by insertion of loxP sites into the 5' and 3' region of exon 2 of the mouse Wnk1 gene.
The Wnk1^{ff} mice were crossed to mice carrying Cre under the control of the progesterone receptor (PGR^{Cre}) to generate conditional uterine Wnk1 ablated mice (Wnk1^{dd}, Fig. S1) \(^{17}\).

**Fertility trial**

Seven-week old Wnk1^{ff} (control) and Wnk1^{dd} (experimental) mice were housed with wild-type CD1 males for a period of 6 months, during which the mice were monitored daily for pregnancy and delivery. Upon the first observation of delivery, the total number of pups including both live and dead were recorded.

**Implantation determination and pseudopregnancy**

Mice of age 6-10 weeks were housed with wild-type C57BL/6J males, and monitored each morning until vaginal plug is observed (indicating that mating has occurred). The first noon following which the vaginal plug was seen was defined as gestation day (GD) 0.5. Mice were anesthetized by isoflurane inhalation on GD 4.5 and 5.5, followed by retro-orbital administration of 200 µL of 1% Evans Blue Dye prepared in phosphate buffered saline (PBS). Mice were then euthanized, and the uterine horns harvested for implantation determination and imaging. For mice sacrificed on GD 4.5, the uterine horns were flushed using a p200 pipette if no implantation sites were observed, and the eluant was examined under brightfield microscope to determine presence of blastocysts. The uterine horns were then fixed 48 hours in 4% paraformaldehyde (PFA) prepared in PBS for histology and immunohistochemistry, or frozen on dry ice for RNA and protein extraction. For RNA-seq, pseudopregnancy was induced by mating the females to vasectomized wild-type male mice, and all procedures were conducted as described above except for the Evans Blue Dye injection, as no implantations were expected.
**Superovulation assay**

Three-week old Wnk1^f/f and Wnk1^d/d mice were subjected to a superovulation regimen, which began with intraperitoneal administration of 5 IU of pregnant mare’s serum gonadotropin (catalog no. 493-10-2.5, Lee Biosolutions), followed by 5 IU of human chorionic gonadotropin (catalog no. 869031, EMD Millipore) 48 hours later. Superovulated mice were placed with wild-type CD1 males overnight. Mating was confirmed by presence of vaginal plug the next morning (GD 0.5), and mice were euthanized on GD 1.5 followed by oviduct flushing. The number of embryos was determined by counting under a brightfield microscope.

**Serum collection**

On GD or pseudopregnancy day (PPD) 4.5, mice were anesthetized by intraperitoneal administration of Fetal Plus (1mg/10g body mass) and whole blood was collected via retro-orbital puncture. Blood was allowed to clot at room temperature for approximately 30 minutes, then centrifuged at 1000 X G for 10 minutes at 4°C. The supernatant (serum) was moved into a fresh tube and stored at -80°C until hormone testing. Hormone testing was conducted by the Ligand Core Laboratory of University of Virginia, Center for Research in Reproduction.

**High frequency ultrasound imaging**

On GDs 8.5, 10.5 and 12.5, high frequency ultrasound imaging was used to evaluate the uterus and embryo development. Dams were anesthetized by isoflurane inhalation and placed onto an electric heating pad to maintain body temperature. Abdominal hair was removed using depilatory cream (Nair™ Church & Dwight Co. Trenton, NJ), and eye lubricant was applied to prevent desiccation. Dams were manipulated into a supine position for the scan while heart rate and body
temperature were continuously monitored. Images were visualized and captured using the VisualSonics VevoR 2100 Imaging System with a 550s scan head (Fujifilm VisualSonics Inc., Toronto, ON) at 55 megahertz. Each scanning session was limited to maximum 15 minutes, after which the dams were monitored until full recovery.

