1 **AKT1-FOXO4 AXIS REGULATES HEMOCHORIAL PLACENTATION** 2 Keisuke Kozai^{1,*,§}, Ayelen Moreno-Irusta^{1,*}, Khursheed Iqbal¹, Mae-Lan Winchester^{2†}, 3 Regan L. Scott¹, Mikaela E. Simon¹, Masanaga Muto^{1‡}, Marc R. Parrish², and Michael 4 J. Soares^{1,2,3,¥} 5 6 7 ¹Institute for Reproduction and Perinatal Research, Department of Pathology & Laboratory Medicine, University of Kansas Medical Center, Kansas City, KS 8 9 ²Department of Obstetrics and Gynecology, University of Kansas Medical Center, 10 11 Kansas City, KS 12 ³Center for Perinatal Research, Children's Mercy Research Institute, Children's Mercy, 13 Kansas City, MO 14 15 *Contributed equally 16 [¥]Correspondence: msoares@kumc.edu 17 18 [§]Present address: Department of Obstetrics and Gynecology, University of Missouri-Kansas 19 City School of Medicine, Kansas City, MO 20 21 [†]Present address: Department of Obstetrics and Gynecology, University Hospital, Case 22Western Reserve University, Beachwood, OH 44122 23 24 25 ^{*}Present address: Department of Stem Cells and Human Disease Models, Research Center for Animal Life Science, Shiga University of Medical Science, Seta, Tsukinowa-cho, Otsu, 26 27 Shiga 520-2192, Japan 28 **Running title:AKT1 and uteroplacental development** 29 30 31 Keywords: AKT1, FOXO4, trophoblast, placenta, pregnancy, rat, genome editing 32

33 ABSTRACT

34 AKT1 is a serine/threonine kinase implicated in fetal, placental, and postnatal growth. In 35 this study, we investigated roles for AKT1 in placental development using a 36 genome-edited/loss-of-function rat model. Both heterozygous and homozygous Akt1 mutant 37 rats were viable and fertile. Disruption of AKT1 resulted in placental, fetal, and postnatal 38 growth restriction. Akt1 null placentas showed deficits in both junctional zone and labyrinth 39 zone size. Robust differences in the transcriptome of wild type versus Akt1 null junctional 40 zones were identified. Forkhead box O4 (Foxo4), which encodes a transcription factor and known AKT substrate, was abundantly expressed in the junctional zone. FOXO4 expression 41 42 was prominent in the junctional zone and invasive trophoblast cells of the rat placentation site 43 and enhanced following rat trophoblast stem cell differentiation. Foxo4 gene disruption using 44 genome-editing resulted in placentomegaly, including an enlarged junctional zone. AKT1 and 45 FOXO4 regulate the expression of many of the same transcripts expressed by trophoblast 46 cells; however, in opposite directions. In summary, we have identified AKT1 and FOXO4 as 47 part of a regulatory network controlling hemochorial placenta development. 48

50 **INTRODUCTION**

51	The placenta is an extraembryonic structure essential for normal fetal development
52	(Maltepe & Fisher, 2015; Burton et al., 2016). Placentas possess two main functions: i)
53	transformation of the maternal environment to support viviparity and ii) regulation of the
54	transfer of nutrients to the fetus (Gardner & Beddington, 1988; Soares et al., 2018; Knöfler
55	et al., 2019). These specialized functions are ascribed to trophoblast cells, which differentiate
56	along a multi-lineage pathway and are situated within specific compartments of the placenta
57	(Gardner & Beddington, 1988; Soares et al., 2018; Knöfler et al., 2019; Aplin & Jones,
58	2021). Placentas come in different shapes, sizes, and connectivity to the mother (Wooding &
59	Burton, 2008; Roberts et al., 2016). Placentation in some mammalian species is
60	characterized by trophoblast cells migrating into the maternal uterus where they modify the
61	vasculature facilitating maternal nutrient flow to the placenta (Pijnenborg et al., 1981;
62	Soares et al., 2018). This type of placenta is referred to as hemochorial (Wooding & Burton,
63	2008; Roberts et al., 2016). The human and rat possess hemochorial placentation where
64	invasive trophoblast cells migrate deep into the uterine parenchyma (Pijnenborg et al., 1981;
65	Soares et al., 2018). Regulation of deep hemochorial placentation is poorly understood. The
66	rat represents a useful animal model for investigating the regulation of deep hemochorial
67	placentation (Pijnenborg & Vercruysse, 2010; Soares et al., 2012; Shukla & Soares,
68	2022).
69	

The rat placenta can be divided into two main compartments: i) junctional zone; ii) labyrinth zone (**Ain** *et al.*, **2006**; **Soares** *et al.*, **2012**). The junctional zone compartment of the placenta is situated proximal to the uterine endometrium and is responsible for transforming the maternal environment, whereas the labyrinth zone is located between the junctional zone and fetus where it facilitates nutrient delivery to the fetus (**Knipp** *et al.*,

75	1999; Soares et al., 2012). Junctional zone-specific functions include the production of
76	peptide and steroid hormones that target maternal organs and the generation of invasive
77	trophoblast cells that migrate into and restructure the uterine parenchyma (Soares et al., 1996,
78	2012). The extravillous trophoblast cell column is a structure within the human placentation
79	site, which shares some of these same responsibilities (Soares et al., 2018; Knöfler et al.,
80	2019). The junctional zone and extravillous trophoblast cell column are pivotal to the
81	regulation of maternal adaptations to pregnancy, yet little is known about how they are
82	regulated.
83	
84	In this report, we focus on the phosphatidylinositol 3-kinase (PI3K)/AKT pathway and its
85	involvement in regulating junctional zone biology. AKT1 is one of three AKT
86	serine/threonine kinases and represents an integral component of signal transduction
87	pathways regulating cell proliferation, differentiation, migration, survival, and metabolism
88	(Manning & Toker, 2017; Cole et al., 2019). AKT1 has also been implicated in placentation
89	and trophoblast cell development through rodent mutagenesis experiments and investigations
90	with human trophoblast cells (Kamei et al., 2002; Yang et al., 2003; Qiu et al., 2004; Dash
91	et al., 2005; Kent et al., 2010, 2011, 2012; Plaks et al., 2011; Haslinger et al., 2013;
92	Sharma et al., 2016). Disruptions in AKT signaling have been connected to trophoblast cell
93	dysfunction leading to recurrent pregnancy loss, preeclampsia, and infertility (Pollheimer &
94	Knöfler, 2005; Ferretti et al., 2007; Fisher, 2015; Burton & Jauniaux, 2018). Herein we
95	show that AKT1 inactivation leads to placental and fetal growth restriction in the rat. AKT1
96	acts via phosphorylation of its target proteins leading to functional changes, including
97	activation or inhibition of the target protein function (Manning & Toker, 2017; Cole et al.,
98	2019). We identified forkhead box O4 (FOXO4), a transcription factor, as an AKT1 substrate
99	within the rat junctional zone and in rat trophoblast cells and demonstrated FOXO4

- 100 involvement in junctional zone development and the regulation of trophoblast cell
- 101 differentiation.
- 102

103 **RESULTS**

104 Generation of an Akt1 mutant rat model

105 We examined the role of AKT1 in regulating deep placentation in the rat using

106 CRISPR/Cas9 genome editing. A mutant rat strain possessing a 1,332 bp deletion within the

107 *Akt1* gene was generated (**Fig. 1A and B**). The deletion included part of Exon 4, the entire

region spanning Exon 5 through Exon 6, and part of Exon 7 and led to a frameshift and

109 premature stop codon (Fig. 1A and B). The deletion effectively removed the kinase domain

and regulatory regions of AKT1 (Fig. 1C). The *Akt1* mutation was successfully transmitted

111 through the germline. A rat colony possessing the *Akt1* mutation was established and

112 maintained via heterozygous x heterozygous breeding. Mating of heterozygotes produced the

113 predicted Mendelian ratio (Fig. 1D; Supplemental Table 1). Placental tissues possessing a

homozygous deletion within the *Akt1* locus ($Akt1^{-/-}$) were deficient in AKT1 protein and

exhibited prominent deficits in pan-AKT and phospho-AKT protein expression (Fig. 1E).