**Tissue processing, histology, immunohistochemical and immunofluorescence staining**

For histology, immunohistochemistry and immunofluorescence, the tissues were similarly processed as described below. After 48 hour fixation, tissues were placed into 70% ethanol for a minimum of 48 hours. Tissues were then dehydrated and embedded in paraffin blocks and sectioned to 5 µm thickness onto glass slides. Slides were heated at 60°C for 10 minutes, followed by 5 minutes cooling. Sections were deparaffinized by 3 serial incubations in Citrisolv clearing agent (catalog no. 22-143-975, Thermo Fisher), followed by rehydration through decreasing % of ethanol. For histology, sections were subjected to hematoxylin and eosin (H&E) and Masson’s trichrome staining, followed by dehydration through increasing % of ethanol, incubation in Citrisolv and coverslipping. For immunohistochemistry, sections were subjected to antigen retrieval after rehydration by boiling in the Vector Labs Antigen Unmasking Solution as per manufacturer’s instructions (H-3300, Vector Laboratories, Burlingame, CA, USA). Blocking of endogenous peroxidase was performed by treating the sections with 3% hydrogen peroxide diluted in distilled water for 10 minutes at room temperature. Tissues were blocked in 5% normal donkey serum (NDS) for 60 minutes at room temperature, prior to overnight incubation with the primary antibody at 4°C. The slides were washed twice in PBS for a total of 10 minutes at room temperature and secondary antibody diluted in 1% w/v bovine serum albumin (BSA) prepared in PBS was applied. The ABC reagent was applied to tissue according to the manufacturer’s instructions (Vector Labs ABC PK-6100, Vector Laboratories). Signals were developed using the Vector Labs DAB ImmPACT Staining Kit (Vector Labs SK-4105, Vector Laboratories). Finally, the tissues sections
were counterstained with hematoxylin and dehydrated through increasing ethanol concentration, followed by Citrisolv incubation and coverslipping. For immunofluorescence, tissue sections were subjected to antigen retrieval as described above. Tissues were blocked in 0.4% v/v Triton X-100, 1% BSA and 5% NDS for 30 minutes at room temperature followed by overnight incubation in primary antibody prepared in 0.4% Triton X-100/PBS at 4°C. Sections were washed 3 times 5 minutes in PBS and incubated with secondary antibodies diluted in 0.4% Triton X-100/PBS for 90 minutes at room temperature. Finally, the slides were washed 3 times 5 minutes in PBS, and coverslipped in DAPI containing mounting medium (Vectorshield Hardset™ Antifade Mounting Medium, catalog no. H-1400, Vector Laboratories). Details of antibodies used in this study are provided in table S6.

**RNA extraction and cDNA conversion**

The frozen tissues were disrupted in TRIzol reagent (Thermo Fisher) by bead milling, followed by 2 aqueous phase separations using 1-Bromo-3-chloropane and chloroform. Pure ethanol was added to the aqueous layer, and the RNA was extracted using the Qiagen RNEasy RNA mini prep kit columns as per manufacturer's instructions (Qiagen, Valencia, CA). Resulting RNA concentration and quality was determined using the NanoDrop ND-1000. cDNA was generated by reverse transcription using the M-MLV Reverse Transcriptase (catalog number 28025013 Thermo Fisher) following the manufacturer's instructions.

**qRT-PCR**

qRT-PCR was performed using the SsoAdvanced™ Universal SYBR Green Supermix (1725274, Bio-Rad) with the following primers (from 5’ to 3’, F = forward and R = reverse): Wnk1 – AGGCAGAGATTCAAGAAGAGG (F) and CCCAGGAATCATAGAATCG (R); Msn –
CCATGCCGAAGACGATCA (F) and CCAACTTCCCTCAAACCAATAG (R); and Foxa2 – GAGACTTTGGAGAGCTTTGAG (F) and GATCACTGTGGCCCATCTATT (R). Lif expression was determined using the Taqman Master Mix (Life Technologies) and Taqman probes (Applied Biosystems). The Delta delta Ct values were calculated using 18S RNA control amplification results to acquire the relative mRNA expression for each sample.

RNA-sequencing

For each mouse, 1 µg of total uterine RNA was sent to the NIH Intramural Sequencing Center to create a library using the TruSeq RNA Kit (Illumina, San Diego, CA, USA) following the manufacturer’s instructions. The RNA libraries were sequenced with a HiSeq 2000 System (Illumina). The raw RNA reads (75 nt, paired-end) were processed by filtering with average quality score greater than 20. Reads that passed the initial processing were aligned to the mouse reference genome (mm10; Genome Reference Consortium Mouse Build 38 from December 2011) using TopHat version 2.0.4. Expression values of RNA-seq were expressed as fragments per kilobase of exon per million fragments (FPKM). Differential expression was calculated using Cuffdiff function from Cufflinks version 2.2. Transcripts with the average FPKM > 1 in at least one group, q-value < 0.05 and at least 1.5-fold difference in FPKM were defined as differentially expressed genes (DEGs). The data discussed in this publication have been deposited in NCBI’s Gene Expression Omnibus and are accessible through GEO Series accession number GSE144802.