116 These findings support the successful disruption of the *Akt1* locus and are consistent with

117 AKT1 being the predominant AKT isoform within the placenta (Yang et al., 2003; Kent et

- 118 *al.*, **2011; Haslinger** *et al.*, **2013**).
- 119

120 AKT1 deficiency results in placental, fetal, and postnatal growth restriction

121 Disruption of the *Akt1* locus in the mouse disrupts placental, fetal, and postnatal growth

122 (Chen et al., 2001; Cho et al., 2001; Yang et al., 2003; Plaks et al., 2011; Kent et al., 2012).

123 We observed a similar phenotype in the rat. Gestation day (gd) 18.5 placental and fetal

weights and postnatal pup weights were significantly smaller in the $Aktl^{-/-}$ rat model when

125	compared to $Aktl^{+/+}$ rats (Fig. 2A-H). Junctional and labyrinth zone compartments of the
126	placenta were also significantly smaller in $AktI^{-/-}$ placentas (Fig. 2D, E, I).

127

128 **AKT1 regulates junctional zone and invasive trophoblast cell phenotypes**

Transcript profiles were determined for $Akt1^{+/+}$ and $Akt1^{-/-}$ gd 18.5 junctional zone tissues using RNA-sequencing (**RNA-seq**). The size and morphological phenotypes associated with inactivation of AKT1 were associated with distinct transcript profiles (**Fig. 3; Dataset 1**). Disruption of AKT1 resulted in upregulation of 254 transcripts and downregulation of 333

transcripts (**Dataset 1**). Pathway analysis included signatures for cell cycle, DNA replication,

cellular senescence, and PI3K-AKT signaling pathways (Fig. 3A). Transcripts encoding cell

135 cycle progression were consistently repressed in the $Aktl^{-/-}$ junctional zones (**Fig. 3A and B**).

136 In addition, we also observed prominent downregulation of a member of the expanded

137 prolactin (**PRL**) gene family, *Prl8a4*, and the upregulation of cellular communication

network factor 3 (*Ccn3*, also called *Nov*; **Fig. 3B**).

139

The junctional zone serves as the source of invasive trophoblast cells entering the uterus. 140 Consequently, we investigated the uterine-placental interface of $Akt I^{+/+}$ and $Akt I^{-/-}$ placentas 141 142 and monitored the surface area occupied by intrauterine invasive trophoblast cells and the expression of invasive trophoblast cell-specific transcripts. Akt1^{-/-} invasive trophoblast cells 143 exhibited decreased infiltration into the uterus (Supplemental Fig. 1A-C). We also observed 144 approximately a 50% decrease in the expression of cytokeratin transcripts, which are 145 146 typically expressed by invasive trophoblast cells within the uterine-placental interface 147 (Supplemental Fig. 1D). More prominent decreases in expression of invasive 148 trophoblast-cell specific transcripts were observed in uterine-placental interface tissue associated with $AktI^{-}$ placentas (Supplemental Fig. 1D). 149

150

- Collectively, the data indicate that AKT1 signaling has profound effects on development
 of the junctional zone and the invasive trophoblast cell lineage.
- 153
- 154 FOXO4 is a target of PI3K/AKT signaling
- 155 Forkhead box (FOX) transcription factors are known targets of PI3K/AKT signaling and
- have key roles in regulating developmental processes (Lam *et al.*, 2013; Schmitt-Ney, 2020;
- 157 Herman et al., 2021). We interrogated RNA-seq datasets from wild type gd 18.5 junctional
- zone for FOX family transcription factors. Transcripts for several FOX transcription factors
- 159 were detected (transcripts per million, TPM value ≥ 1.0) (**Fig. 4A**). *Foxo4* transcripts were
- striking in their abundance relative to all other FOX family members. AKT1 did not
- significantly affect expression levels for any of the FOX family transcripts (**Dataset 1**).
- 162 Foxo4 transcripts were specifically localized to the junctional zone and a subset of invasive
- 163 trophoblast cells localized to uterine spiral arteries, termed endovascular invasive trophoblast
- 164 cells (**Fig. 4B and C**). Total and phosphorylated FOXO4 protein were significantly
- diminished in *Akt1* null junctional zones (**Fig. 4D**). We next explored FOXO4 in
- differentiated rat trophoblast stem (TS) cells, a model for trophoblast cell lineages found in
- 167 the junctional zone (Asanoma et al., 2011). Foxo4 transcript and total and phosphorylated
- 168 FOXO4 protein showed striking increases in abundance following TS cell differentiation (Fig.
- 169 **4E-F**). As previously demonstrated, AKT activity increased following trophoblast cell
- differentiation (Kamei et al., 2002; Kent et al., 2010, 2011; Fig. 4G). AKT activation was
- required for optimal FOXO4 phosphorylation (Fig. 4G). Thus, we established a link between
- 172 PI3K/AKT signaling and FOXO4 in trophoblast cell lineage development.
- 173

174 Generation of a *Foxo4* mutant rat model

175	We examined the role of FOXO4 in regulating placentation in the rat using CRISPR/Cas9
176	genome editing. The Foxo4 gene consists of four exons and resides on the X chromosome
177	(Liu et al., 2020). A mutant rat strain possessing a 3,096 bp deletion within the Foxo4 gene
178	was generated (Fig. 5A and B). The deletion included the 3' part of Exon 2 and the 5' part of
179	Exon 3 and led to a frameshift and a premature stop codon (Fig. 5A and B). The deletion
180	effectively disrupted the conserved forkhead DNA binding domain and removed nuclear
181	localization, nuclear export, and transactivation domains of FOXO4 (Fig. 5C). The Foxo4
182	mutation was successfully transmitted through the germline (Fig. 5D; Supplemental Table
183	2). A rat colony possessing the <i>Foxo4</i> mutation was established and maintained via
184	hemizygous male x wild type female breeding, which produced the predicted Mendelian ratio
185	(Supplemental Table 2). Junctional zone tissues possessing a maternally inherited Foxo4
186	mutation ($Foxo4^{Xm-}$) were deficient in FOXO4 protein (Fig. 5E). The results are consistent
187	with paternal silencing of X chromosome-linked genes expressed in extraembryonic tissues
188	(Takagi & Sasaki, 1975; West et al., 1978; Hemberger, 2002). FOXO4 was successfully
189	disrupted in the rat.
190	
191	FOXO4 deficiency results in placentomegaly and a modified junctional zone phenotype
192	Placentation site phenotypes of mice possessing mutations at the Foxo4 locus have not
193	been described (Liu et al., 2020; Hosaka et al., 2004). Rats possessing a maternally inherited
194	mutant <i>Foxo4</i> allele ($Foxo4^{Xm}$) exhibited placentomegaly and decreased placental efficiency
195	(fetal/placental weight ratio) when examined on gd 18.5 (Fig. 6A-C) and gd 20.5
196	(Supplemental Fig. 2A-C). In contrast, a paternally inherited mutant <i>Foxo4</i> allele did not
197	significantly affect placenta or fetal weights (Supplemental Fig. 2D and E). FOXO4
198	deficiency associated placentomegaly included significantly larger junctional and labyrinth
199	zones (Fig. 6D-G, Supplemental Fig. 2F-H) but did not affect the intrauterine invasive