Human phospho-kinase antibody array

Site specific phosphorylation levels of 43 kinases were measured using the Human Phospho-Kinase Array Kit (catalog no. ARY 003B, R&D Systems) according to the manufacturer’s
instructions with the experimental design as described below. Pseudopregnancy was induced in
the Wnk1^{tf} and Wnk1^{dd} mice as previously described, and mice were euthanized on PPD 4.5.
Uterine tissues were frozen at -80\(^\circ\)C until ready to proceed. Lysate were extracted independently
from 6 mice per group by bead milling in the lysis buffer provided within the kit, and protein
concentrations were determined using the BCA Protein Assay Kit (catalog no. 23225, Pierce).
Equal amounts from each mouse were pooled in each group (to a total of 900 µg), and the
remaining steps followed the standard protocol of the kit. Signal intensity was quantified by
ImageJ\textsuperscript{56}.

**Protein extraction from uterine tissues and protein expression analysis**

Tissues were homogenized in RIPA Lysis and Extraction Buffer (Thermo Fisher) supplemented
with protease inhibitor cocktail (cOmplete Mini, EDTA-free, catalog no. 11836170001, Roche
Diagnostics) and phosphatase inhibitor cocktail (phosSTOP, catalog no. 4906837001, Roche
Diagnostics), followed by centrifugation at 10,000 \(\times\) G for 10 minutes at 4\(^\circ\)C, and the supernatant
was moved into fresh eppendorf. Protein concentrations were measured using the BCA Protein
Assay Kit (Pierce). Heat denatured protein samples were resolved using 7.5%, 10% or gradient
4-20% Criterion Tris-HCl precast gels (Bio-Rad), followed by transferring using the Trans-Blot
Turbo Transfer System (Bio-Rad), as according to the manufacturer’s instructions. PVDF and
nitrocellulose membranes were used for target proteins \(> 200\) KDa and \(< 200\) KDa, respectively.
After transfer, the membranes were blocked in 5% w/v non-fat milk or BSA prepared in Tris
buffered saline with 0.1% Tween-20 (TBST). Membranes were incubated with primary antibody
at 4\(^\circ\)C with shaking overnight, followed by three 10 minute washes in TBST the next morning.
Membranes were proceeded to secondary antibody incubation at room temperature for at least
one hour with shaking, and washed another 3 times in TBST. Depending on the expected signal
strength, different peroxidase chemiluminescent substrates were used: KPL LumiGLO\textsuperscript{R} (catalog
no. 546101, Seracare), Clarity Western ECL Substrate (catalog no. 1705060, Bio-Rad), and Amersham ECL Prime Western Blotting Detection Reagent (catalog no. RPN2232, GE Healthcare Life Sciences). Antibody sources and dilutions are summarized in supplemental table 6. For each western blot, GAPDH or β-tubulin were detected as the loading control, and in cases where the target protein is in the same region as the loading control proteins, a duplicate gel was ran and transferred in parallel. For each set of samples, a representing GAPDH or β-tubulin blot is shown. Details of antibodies used in this study are provided in table S6.

Tissue clearing and three-dimensional reconstruction

Uterine tissues were fixed in 4% PFA for 16 hours, followed by 3 rinses in PBS. Tissues were then incubated in hydrogel monomer solution AP40 (4% v/v acrylamide and 0.25% w/v VA-044 in PBS) for 72 hours at 4°C, protected from light. Oxygen was then removed from the samples using a chamber connected to vacuum and nitrogen, followed by incubation at 37°C for 3 hours to initiate tissue-hydrogel hybridization. Hydrogel was removed from the tissues via 3 PBS washes, and tissues were subsequently incubated in 8% SDS prepared in PBS for 7 days at 37°C with shaking, and the SDS solution replaced twice during incubation. The tissues were then washed 5 times one hour in PBS and blocked in 5% NDS prepared in PBS/triton X-100 with 0.01% of sodium azide. The samples were then incubated in primary antibody prepared in antibody diluent (2% v/v NDS, 0.01% w/v sodium azide in PBST) for 6 days at room temperature with constant rotation, followed by 5 one hour washes in 0.1% v/v Triton in PBS (PBS-T). Secondary antibody was similarly prepared in antibody diluent and incubated for another 6 days at room temperature with constant rotation and protected from light, replacing antibody half way through incubation. Finally, the samples were washed an additional 5 times one hour in PBS-T and incubated in Refractive Index Matching Solution (80% w/v Histodenz (catalog no. D2158, Sigma-Aldrich) prepared in 0.02M phosphate buffer, pH7.5 with 0.1% Tween-20 and 0.01% sodium azide,
refractive index = 1.46) for 1-3 days, and samples were mounted in fresh Reflective Index Mounting Solution using a 1 mm deep iSpacer (www.sunjinlabs.com). Details of antibodies used in this study are provided in table S6.

Cell culture

Human endometrial epithelial cell line HEC1A and telomerase-transformed human endometrial stromal cells (THESC) were obtained from American Type Culture Collection (ATCC, Rockville, MD, USA). HEC1A cells were cultured in McCoy’s 5A modified medium (catalog no. 16600082, Gibco) and the THESC cells were maintained in DMEM/F12 (1:1) (catalog no. 11330-032, Gibco), both supplemented with 10% fetal bovine serum (FBS, catalog no. 10437-028, Gibco) and 100 U/mL penicillin and 100 µg/mL streptomycin, unless otherwise stated.

siRNA transfection and drug treatments

Cells were transfected with siRNAs using the Lipofectamine RNAiMax transfection reagent (catalog no. 13778150, Thermo Fisher) following the manufacturer’s protocol. Cells were transfected with 24 – 72 nM siRNA in transfection medium supplemented with 2% charcoal-stripped FBS (catalog no. 12676-029, Gibco) for 24-48 hours before replacing with fresh growth medium. Proteins were harvested from cells 72 hours after transfection unless otherwise stated. The siRNAs used in this study were: nontargeting siRNA (siCTRL, catalog no. D-001810-10-20, Dharmacon), Wnk1 targeting siRNA (siWNK1, catalog no. L-005362-02-0005, Dharmacon), Mtor targeting siRNA (simTOR, catalog no. L-003008-00-0005, Dharmacon), and Ppp2r1a targeting siRNA (siPPP2R1A, catalog no. L-060647-00-0005, Dharmacon). AKT and mTOR inhibitors GDC0941 and rapamycin (catalog no. S1065 and S1039, respectively, Selleckchem) were
dissolved in DMSO, and cells were treated with 5 µM GDC 0941 and 10 – 40 µM rapamycin for 24 hours, while the control cells received equivalent volumes of DMSO.

**Immunofluorescence of cultured cells**

Cells were cultured in 4-chambered coverglass (catalog no. 155382, Thermo Fisher) as described above. Following transfection and/or drug treatment for the appropriate time period, cells were rinsed in cold PBS, fixed in 4% PFA and permeabilized in 0.5% Triton X-100/PBS for 10 and 5 minutes, respectively, at room temperature. Cells were then incubated in blocking buffer (5% v/v NDS, 0.2% v/v fish gelatin (catalog no. G7765, Sigma-Aldrich), 0.2% v/v Tween-20 in PBS) for 30 minutes at 37°C. Primary antibody was diluted in blocking buffer and added to the cells for 60 minutes, followed by secondary antibody for another 60 minutes; both incubation steps were performed at 37°C in a humidified chamber. Finally, cells were rinsed 3 times with 0.2% Tween-20/PBS and coverslipped using a DAPI containing mounting medium (Vectorshield Hardset™ Antifade Mounting Medium, catalog no. H-1400, Vector Laboratories). Details of antibodies used in this study are provided in table S6.