200	trophoblast cell lineage (Supplemental Fig. 3). Transcript profiles were determined for wild
201	type ($Foxo4^{Xm+}$) and $Foxo4^{Xm-}$ gd 18.5 junctional zone tissues using RNA-seq (Fig. 6H).
202	Disruption of FOXO4 resulted in upregulation of 369 transcripts and downregulation of 845
203	transcripts (Dataset 2). Pathway analysis included signatures for PI3K-AKT signaling, cell
204	cycle, DNA replication, extracellular matrix receptor interaction, and complement and
205	coagulation pathways (Fig. 6H). Junctional adhesion molecule-like (JAML), lipoprotein(a)
206	like 2 (LPAL2), erb-b2 receptor tyrosine kinase 3 (ERBB3), and growth factor receptor
207	bound protein 7 (GRB7) were each conspicuous in their prominent downregulation in
208	FOXO4 deficient junctional zone tissue (Fig. 6I). JAML contributes to epithelial barrier
209	function, modulates immune cell trafficking, and angiogenesis (Kummer & Ebnet, 2018),
210	whereas LPAL2 is a long noncoding RNA contributing to inflammatory and oxidative stress
211	responses (Han et al., 2018). JAML and LPAL2 have not previously been linked to
212	trophoblast or placental biology. ERBB3 is a receptor for neuregulin 1 and promotes
213	trophoblast cell survival (Fock et al., 2015) and GRB7 is an adaptor protein participating in
214	signal transduction activated through ERBB3 (Fiddes et al., 1998). Interestingly, numerous
215	junctional zone transcripts regulated by FOXO4 were reciprocally regulated by AKT1 (88%
216	of transcripts upregulated in Akt1 null were downregulated in Foxo4 mutant tissues; 38% of
217	transcripts downregulated in Akt1 null were upregulated in Foxo4 mutant tissues; Fig. 7). The
218	reciprocal relationship between AKT1 and FOXO4 is evident at structural and molecular
219	levels.
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FOXO4 contributes to the regulation of the trophoblast cell lineage

We next modeled junctional zone cell biology using rat TS cells. The consequences of FOXO4 disruption in differentiating rat TS cells were examined. FOXO4 expression was inhibited via ectopic expression of short hairpin RNAs specific to *Foxo4* (**Fig. 8A and B**).

225	Although, a morphologic phenotype was not evident, prominent differences in the
226	transcriptomes of TS cells expressing control versus Foxo4 shRNAs were observed (Fig. 8C).
227	Disruption of FOXO4 resulted in upregulation of 260 transcripts and downregulation of 443
228	transcripts (Dataset 3). Pathway analysis included signatures for PI3K-AKT signaling,
229	longevity regulating, calcium signaling, and glutathione metabolism pathways (Fig. 8C).
230	Among the dysregulated transcripts was an upregulation of matrix metallopeptidase 12, a
231	known constituent of endovascular invasive trophoblast cells (Harris et al., 2010;
232	Chakraborty et al., 2016) and downregulation of trophoblast specific protein alpha, a
233	transcript characteristic of spongiotrophoblast cells within the junctional zone (Iwatsuki et
234	al., 2000). FOXO4 is a known regulator of responses to oxidative stress (Liu et al., 2020).
235	Several transcripts associated with inflammatory and cellular stress responses, including
236	thioredoxin interacting protein, glutathoione S-transferase mu 1, arachidonate 5-lipoxygenase
237	activating protein, interferon kappa, nuclear protein 1, and Lpal2, were prominently
238	downregulated (Sies & Cadenas, 1985; LaFleur et al., 2001; Mashima & Okuyama, 2015;
239	Han et al., 2018; Huang et al., 2021; Nirgude & Choudhary, 2021; Qayyum et al., 2021;
240	Satapathy & Wilson, 2021).
241	
242	Key findings

243 Collectively, the results indicate that AKT1 drives placental growth, including regulation

of deep intrauterine trophoblast cell invasion. These actions are accomplished, at least in part,

through modulation of FOXO4, which acts to restrain placental growth and coordinate

responses to physiological stressors.

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250 **DISCUSSION**

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251	The rat and human possess a type of hemochorial placentation where specialized
252	trophoblast cells penetrate deep into the uterus and transform the uterine parenchyma,
253	including the vasculature (Pijnenborg et al., 1981; Pijnenborg & Vercruysse, 2010; Soares
254	et al., 2012). Central to deep placentation is the source of invasive trophoblast cells, which in
255	the rat is a compartment within the placenta referred as the junctional zone and, in the human,
256	the extravillous trophoblast cell column (Soares et al., 2012, 2018; Knöfler et al., 2019). In
257	addition to an intrauterine role, the cellular constituents of these placental compartments
258	produce hormones directed to maternal tissues with actions that ensure in utero survival and
259	promotion of fetal growth (Soares et al., 1996; John, 2017). In this report, AKT1 and
260	FOXO4 were identified as regulators of rat junctional zone development. In vivo disruption
261	of AKT1 and FOXO4 led to opposite effects on junctional zone development. AKT1
262	deficiency resulted in growth restriction of the junctional zone, phenotypic alteration of the
263	invasive trophoblast cell lineage, as well as compromised fetal and postnatal growth, and
264	AKT1 was capable of phosphorylating FOXO4 in rat trophoblast cells. Deficiency of FOXO4
265	resulted in an expanded junctional zone. Deficits in AKT1 or FOXO4 also impacted
266	transcriptomic profiles of the junctional zone. The findings indicate that AKT1 and FOXO4
267	are part of a gene regulatory network controlling hemochorial placentation.
268	
269	AKT1 signaling influenced placental development Akt1 null mutations in the mouse and

AKT1 signaling influenced placental development. *Akt1* null mutations in the mouse and rat yield similar phenotypes characterized by placental, fetal, and postnatal growth restriction (Chen *et al.*, 2001; Cho *et al.*, 2001; Yang *et al.*, 2003; Plaks *et al.*, 2011; Kent et al., 2012). Smaller junctional zones accompanying AKT1 deficiency were associated with a downregulation of transcripts encoding proteins driving cell proliferation. These results imply

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that size differences of junctional zone compartments in the wild type versus Akt1 nulls were

275related to, at least in part, diminished trophoblast cell proliferation in Akt1 junctional zone 276 tissues. The data also fit well with known actions of AKT signaling promoting cell 277 proliferation in a wide range of cell types (Manning & Cantley, 2007; Manning & Toker, 278 2017; Cole et al., 2019). AKT1 disruption also altered differentiated junctional zone 279 trophoblast cell phenotypes. As cellular constituents of the junctional zone differentiate, they 280 acquire the capacity to express several members of the expanded prolactin (**PRL**) family of 281 hormones/cytokines (Soares, 2004; Alam et al., 2006; Soares et al., 2007). The expression 282 of PRL8A4, a member of the expanded PRL family, was dramatically downregulated in Akt1 283 null junctional zones. PRL8A4 is an orphan ligand with little known of its significance to the 284 biology of pregnancy other than as a signature feature of the differentiated junctional zone 285 phenotype (Iwatsuki et al., 1998; Soares et al., 2007). AKT signaling has previously been 286 implicated in the regulation of the differentiation of rodent and human trophoblast cells 287 (Kamei et al., 2003; Kent et al., 2010, 2011; Haslinger et al., 2013). Disruption of Akt1 also 288 interfered with invasive trophoblast cell development. Trophoblast cell infiltration into the 289 uterine-placental interface was diminished in Akt1 nulls as was the expression of transcripts 290 indicative of the invasive trophoblast cell lineage. Involvement of AKT signaling has also 291 been implicated in the development of the human extravillous trophoblast cell lineage 292 (Pollheimer & Knöfler, 2005; Haslinger et al., 2013; Morey et al., 2021). AKT signaling 293 could affect invasive trophoblast cell development through its actions on their origin in the 294 junctional zone and EVT cell column or alternatively, their maturation as they invade into the 295 uterus. Finally, the impact of AKT signaling in invasive trophoblast cell development may be 296 more profound than observed with AKT1 deficiency due to compensatory activities of AKT2 297 and AKT3 (Kent et al., 2011; Haslinger et al., 2013).