**WNK1 Immunoprecipitation Mass-spectrometry**

HEC1A cells were grown to 70% confluency, followed by collection using trypsin. Cells were washed 2 X in cold PBS, followed by resuspension in cell lysis buffer (50 mM Tris-HCl pH 7.5, 150 mM NaCl, 1 mM EDTA, 1% NP-40, 1% sodium deoxycholate, 0.1% SDS, with protease and phosphatase inhibitors added fresh to 1 X). Cells were incubated on ice for 10 minutes, followed by sonication on medium power (3 X 5 seconds). Lysate was centrifuged at 13,000 X G for 10 minutes at 4°C. WNK1 targeting antibody was added at 1:100 to the supernatant, and incubated with rotation at 4°C overnight. Prewashed beads (50% protein A and 50% protein G, catalog
numbers 10002D and 10004D, respectively, Thermo Fisher) were added to the immunocomplex and incubated for 30 minutes at room temperature with rotation. Beads were pelleted using a magnetic separation rack, followed by 3 washes in lysis buffer. Beads were heated to 100°C with 3 X SDS buffer (150 mM Tris-HCl pH 6.8, 6% SDS, 0.3% BPB, 30% glycerol, 3% B-mercaptoethanol) for 5 minutes, before electrophoresis through a 7.5% Criterion Tris-HCl precast gel (Bio-Rad). Gel regions were excised from the SDS-PAGE gel and minced, and digests were performed with a ProGest robotic digester (Genomic Solutions) where the gel pieces were destained by incubation in 25 mM ammonium bicarbonate with 50% acetonitrile (v/v) twice for a total of 30 minutes. The gel pieces were dehydrated in acetonitrile, followed by drying under a nitrogen stream, and further incubated with 250 ng trypsin (Promega) for 8 hours at 37°C. The digests were collected, and peptides were re-extracted three times. The extractions were pooled for each sample, lyophilized and resuspended in 20 µL 0.1% formic acid. The protein digests were analyzed by LC/MS on a Q Exactive Plus mass spectrometer (Thermo Fisher) interfaced with a nanoAcquity UPLC system (Waters Corporation), and equipped with a 75 µm x 150 mm BEH dC18 column (1.8 µm particle, Waters Corporation) and a C18 trapping column (18 µm x 20 mm) with a 5 µm particle size at a flow rate of 400 nL/min. The trapping column was positioned in-line of the analytical column and upstream of a micro-tee union which was used both as a vent for trapping and as a liquid junction. Trapping was performed using the initial solvent composition. A volume of 5 µL of digested sample was injected into the column, and peptides were eluted by using a linear gradient from 99% solvent A (0.1% formic acid in water (v/v)) and 1% solvent B (0.1%formic acid in acetonitrile (v/v)), to 40% solvent B over 60 minutes. For the mass spectrometry, a data dependent acquisition method was employed with an exclusion time of 15 seconds and an exclusion of +1 charge states. The mass spectrometer was equipped with a NanoFlex source and was used in the positive ion mode. Instrument parameters were as follows: sheath gas, 0; auxiliary gas, 0; sweep gas, 0; spray voltage, 2.7 kV; capillary temperature, 275°C; S-lens, 60; scan range (m/z) of 200 to 2000; 2 m/z isolation window; resolution: 70,000; automated
gain control (AGC), 2 X 10^5 ions; and a maximum IT of 200 ms. Mass calibration was performed before data acquisition using the Pierce LTQ Velos Positive Ion Calibration mixture (Thermo Fisher). Peak lists were generated from the LC/MS data using Mascot Distiller (Matrix Science) and the resulting peak lists were searched using the Spectrum Mill software package (Agilent) against the SwissProt database. Searches were performed using trypsin specificity and allowed for one missed cleavage and variable methionine oxidation. Mass tolerance were 20 ppm for MS scans and 50 ppm for MSMS scans.

**Generation of mammalian YFP-WNK1 expression constructs**

The coding region of the WNK1 sequence (NM_014823.3) with attL sites and N-terminal TEV cleavage site was synthesized by GeneWiz Inc. and cloned into pUC57 (Kanamycin) plasmid. Gateway Cloning using LR Clonase II mix (Thermo Fisher) was used to transfer the WNK1 sequence into the Vivid Colors pcDNA6.2/N-YFP vectors (Thermo Fisher), which created the mammalian expression vectors with YFP fused to the N-terminal end of WNK1 (Fig. S7, c4161).

**Co-Immunoprecipitation**

HEC1A cells were transfected with cYFP or c4161 for 48 hours, followed by trypsinization, 3 washes and resuspension in lysis buffer (50 mM Tris pH8.0, 400 mM NaCl, 0.1% NP-40 and 0.5 mM DTT, with protease and phosphatase inhibitors freshly added to 1 X). The lysate was incubated at 4°C with rotation for 30 minutes. Lysates were centrifuged at 21,100 X G for 10 minutes at 4°C, and the supernatant was added to 1.5 volumes of 25% glycerol, followed by centrifugation at 21,100 X G for 10 minutes at 4°C. Anti-GFP resin slurry was added to the supernatant and nutated for 1 hour at 4°C. Beads were centrifuged at 1,000 X G for 5 minutes, 4°C, followed by 6 washes in 100 µL of PBST in Bio-Spin columns (catalog number 7326204,
Bio-Rad). The bound immunocomplexes were eluted via 0.1 M glycine, pH 2.0, and eluent was neutralized using 2M Tris-HCl, pH 8.0.

Confocal Microscopy

All fluorescent images presented in this study were captured using the Zeiss LSM 780 UV confocal microscope.

Data analysis

GraphPad Prism versions 7 and 8 were used for data analysis. Each set of data points were first subjected for normality tests. Student’s t tests and Mann-Whitney tests were performed for normally distributed data and non-normally distributed data, respectively. In each case, a p-value less than 0.05 was considered as significant. Functional annotation for the differentially expressed genes derived from RNA-seq were analyzed by Ingenuity Pathway Analysis (IPA) and Database for Annotation, Visualization, and Integrated Discovery (DAVID)\textsuperscript{57}. 

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Author Contributions

Conceptualization, R.A.C., S.P.W. and F.J.D.; Methodology, R.A.C. and F.J.D.; Validation, R.A.C.; Formal Analysis, R.A.C. and T.W.; Investigation, R.A.C.; Resources, S.L.Y., J.L., C.L.H. and F.J.D.; Data Curation, R.A.C. and T.W.; Writing – Original Draft, R.A.C.; Writing – Review & Editing, R.A.C. and F.J.D.; Visualization, R.A.C.; Supervision, F.J.D.; Project Administration, R.A.C. and F.J.D.; Funding Acquisition, F.J.D.
Declaration of Interests

The authors declare no competing interests.
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**Figure 1.** WNK1 is expressed in the uterus during the window of implantation in both humans and mice. (A and B) Immunohistochemical staining of WNK1 in proliferative and mid-secretory phased endometrial tissues from fertile women (A), and during receptive gestation day (GD) 4.5 and post implantation GD 5.5 in the uterus of wild-type mice (B). IgG served as negative controls, scale bars = 50 µm.
**Figure 2. WNK1 ablation altered uterine morphology and microenvironment.** (A) Three-dimensional reconstruction of Wnk1\(^{flf}\) and Wnk1\(^{dd}\) uteri on GD 4.5 using tissue clearing and confocal microscopy. The glands, myometrium and embryo were marked by FOXA2 (green), ACTA2 (red) and OCT4 (purple), respectively. Images were captured by tile-scanning and Z-stacking, and reassembled *in silico* using Imaris software. White arrow indicates position of the embryo. Scale bars = 500 µm. (B) Immunofluorescence of uterine cross section showing glands (FOXA2, green) and myometrium (ACTA2, red) from Wnk1\(^{flf}\) and Wnk1\(^{dd}\) uteri. White arrows indicate gland extension into myometrium. Scale bars = 50 µm. (C) Adenomyosis biomarker Msn mRNA expression as determined by qRT-PCR (n = 4). (D) Quantification of number of glands per cross section for Wnk1\(^{flf}\) and Wnk1\(^{dd}\) mice (n = 6), and (E) quantification of Foxa2 mRNA expression as determined by qRT-PCR (n = 6). (F) Expression of mitotic markers CCND1 and H3S10P in the uterus of 26-week-old Wnk1\(^{flf}\) and Wnk1\(^{dd}\) mice, scale bars = 100 µm. (G) Masson’s trichrome staining of uteri cross section from 26 and 50 week old Wnk1\(^{flf}\) and Wnk1\(^{dd}\) mice, scale bars = 100 µm. All quantitative results shown are average ± SD, \( * p < 0.05 \).
Figure 3. Uterine WNK1 ablation compromised fertility and impaired implantation in mice.