298

299	AKT1 regulates cellular function through its actions as a serine/threonine kinase and thus,
300	phosphorylation of its substrates (Manning & Cantley, 2007; Manning & Toker, 2017;
301	Cole et al., 2019). The forkhead box (FOX) family of transcription factors are
302	well-established targets of AKT action (Lam et al., 2013; Schmitt-Ney, 2020; Herman et al.,
303	2021). Among FOX family members, FOXO4 expression was uniquely elevated in the rat
304	junctional zone. FOXO4 phosphorylation state in trophoblast cells was affected by AKT
305	signaling. AKT-mediated phosphorylation of FOXO4 leads to FOXO4 exit from the nucleus
306	and inactivation (Schmitt-Ney, 2020; Herman et al., 2021), which might suggest that an
307	AKT1 deficiency would result in the stabilization of FOXO4 protein in the junctional zone.
308	Instead, AKT1 deficiency led to depletion of junctional zone total and phosphorylated
309	FOXO4 proteins. Consequently, the observed Akt1 null placental phenotype was associated
310	with the depletion of both AKT1 and FOXO4 proteins. In addition to regulation by AKT
311	signaling, FOXO4 activity/stability is stimulated by Jun kinase and monoubiquitylation
312	(Essers et al., 2004; van der Horst et al., 2006; Brenkman et al., 2008; Liu et al., 2020),
313	while inhibited by acetylation and polyubiquitylation (Fukuoka et al., 2003; Huang &
314	Tindall, 2011; Liu et al., 2020). Whether AKT1 indirectly affects FOXO4 protein via
315	impacting these other FOXO4 regulators remains to be determined.
316	
317	FOXO4 is a transcription factor implicated in the regulation of the cell cycle, apoptosis,
318	responses to oxidative stress, and a range of disease processes (Liu et al., 2020). The original

319 characterization of the *Foxo4* null mouse concluded that FOXO4 did not have a singular role

- in the pathophysiology of the mouse (Hosaka *et al.*, 2004). The absence of a reported
- 321 phenotype for the *Foxo4* null mouse model was attributed to the compensatory actions of
- 322 other members of the FOXO family, including FOXO1, FOXO3, and possibly FOXO6

323 (Hosaka *et al.*, 2004; Liu *et al.*, 2020). We describe a prominent placental phenotype for the

324 Foxo4 null rat model. The placental anomalies associated with FOXO4 deficiency were 325 compatible with the production of viable offspring. The absence of a fertility defect in the 326 Foxo4 null mouse model likely precluded a closer examination of placentation (Hosaka et al., 327 **2004**). However, it is also possible that elements of FOXO4 action are species specific. 328 FOXO4 is prominently expressed in the junctional zone and to a lesser extent in invasive 329 trophoblast cells. Disruption of FOXO4 led to an expansion of both junctional and labyrinth 330 zone placental compartments, which probably reflects cell autonomous and non-cell 331 autonomous actions, respectively. A striking reciprocal pattern of AKT1 versus FOXO4 gene 332 regulation within the junctional zone was demonstrated and included differentially regulated 333 transcripts encoding proteins involved in the regulation of cell proliferation and cell death. 334 We surmise that AKT1 promotes junctional zone growth via stimulating the expression of 335 transcripts connected to cell cycle progression and inhibited transcripts connected to cell 336 death, whereas the converse is true for FOXO4. These biological roles are consistent with the 337 known actions of AKT1 and FOXO4 in other cell systems (Manning & Toker, 2017; Liu et 338 *al.*, 2020; Herman *et al.*, 2021). A transcriptional regulatory network involving FOXO4 has 339 also been identified in human extravillous trophoblast cells (Morey et al., 2021). Importantly, 340 FOXO4 also regulates trophoblast cell responses to oxidative stress and is thus, positioned to 341 contribute to placental adaptations to a compromised maternal environment and in disease 342 states affecting placentation.

343

344 MATERIALS AND METHODS

345 Animals

Holtzman Sprague-Dawley rats were maintained in an environmentally controlled facility with lights on from 0600 to 2000 h with food and water available ad libitum. Time-mated pregnancies were established by co-housing adult female rats (8-10 weeks of age) with adult

349	male rats (>10 weeks of age). Detection of sperm or a seminal plug in the vagina was
350	designated gd 0.5. Pseudopregnant females were generated by co-housing adult female rats
351	(8-10 weeks of age) with adult male vasectomized males (>10 weeks of age). The detection
352	of seminal plugs was designated pseudopregnancy day 0.5. Four to five-week-old donor rats
353	were superovulated by intraperitoneal injection of pregnant mare serum gonadotropin (30
354	units, G4877, Sigma-Aldrich, St. Louis, MO), followed by an intraperitoneal injection of
355	human chorionic gonadotropin (30 units, C1063, Sigma-Aldrich) ~46 h later, and
356	immediately mated with adult males. Zygotes were flushed from oviducts the next morning
357	(gd 0.5). The University of Kansas Medical Center (KUMC) Animal Care and Use
358	Committee approved all protocols involving the use of rats.
359	
360	Tissue collection and analysis
361	Rats were euthanized by CO ₂ asphyxiation at designated days of gestation. Uterine
362	segments containing placentation sites were frozen in dry ice-cooled heptane and stored at
363	-80°C until processed for histological analyses. Alternatively, placentation sites were
364	dissected into placentas, the adjacent uterine-placental interface tissue (also referred to as the
365	metrial gland), and fetuses as previously described (Ain et al., 2006). Placentas were weighed
366	and dissected into junctional zone and labyrinth zone compartments (Ain et al., 2006).
367	Placental compartments and uterine-placental interfaces were frozen in liquid nitrogen and
368	stored at -80°C until used for biochemical analyses. Fetuses were weighed, genotyped, and
369	sex determined by polymerase chain reaction (PCR) (Dhakal & Soares, 2017).
370	
371	Generation of Akt1 and Foxo4 mutant rat models
372	Mutations at Akt1 and Foxo4 loci were generated using CRISPR/Cas9 genome editing
373	(Kaneko, 2017; Iqbal et al., 2021). Guide RNAs targeting Exon 4 (target sequence:

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374	GCCGTTTGAGTCCATCAGCC; nucleotides 356-375) and Exon 7 (target sequence:
375	TTGTCATGGAGTACGCCAAT; nucleotides 712-731) of the Akt1 gene (NM_033230.3) or
376	targeting Exon 2 (target sequence: CCAGATATACGAATGGATGGTCC; nucleotides
377	517-539) and Exon 3 (target sequence: GTTCATCAAGGTACATAACGAGG; nucleotides
378	631-653) of the Foxo4 gene (NM_001106943.1) were electroporated into single-cell rat
379	embryos using the NEPA21 electroporator (Nepa Gene Co Ltd, Ichikawa City, Japan).
380	Electroporated embryos were transferred to oviducts of day 0.5 pseudopregnant rats. Initially,
381	offspring were screened for Akt1 or Foxo4 mutations from genomic DNA from tail-tip
382	biopsies using the REDExtract-N-Amp TM Tissue PCR kit (XNAT, Millipore Sigma,
383	Burlington, MA). PCR was performed on the purified DNA samples using primers flanking
384	the guide RNA sites, and products resolved by agarose gel electrophoresis and ethidium
385	bromide staining. Genomic DNA containing potential mutations was amplified by PCR, gel
386	purified, and precise boundaries of deletions determined by DNA sequencing (Genewiz Inc.,
387	South Plainfield, NJ). Founders with Akt1 or Foxo4 mutations were backcrossed to wild type
388	rats to demonstrate germline transmission. Routine genotyping was performed by PCR on
389	genomic DNA with specific sets of primers (Supplemental Table 3).