(A) Numbers of litters and pups produced by Wnk1^{ff} and Wnk1^{dd} dams during the 6-month breeding trial with wild-type males. (B) Average number of litters produced, (C) average number of pups per litter produced and (D) time of last delivery, n = 7 for Wnk1^{ff} mice and n = 4 for Wnk1^{dd} mice. Results shown are mean ± SD, * p < 0.05. (E) Percentage of mated Wnk1^{ff} and Wnk1^{dd} mice with implantation (green), without implantation (pink), and without implantation but presented embryos in the uterus (blue) on GD 4.5 and GD 5.5, n = 19 and 12 for Wnk1^{ff} mice on GD 4.5 and GD 5.5, respectively; and n = 17 and 13 for Wnk1^{dd} mice on GD 4.5 and GD 5.5, respectively. Red arrows indicate position of implantation sites. (F) Gross uterine morphology of Wnk1^{ff} and Wnk1^{dd} mice on GD 4.5, scale bars = 1 cm. (G) Hematoxylin and eosin staining of uterine cross sections at implantation site on GD 5.5 in Wnk1^{ff} and Wnk1^{dd} uteri, arrow indicates presence of maternal epithelium, and E = embryo. Scale bars = 100 µm. (H) Implantation marker Lif mRNA expression in the uteri as determined by qRT-PCR on GD 3.5 and GD 4.5 for Wnk1^{ff} and Wnk1^{dd} mice. Results shown are mean ± SD, * p < 0.05. (I) Expression of proliferative marker Ki-67, implantation markers PGR and FOXO1 on GD 4.5 in the stroma and epithelium of Wnk1^{ff} and Wnk1^{dd} mice. LE = luminal epithelium and S = stroma, scale bars = 25 µm.
Figure 4. Delayed implantation negatively impacted pregnancy and embryo development in the Wnk1<sup>dd</sup> mice. (A – B) Number of implantation sites on GD 4.5 (A) and GD 5.5 (B) in Wnk1<sup>ff</sup> and Wnk1<sup>dd</sup> mice (n = 6). (C – D) Uterine gross morphology on GD 5.5 (C) and GD 8.5 (D), with implantation sites on GD 5.5 marked by red arrows, and blue arrows indicate resorption and abnormal decidualization on GD 8.5. Scale bars = 1 cm. (E) Hematoxylin and eosin staining of cross section through the centre of decidual ball on GD 8.5 from Wnk1<sup>ff</sup> and Wnk1<sup>dd</sup> mice, with black arrows and dashed line indicating vascularization and placentation, respectively. Scale bars = 2 mm. (F) Ultrasound scans of uterus and embryo during mid-pregnancy at GD 8.5, 10.5 and 12.5. Scale bars = 2 mm. (G and H) Quantification of gestation sac size by length and width on GD 8.5 (G), and embryo size by crown-rump length (CRL) on GD 8.5 and 10.5, as measured from ultrasound scans (n = 6). Results shown are mean ± SD, *p < 0.05.
Figure 5. Loss of uterine WNK1 altered the transcriptome and kinase phosphorylation network during the receptive phase. (A) Activity of upstream regulators as predicted by Ingenuity Pathway Analysis based on the altered uterine transcriptome of Wnk1<sup>d/d</sup> mice on PPD 4.5. See Table S4 for complete list. (B) Comparison of differentially expressed genes (DEGs) between the uteri of WNK1 ablated (blue) and FOXO1 ablated (pink) mice identified 313 common genes. (C) Percentage of the 313 genes categorized into commonly upregulated (pink), commonly downregulated (green), or upregulated in one and downregulated in the other (yellow and blue). (D and E) Kinome phosphorylation status in Wnk1<sup>d/d</sup> and Wnk1<sup>fr</sup> uteri on pseudopregnancy day (PPD) 4.5, with selected alterations shown in (E). All kinases with > 1.5 fold change in signal intensity as quantified by ImageJ is shown in Fig. S4. Results were acquired using pooled uterine lysate from 6 mice in each group. (F and G) Expression of phosphorylated and total AKT in Wnk1<sup>fr</sup> and Wnk1<sup>d/d</sup> uteri on GD 4.5 as shown by immunohistochemistry ((F), LE = luminal epithelium and S = stroma), and on GD 3.5 and PPD 4.5 as shown by western blotting (G), scale bars = 25 µm. (H) Western blot analysis showing levels of phosphorylated and total FOXO1 in Wnk1<sup>fr</sup> and Wnk1<sup>d/d</sup> uteri on PPD 4.5.
A

B

C

D

E
Figure 6. WNK1 ablation led to FOXO1 nuclear exclusion via AKT phosphorylation which is associated with decreased PP2A phosphatase expression. (A) Western blot showing levels of phosphorylated and total AKT and FOXO1 in HEC1A and THESC cells transfected with 24 nM siCTRL or siWNK1. (B) Immunofluorescence showing FOXO1 subcellular localization (green), with nuclei presented in DAPI in HEC1A and THESC control cells (1, 4), siWNK1 transfected cells (2, 5), and GDC0941 treated, siWNK1 transfected cells (3, 6), scale bars = 20 µm. (C) Expression of FOXO1, AKT and PI3K members in HEC1A and THESC cells transfected with siCTRL, siWNK1, and treated with AKT inhibitor GDC0941. (D and E) Expression of PI3K proteins (D), and mTOR, PP2A subunits and PTEN (E) in Wnk1^f/f and Wnk1^d/d uteri on PPD 4.5.
Figure 7. WNK1 regulates AKT signaling through direct interaction with PPP2R1A. (A) WNK1 and YFP expression in HEC1A cells transfected with YFP expressing control plasmid (cYFP) or YFP-tagged WNK1 expression construct (c4161). (B) Co-immunoprecipitation of WNK1 and PPP2R1A with YFP in HEC1A whole cell lysate, as indicated by western blotting. (C) Expression of PPP2R1A and PP2A subunits in HEC1A cells transfected with 24 nM siCTRL or siWNK1 for 72 hours. (D) Expression of PP2A subunits A, B and C, AKT and FOXO1 in HEC1A cells transfected with 72 nM siCTRL or siPPP2R1A for 72 hours. (E) Diagram illustrating the WNK1-PP2A-AKT-FOXO1 signaling axis.
Supplemental Information Titles and Legends