390

391 Western blot analysis

392 Tissue lysates were prepared with radioimmunoprecipitation assay lysis buffer system

393 (sc-24948A, Santa Cruz Biotechnology, Santa Cruz, CA). Protein concentrations were

determined using the *DC*TM Protein Assay Kit (5000112, Bio-Rad Laboratories, Hercules,

395 CA). Proteins (20 µg/lane) were separated by SDS-PAGE. Separated proteins were

electrophoretically transferred to polyvinylidene difluoride membranes (10600023, GE

Healthcare, Milwaukee, WI) for 1 h at 100 V on ice. Membranes were subsequently blocked

with 5% milk or 5% bovine serum albumin for 1 h at room temperature and probed separately

399	with specific primary antibodies to AKT1 (1:1,000 dilution, 75692, Cell Signaling
400	Technology, Danvers, MA), pan-AKT (1:1,000 dilution, 4691, Cell Signaling Technology),
401	phospho-Ser ⁴⁷³ AKT (1:2,000 dilution, 4060, Cell Signaling Technology), FOXO4 (1:2,000
402	dilution, 21535-1-AP, Proteintech, Rosemont, IL), phospho-Ser ²⁶² FOXO4 (1:3,000 dilution,
403	ab126594, Abcam, Cambridge, MA), and glyceraldehyde 3-phosphate dehydrogenase
404	(GAPDH, 1:5,000 dilution, ab8245, Abcam) in Tris-buffered saline with Tween 20 (TBST)
405	overnight at $4\square$. After primary antibody incubation, the membranes were washed in TBST
406	three times for ten min each at room temperature. After washing, the membranes were
407	incubated with anti-rabbit or anti-mouse immunoglobulin G (IgG) conjugated to horseradish
408	peroxidase [HRP, 1:5,000 dilution or 1:20,000 dilution (phospho-Ser ²⁶² FOXO4), 7074S and
409	7076S, Cell Signaling Technology] in TBST for 1 h at room temperature, washed in TBST
410	three times for ten min each at room temperature, immersed in Immobilon Crescendo
411	Western HRP Substrate (WBLUR0500, Sigma-Aldrich), and luminescence detected using
412	Radiomat LS film (Agfa Healthcare, Mortsel, Belgium).
413	
414	Transcript analysis
415	Total RNA was extracted from tissues using TRI Reagent Solution (AM9738,
416	Thermo-Fisher, Waltham, MA) according to the manufacturer's instructions. Total RNA (1

- 417 µg) was reverse transcribed using a High-Capacity cDNA Reverse Transcription Kit
- 418 (4368813, Thermo-Fisher). Complementary DNAs were diluted 1:10 and subjected to reverse
- 419 transcription-quantitative PCR (**RT-qPCR**) using PowerUp SYBR Green Master Mix
- 420 (A25742, Thermo-Fisher) and primers provided in Supplemental Table 4. QuantStudio 5
- 421 Flex Real-Time PCR System (Applied Biosystems, Foster City, CA) was used for
- 422 amplification and fluorescence detection. PCR was performed under the following
- 423 conditions: $95\square$ for 10 min, followed by 40 cycles of $95\square$ for 15 sec and $60\square$ for 1 min.

424 Relative mRNA expression was calculated using the delta-delta Ct method. *Gapdh* was used
425 as a reference transcript.

426

427 **RNA-seq analysis**

428 Transcript profiles were generated from wild type and $Akt I^{-/-}$, and $Foxo 4^{Xm-}$ junctional

429 zone tissues, and rat differentiated TS cells expressing control or *Foxo4* shRNAs.

430 Complementary DNA libraries from total RNA samples were prepared with Illumina TruSeq

431 RNA preparation kits according to the manufacturer's instructions (Illumina, San Diego, CA).

432 RNA integrity was assessed using an Agilent 2100 Bioanalyzer (Santa Clara, CA). Barcoded

433 cDNA libraries were multiplexed onto a TruSeq paired-end flow cell and sequenced (100-bp

434 paired-end reads) with a TruSeq 200-cycle SBS kit (Illumina). Samples were run on an

435 Illumina NovaSeq 6000 sequencer at the KUMC Genome Sequencing Facility. Reads from

436 *.fastq files were mapped to the rat reference genome (Ensembl Rnor_5.0.78) using CLC

437 Genomics Workbench 12.0 (Qiagen, Germantown, MD). Transcript abundance was expressed

438 as transcript per million mapped reads (TPM) and a *P* value of 0.05 was used as a cutoff for

439 significant differential expression. Statistical significance was calculated by empirical

440 analysis of digital gene expression followed by Bonferroni's correction. Pathway analysis

441 was performed using Datbase for Annotation, Visualization, and Integrated Discovery

- 442 (DAVID; Huang *et al.*, 2009).
- 443

444 Immunohistochemistry

445 Placentation sites were embedded in optimum cutting temperature (**OCT**) compound and

sectioned at 10 μ m thickness. Sections were fixed in 4% paraformaldehyde, washed in

447 phosphate buffered saline (pH 7.4) three times for five min each, blocked with 10% normal

448 goat serum (50062Z, Thermo-Fisher), and incubated overnight with primary antibodies: pan

463	In situ hybridization
462	
461	described (Nteeba et al., 2020).
460	within the uterine-placental interface was quantified using ImageJ software, as previously
459	camera (Roper). The area occupied by cytokeratin-positive cells (invasive trophoblast cells)
458	Nikon 80i upright microscope (Nikon) with a Photometrics CoolSNAP-ES monochrome
457	Birmingham, AL) and examined microscopically. Fluorescence images were captured on a
456	then mounted with Fluoromount-G mounting media (0100-01, Southern Biotech,
455	3-amino-9-ethylcarbazole (AEC) substrate kit (SK-4200, Vector Laboratories). Sections were
454	dilution, A9044, Sigma-Aldrich) for 3 h at room temperature and color development with a
453	dilution, A11011, Thermo-Fisher) or rabbit anti-mouse IgG conjugated to HRP (1:500
452	corresponding secondary antibodies: Alexa 568-conjugated goat anti-rabbit IgG (1:500
451	After washing with phosphate buffered saline (pH 7.4), sections were incubated with
450	(1:300 dilution, sc-6260, Santa Cruz Biotechnology) to distinguish placental compartments.
449	cytokeratin (1:300 dilution, F3418, Sigma-Aldrich) to identify trophoblast cells and vimentin

Distributions of transcripts for *Foxo4* and *Prl7b1* were determined on cryosections of rat
placentation sites. RNAscope Multiplex Fluorescent Reagent Kit version 2 (Advanced Cell
Diagnostics, Newark, CA) was used for in situ hybridization analysis. Probes were prepared
to detect *Foxo4* (NM_001106943.1, 1038981-C1, target region: 750-1651) and *Prl7b1*(NM_153738.1, 860181-C2, target region: 28-900). Fluorescence images were captured on a
Nikon 80i upright microscope (Nikon) with a Photometrics CoolSNAP-ES monochrome
camera (Roper).

471

472 Rat TS cell culture

473 Blastocyst-derived rat TS cells (Asanoma *et al.*, 2011) were cultured in Rat TS Cell

474	Medium [RPMI 1640 medium (11875093, Thermo-Fisher), 20% (vol/vol) fetal bovine serum
475	(FBS, Thermo-Fisher), 100 μ M 2-mercaptoethanol (M7522, Sigma-Aldrich), 1 mM sodium
476	pyruvate (11360-070, Thermo-Fisher), 100 μ M penicillin and 50 U/mL streptomycin
477	(15140122, Thermo-Fisher)] supplemented with 70% rat embryonic fibroblast
478	(REF)-conditioned medium prepared as described previously (Asanoma et al., 2011), 25
479	ng/ml fibroblast growth factor 4 (FGF4; 100-31, Peprotech), and 1 μ g/mL heparin (H3149,
480	Sigma-Aldrich). For induction of differentiation, rat TS cells were cultured for 15 days in rat
481	TS medium containing 1% (vol/vol) FBS without FGF4, heparin, and REF-conditioned
482	medium. In some experiments, rat TS cells were exposed to a phosphatidylinositol 3-kinase
483	(PI3K) inhibitor (LY294002, 10 μ M, 9901, Cell Signaling Technology).
484	
485	Lentivirus construction and production
486	Lentivirus construction and production were described previously (Muto et al., 2021;
486 487	Lentivirus construction and production were described previously (Muto <i>et al.</i> , 2021; Varberg et al. , 2021). Briefly, the lentivirus encoding the shRNA targeting <i>Foxo4</i>
487	Varberg et al., 2021). Briefly, the lentivirus encoding the shRNA targeting <i>Foxo4</i>
487 488	Varberg et al., 2021). Briefly, the lentivirus encoding the shRNA targeting <i>Foxo4</i> (Supplemental Table 5) was constructed using a pLKO.1 vector. shRNA oligo sequences
487 488 489	Varberg et al., 2021). Briefly, the lentivirus encoding the shRNA targeting <i>Foxo4</i> (Supplemental Table 5) was constructed using a pLKO.1 vector. shRNA oligo sequences used in the analyses are provided in Supplemental Table 8. Lentiviral packaging vectors were
487 488 489 490	Varberg et al., 2021). Briefly, the lentivirus encoding the shRNA targeting <i>Foxo4</i> (Supplemental Table 5) was constructed using a pLKO.1 vector. shRNA oligo sequences used in the analyses are provided in Supplemental Table 8. Lentiviral packaging vectors were obtained from Addgene and included pMDLg/pRRE (plasmid 12251), pRSV-Rev (plasmid
487 488 489 490 491	Varberg et al., 2021). Briefly, the lentivirus encoding the shRNA targeting <i>Foxo4</i> (Supplemental Table 5) was constructed using a pLKO.1 vector. shRNA oligo sequences used in the analyses are provided in Supplemental Table 8. Lentiviral packaging vectors were obtained from Addgene and included pMDLg/pRRE (plasmid 12251), pRSV-Rev (plasmid 12253), pMD2.G (plasmid 12259). Lentiviral particles were produced using Attractene
487 488 489 490 491 492	Varberg et al., 2021). Briefly, the lentivirus encoding the shRNA targeting <i>Foxo4</i> (Supplemental Table 5) was constructed using a pLKO.1 vector. shRNA oligo sequences used in the analyses are provided in Supplemental Table 8. Lentiviral packaging vectors were obtained from Addgene and included pMDLg/pRRE (plasmid 12251), pRSV-Rev (plasmid 12253), pMD2.G (plasmid 12259). Lentiviral particles were produced using Attractene
487 488 489 490 491 492 493	Varberg et al., 2021). Briefly, the lentivirus encoding the shRNA targeting <i>Foxo4</i> (Supplemental Table 5) was constructed using a pLKO.1 vector. shRNA oligo sequences used in the analyses are provided in Supplemental Table 8. Lentiviral packaging vectors were obtained from Addgene and included pMDLg/pRRE (plasmid 12251), pRSV-Rev (plasmid 12253), pMD2.G (plasmid 12259). Lentiviral particles were produced using Attractene (301005, Qiagen) in human embryonic kidney (HEK) 293FT (Thermo-Fisher) cells.
487 488 489 490 491 492 493 494	Varberg et al., 2021). Briefly, the lentivirus encoding the shRNA targeting <i>Foxo4</i> (Supplemental Table 5) was constructed using a pLKO.1 vector. shRNA oligo sequences used in the analyses are provided in Supplemental Table 8. Lentiviral packaging vectors were obtained from Addgene and included pMDLg/pRRE (plasmid 12251), pRSV-Rev (plasmid 12253), pMD2.G (plasmid 12259). Lentiviral particles were produced using Attractene (301005, Qiagen) in human embryonic kidney (HEK) 293FT (Thermo-Fisher) cells. Lentiviral transduction

499 Statistical analysis

- 500 Student's *t*-test, Welch's *t*-test, Dunnett's test, or Steel test were performed, where
- ⁵⁰¹ appropriate, to evaluate the significance of the experimental manipulations. Results were
- 502 deemed statistically significant when P < 0.05.
- 503

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- 511

512 AUTHOR CONTRIBUTIONS

- 513 K.K., A.M.-I., and M.J.S. conceived and designed the research; K.K., A.M.-I., K.I.,
- 514 M.-L.W., R.L.S., M.E.S., M.M. performed experiments; K.K., A.M.-I., K.I., M.R.P., and
- 515 M.J.S. analyzed the data and interpreted results of experiments; K.K., A.M.-I., and M.J.S.
- 516 prepared figures and manuscript; All authors read, contributed to editing, and approved the
- 517 final version of manuscript.
- 518

519 CONFLICT OF INTEREST

- 520 There is no conflict of interest that could be perceived as prejudicing the impartiality of 521 the research reported.
- 522

523 DATA AVAILABILITY

524	RNA-seq datasets are available at the Gene Expression Omnibus (GEO) database,
525	https://www.ncbi.nlm.nih.gov/geo/ (accession number GSE205831). All data generated and
526	analyzed in this study are included in the published article and supporting files. Resources
527	generated from the research are available from the corresponding author upon reasonable
528	request. AKT1 and FOXO4 mutant rat models are available through the Rat Resource and
529	Research Center (Columbia, MO).
530	
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792	FIGURE LEGENDS
793	Figure 1. In vivo genome editing of the rat Akt1 locus. A) Schematic representation of the
794	rat <i>Akt1</i> gene (<i>Akt1</i> ^{+/+}) and guide RNA target sites within Exons 4 and 7 (NM_033230.3). Red
795	bars beneath Exons 4 and 7 correspond to the 5' and 3' guide RNAs used in the genome
796	editing. B) The mutant <i>Akt1</i> allele (<i>Akt1^{-/-}</i>) possesses a 1,332 bp deletion. Parts of Exons 4
797	and 7 and all of Exons 5 and 6 are deleted, leading to a frameshift and premature Stop codon

in Exon 7. C) Amino acid sequences for $AKT1^{+/+}$ and $AKT1^{-/-}$. The red sequence corresponds

807	Figure 2. <i>Akt1^{-/-}</i> placentas and fetuses are growth restricted, and <i>Akt1^{-/-}</i> rats exhibit
806	
805	used as a loading control.
804	phospho (p)-AKT (Ser ⁴⁷³) protein in $Akt1^{+/+}$ and $Akt1^{-/-}$ placentas at gd 18.5. GAPDH was
803	mutant (-/-) genotypes were detected by PCR. E) Western blot analysis of AKT1, pan-AKT,
802	to generate homozygous mutants. Wild type (+/+), heterozygous (+/-), and homozygous
801	Offspring were backcrossed to wild type rats, and heterozygous mutant rats were intercrossed
800	correspond to the pleckstrin homology (PH), kinase, and regulatory domains, respectively. D)
799	to the frameshift in Exon 7. The blue, red, and green highlighted amino acid sequence regions

postnatal growth restriction. Placentas (A) and fetuses (B) were dissected from $Akt1^{+/-}$

809 intercrosses at gd 18.5 and weighed; C, fetus/placenta ratio. Placentas were then separated

810 into junctional zone (JZ; D) and labyrinth zone (LZ;E) compartments, and weighed; F, JZ/JZ

ratio. Graphs represent means \pm SEM. $Akt l^{+/+}$, n = 22 from 6 dams; $Akt l^{-/-}$, n = 17 from 6

dams. Asterisks denote statistical differences (**P < 0.01; ***P < 0.001) as determined by

Student's or Welch's *t*-test. Body weights of $Akt1^{+/+}$ and $Akt1^{-/-}$ pups were measured from two

to eight weeks after birth: males (G) and females (H). Graphs represent means \pm SEM. n =

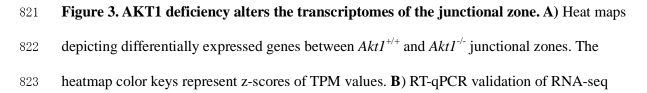
815 13-23/group. Asterisks denote statistical differences (*P < 0.05; ***P < 0.001) as determined

by Student's or Welch's *t*-test. I) Vimentin immunostaining of gd 18.5 $Akt 1^{+/+}$ and $Akt 1^{-/-}$

817 placentation sites. The junctional zone (JZ) is negative for vimentin immunostaining,

818 whereas the uterine-placental interface (UPI) and labyrinth zone (LZ) stain positive for

vimentin. Scale bars=1000 μm



824	results (n=6/group).	Graphs represent	means \pm SEM.	Asterisks der	note statistical	difference

- 825 (**P*<0.05; ***P*<0.01; ****P*<0.001) as determined by Student's or Welch's *t*-test.
- 826

827	Figure 4. FOXO4 is a target of PI3K/AKT signaling. A) Expressions of transcripts for
828	several FOX transcription factors in the junctional zone. \mathbf{B}) In situ localization of transcripts
829	for Foxo4 with Prl7b1 (invasive trophoblast-specific transcript) in the placentation site at gd
830	18.5 of rat pregnancy. Scale bars = 1000 μ m. C) RT-qPCR measurements of <i>Foxo4</i>
831	transcripts in the uterine-placental interface, junctional and labyrinth zones during the second
832	half of gestation (n = $6-9$ /group). Graphs represent means \pm SEM. Asterisks denote statistical
833	difference (** $P < 0.01$) as determined by Steel test. UPI: uterine-placental interface; JZ:
834	junctional zone; and LZ: labyrinth zone. D) Western blot analysis of phospho (p)-FOXO4
835	(Ser ²⁶²) and FOXO4 proteins in $Akt1^{+/+}$ and $Akt1^{-/-}$ placentas at gd 18.5. GAPDH was used as
836	a loading control. E) RT-qPCR measurements of Foxo4 transcripts in the stem state and
837	following induction of differentiation (n = 4-6/group). Graphs represent means \pm SEM.
838	Asterisks denote statistical difference (vs Stem, $*P < 0.05$) as determined by Dunnett's test.
839	Western blot analysis of phospho (p)-FOXO4 (Ser ²⁶²), FOXO4, p-AKT (Ser ⁴⁷³), and
840	pan-AKT proteins in the stem and differentiating (day 15 of differentiation, D15) states (F),
841	and in the differentiating state (D15) following treated with vehicle (0.1% DMSO) or a
842	phosphatidylinositol 3-kinase (PI3K) inhibitor (LY294002, 10 µM) for 1 h (G). GAPDH was
843	used as a loading control.
844	
845	Figure 5. In vivo genome editing of the rat Foxo4 locus. A) Schematic representation of

- the rat *Foxo4* gene ($Foxo4^{Xm+}$) and guide RNA target sites within Exons 2 and 3
- 847 (NM_001106943.1). Red bars beneath Exons 2 and 3 correspond to the 5' and 3' guide RNAs
- used in the genome editing. **B**) The mutant *Foxo4* allele ($Foxo4^{Xm}$) possesses a 3,096 bp

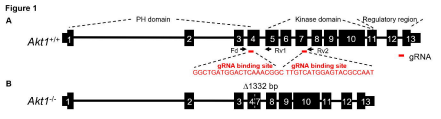
849	deletion. Parts of Exons 2 and 3 are deleted, leading to a frameshift and premature Stop
850	codon in Exon 3. C) Amino acid sequences for $FOXO4^{Xm+}$ and $FOXO4^{Xm-}$. The red sequence
851	corresponds to the frameshift in Exon 3. The green, red, dark blue, and light blue highlighted
852	amino acid sequence regions correspond to the forkhead winged-helix DNA-binding domain
853	(FHD), nuclear localization sequence (NLS), nuclear export sequence (NES), and
854	transactivation domain (TAD), respectively. D) Heterozygous mutant female rats were
855	crossed to wild type male rats to generate hemizygous null male rats. Wild type (+/+),
856	heterozygous (+/-), and hemizygous null (-/y) genotypes were detected by PCR. E) Western
857	blot analysis of FOXO4 protein in the junctional zone of $Foxo4^{Xm+}$ (X ^{m+} Y and X ^{m+} X ^{p+}) and
858	<i>Foxo4</i> ^{Xm-} ($X^{m-}Y$ and $X^{m-}X^{p+}$) placentas at gd 18.5. GAPDH was used as a loading control.
859	
860	Figure 6. Foxo4 hemizygous null and Foxo4 maternally inherited heterozygous
861	conceptuses exhibit placental overgrowth, and FOXO4 deficiency alters the
861 862	conceptuses exhibit placental overgrowth, and FOXO4 deficiency alters the transcriptomes of the junctional zone. Placentas (A) and fetuses (B) were dissected from
862	transcriptomes of the junctional zone. Placentas (A) and fetuses (B) were dissected from
862 863	transcriptomes of the junctional zone. Placentas (A) and fetuses (B) were dissected from <i>Foxo4</i> heterozygous females mated with wild type males at gd 18.5 and weighed; C)
862 863 864	transcriptomes of the junctional zone. Placentas (A) and fetuses (B) were dissected from <i>Foxo4</i> heterozygous females mated with wild type males at gd 18.5 and weighed; C) fetus/placenta ratio. Placentas were then separated into junctional zone (JZ , D) and labyrinth
862 863 864 865	transcriptomes of the junctional zone. Placentas (A) and fetuses (B) were dissected from <i>Foxo4</i> heterozygous females mated with wild type males at gd 18.5 and weighed; C) fetus/placenta ratio. Placentas were then separated into junctional zone (JZ, D) and labyrinth zone (LZ, E) compartments, and weighed; F, JZ/LZ weight ratio. Graphs represent means \pm
862 863 864 865 866	transcriptomes of the junctional zone. Placentas (A) and fetuses (B) were dissected from <i>Foxo4</i> heterozygous females mated with wild type males at gd 18.5 and weighed; C) fetus/placenta ratio. Placentas were then separated into junctional zone (JZ, D) and labyrinth zone (LZ, E) compartments, and weighed; F, JZ/LZ weight ratio. Graphs represent means \pm SEM. X ^{m+} Y, n = 25; X ^{m-} Y, n = 31; X ^{m+} X ^{p+} , n = 14; X ^{m-} X ^{p+} , n = 22 from 8 dams. Asterisks
862 863 864 865 866 867	transcriptomes of the junctional zone. Placentas (A) and fetuses (B) were dissected from <i>Foxo4</i> heterozygous females mated with wild type males at gd 18.5 and weighed; C) fetus/placenta ratio. Placentas were then separated into junctional zone (JZ, D) and labyrinth zone (LZ, E) compartments, and weighed; F, JZ/LZ weight ratio. Graphs represent means \pm SEM. X ^{m+} Y, n = 25; X ^{m-} Y, n = 31; X ^{m+} X ^{p+} , n = 14; X ^{m-} X ^{p+} , n = 22 from 8 dams. Asterisks denote statistical differences (*** <i>P</i> < 0.001) as determined by Student's or Welch's <i>t</i> -test. G
862 863 864 865 866 867 868	transcriptomes of the junctional zone. Placentas (A) and fetuses (B) were dissected from <i>Foxo4</i> heterozygous females mated with wild type males at gd 18.5 and weighed; C) fetus/placenta ratio. Placentas were then separated into junctional zone (JZ, D) and labyrinth zone (LZ, E) compartments, and weighed; F, JZ/LZ weight ratio. Graphs represent means \pm SEM. X ^{m+} Y, n = 25; X ^{m-} Y, n = 31; X ^{m+} X ^{p+} , n = 14; X ^{m-} X ^{p+} , n = 22 from 8 dams. Asterisks denote statistical differences (*** <i>P</i> < 0.001) as determined by Student's or Welch's <i>t</i> -test. G) Vimentin immunostaining of gd 18.5 <i>Foxo4</i> ^{Xm+} and <i>Foxo4</i> ^{Xm} placentation sites. The
862 863 864 865 866 867 868 869	transcriptomes of the junctional zone. Placentas (A) and fetuses (B) were dissected from <i>Foxo4</i> heterozygous females mated with wild type males at gd 18.5 and weighed; C) fetus/placenta ratio. Placentas were then separated into junctional zone (JZ, D) and labyrinth zone (LZ, E) compartments, and weighed; F, JZ/LZ weight ratio. Graphs represent means \pm SEM. X ^{m+} Y, n = 25; X ^{m-} Y, n = 31; X ^{m+} X ^{p+} , n = 14; X ^{m-} X ^{p+} , n = 22 from 8 dams. Asterisks denote statistical differences (*** <i>P</i> < 0.001) as determined by Student's or Welch's <i>t</i> -test. G) Vimentin immunostaining of gd 18.5 <i>Foxo4</i> ^{Xm+} and <i>Foxo4</i> ^{Xm} placentation sites. The junctional zone (JZ) is negative for vimentin immunostaining, whereas the uterine-placental
862 863 864 865 866 867 868 869 869	transcriptomes of the junctional zone. Placentas (A) and fetuses (B) were dissected from <i>Foxo4</i> heterozygous females mated with wild type males at gd 18.5 and weighed; C) fetus/placenta ratio. Placentas were then separated into junctional zone (JZ, D) and labyrinth zone (LZ, E) compartments, and weighed; F, JZ/LZ weight ratio. Graphs represent means \pm SEM. X ^{m+} Y, n = 25; X ^{m-} Y, n = 31; X ^{m+} X ^{p+} , n = 14; X ^{m-} X ^{p+} , n = 22 from 8 dams. Asterisks denote statistical differences (*** <i>P</i> < 0.001) as determined by Student's or Welch's <i>t</i> -test. G) Vimentin immunostaining of gd 18.5 <i>Foxo4</i> ^{Xm+} and <i>Foxo4</i> ^{Xm} placentation sites. The junctional zone (JZ) is negative for vimentin immunostaining, whereas the uterine-placental interface (UPI) and labyrinth zone (LZ) stain positive for vimentin. Scale bars=1000 µm. H)

difference (**P<0.01; ***P<0.001) as determined by Student's or Welch's t-test.

875

876	Figure 7. Reciprocal relationship between AKT1 and FOXO4 in the junctional zone.
877	Venn diagram and heatmaps representing overlap of differentially expressed genes inversely
878	regulated by AKT1 and FOXO4. The heatmap color keys represent z-scores of TPM values.
879	
880	Figure 8. FOXO4 alters the rat TS cell. FOXO4 knockdown efficiency was validated by
881	RT-qPCR (\mathbf{A} , $n = 4$ /group) or western blot (\mathbf{B}) analyses in rat TS cells at day 15 of
882	differentiation following transduction with lentivirus containing a control shRNA or one of
883	two independent <i>Foxo4</i> -specific shRNAs. Graphs represent means \pm SEM. Asterisks denote
884	statistical difference (vs Control shRNA, $*P < 0.05$) as determined by Student's or Welch's
885	<i>t</i> -test. C) Heatmap depicting differentially expressed genes between control and <i>Foxo4</i>
886	shRNA-treated rat TS cells. D) RT-qPCR validation of RNA-seq results (Control shRNA, n =
887	4; <i>Foxo4</i> shRNA 1, n = 4; <i>Foxo4</i> shRNA 2, n = 4). Graphs represent means ± SEM. Asterisks
888	denote statistical difference (compared to Control shRNA, * $P < 0.05$; ** $P < 0.01$; *** $P < 0.$

889 0.001) as determined by Student's or Welch's *t*-test.

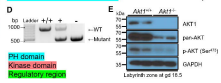


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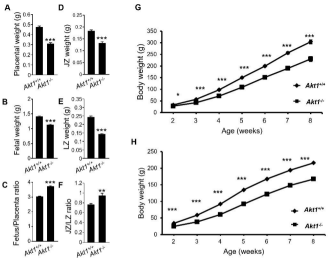
AKT1+/+

MNDVAIVKEGWLHKRGEYIKTWRPRYFLLKNDGTFIGYKERP QDVEORESPLNNFSVAQCQLMKTERPRPNTFIIRCLQWTTVIE RTFHVETPEEREEWTAIOTVADGLRQEEETMDFRSGSPSD NSGAEEMEVALAKPKHRVTMNEFEYLKLLGKGTFGKVILVKEK ATGRYYAMKILKKEVIXAKDEVAHTLTENRVLQNSRHPFLTAL KYSFQTHDRLCFVMEYANGGELFFHLSRERVFSEDRARFYGA EIVSALDYLHSEKNVVYRDLKLENLMLDKDGHIKITDFGLCKEG IKDGATMKTFCGTPEYLAPEVLEDNDYGRAVDWWGLGVVMY GMMCGRLPFYNQDHEKLFELILMELRFRATLGPEAKSLLSGL LKKDPTORLGGGSEDAKEIMQHRFFANIVWQDVYEKKLSPPF KPQJTSEIDTRYFDEEFTAQMITTPPDQDDSMECVDSERRH HFPOFSYASGTA AKT1-

MNDVAJVKEGWLHKRGEYIKTWRPRYFLLKNDGTFIGYKERP QDVEQRESPLNNFSVAQCQLMKTERPRPNTFIIRCLQWTTVIE RTFHVETPEEREEWTTAIQTVADGLKQWGRALLPPVS Stop







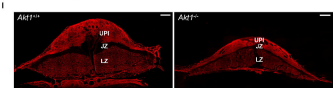


Figure 3

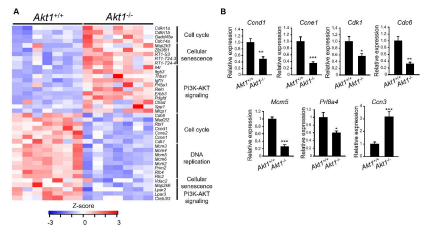


Figure 4

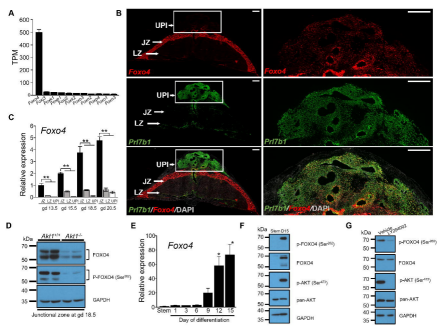
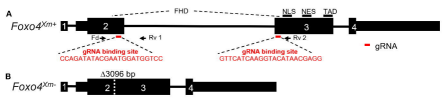


Figure 5



FOXO4Xm-

С

FOXO4Xm+

MEPENKKSATEAAAIIDLDPDFEPOSRPRSCTWPLPRPELATE PPEPSEVEPSLGQKVPTEGHSEPTLLPSRLPEPAGGPQPEILG AVTOPRKGGSRRAWGNQSYAELISOAIESAPEKRLTLAQIYE WMVRTVPYERDKGDSNSSAGWKNSIRHINI SLHSKFIKVHNEA IGKSSWMLNPDGGKGGKAPRRAASMDSSSKLLRGRSKG PKKKPSVLPAPPEGATPRSPLGHFAKWSSSPCPRINEEADV WTTFRPRSSSNASTVSTRLSPMRPESEVLAEEEMPASASSYA GGVPPTLSEDLELLDGLNLASPHSLLSRSNLSSFSLQHPGLAG PLHSYGASLFGPIDGSLSAGEGCFSSSQSLEALLTSDTPPPA DVLMTQVDPILSQAPTLLLLGGMPSSSKLATGVSLCPTPLEGP GPSNLVPTLSWAPPPVMPGAPIPKVLGTPVLASPTEDFSHDR MPQDLDLDMYMENLECOMDNIBDLMDGEGLDFNFEPDP MEPENKKSATEAAAIIDLDPDFEPQSRPRSCTWPLPRPELATE PPEPSEVEPSLGQKVPTEGHSEPTLLPSRLPEPAGGPQPEILG AVTGPRKGGSRRAWGNQSYAELISQAIESAPEKRLTLAGRP PARALGGC

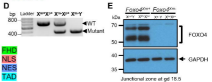


Figure 6

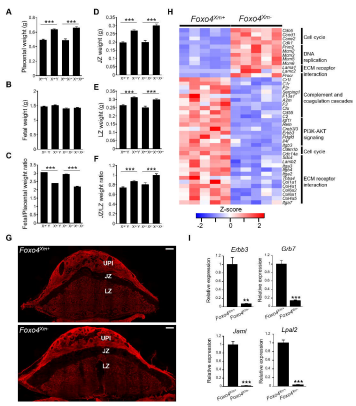
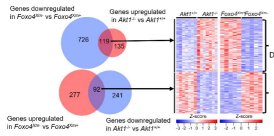


Figure 7



Upregulated in *Akt1^{-/-}* vs *Akt1^{+/+}* and Downregulated in *Foxo4^{Xm-}* vs *Foxo4^{Xm+}*

Donwregulated in *Akt1^{-/-}* vs *Akt1^{+/+}* and Upregulated in *Foxo4^{Xm-}* vs *Foxo4^{Xm+}*

Figure 8