Figure S1. Generation of uterine WNK1 ablation mouse model. (A) The Wnk1\textsuperscript{ff} mice harbours 2 loxP sites (red block “L”) flanking the second exon of the Wnk1 gene, which were crossed to mice carrying Cre under the control of PGR promoter. Resultant PGR\textsuperscript{Cre/+}; Wnk1\textsuperscript{ff} mice would have exon2 excised from the genome. (B and C) Decreased Wnk1 gene and WNK1 protein expression were confirmed using qPCR (B) and western blotting (C), results shown are mean ± SD, * p < 0.05.

Figure S2. Delayed implantation in the Wnk1\textsuperscript{ff} mice was not due to aberrant ovarian-pituitary function. (A) Ovulation and fertilization was examined by inducing super-ovulation in the Wnk1\textsuperscript{ff} and Wnk1\textsuperscript{dd} mice, followed by mating to wild-type male mice. Number of 2-cell embryos were quantified by oviductal flushing and counting under brightfield microscope. (B and C) Serum progesterone (P4) and estradiol (E2) on GD 4.5 in Wnk1\textsuperscript{ff} and Wnk1\textsuperscript{dd} mice. Results shown are mean ± SD.

Figure S3. Principle component analysis of uterine transcriptome during receptivity from 4 Wnk1\textsuperscript{ff} mice (red) and 5 Wnk1\textsuperscript{dd} mice (blue).

Figure S4. Quantification and validation of the phosphokinase array. (A) The phosphokinase array was subjected to densitometrical quantification using ImageJ, and phosphorylation change which exceeded 1.5 fold are shown. (B) Phosphorylation and total levels of AKT, GSK-3α/β, and PRAS40 in the Wnk1\textsuperscript{ff} and Wnk1\textsuperscript{dd} uteri on PPD 4.5 were assayed by western blot, with each lane representing one mouse.
**Figure S5. mTOR is activated in the Wnk1\textsuperscript{dd} uteri, but does not regulate AKT activity.** (A) Rapamycin, an mTOR inhibitor was unable to reverse WNK1 ablation induced FOXO1 nuclear exclusion, indicating that AKT activity is maintained. (B) Rapamycin treatment did not rescue WNK1 ablation induced phosphorylation of AKT and FOXO1 in both HEC1A (left) and THESC (right) cells. (C) Expression and phosphorylation of FOXO1 and AKT in HEC1A cells transfected with siCTRL, siWNK1, simTOR, or both siWNKT and simTOR. Results showed no rescue of FOXO1 and AKT phosphorylation by mTOR knock-down.

**Figure S6. WNK1 immunoprecipitation confirmation.** HEC1A cells were subjected to immunoprecipitation (IP) using IgG (negative control) or WNK1 targeting antibody. The pulldown lysate showed enrichment of WNK1 in the WNK1 IP, but not in the IgG (black arrow indicates expected position of WNK1), as determined by western blotting.

**Fig. S7. Plasmid map of the WNK1 mammalian expression construct pcDNA-6.2-N-YFP-WNK1 (c4161).**

**Table S1. Serum progesterone levels in Wnk1\textsuperscript{ff} and Wnk1\textsuperscript{dd} mice on PPD 4.5 to confirm pseudopregnancy.**

**Table S2. List of differentially expressed genes in the Wnk1\textsuperscript{dd} uteri compared to Wnk1\textsuperscript{ff} uteri on PPD 4.5.**
Table S3. Functional annotation of altered transcriptome in Wnk1<sup>dd</sup> uteri at PPD 4.5 (DAVID).

Table S4. List of upstream regulators with altered activities in Wnk1<sup>dd</sup> uteri at PPD 4.5 (IPA).

Table S5. List of potential WNK1 interacting partners identified by WNK1 IP-MS.

Table S6. List of antibodies, sources, and experimental conditions used in this study.
Data Availability

All data generated or analyzed during this study are included in this published article or in the data repositories listed in References.

Supplemental tables and figures can be found at:

Gene expression data can be found at: