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# Apposition to endometrial epithelial cells activates mouse blastocysts for implantation

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# Apposition to endometrial epithelial cells activates mouse blastocysts for implantation

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#### Table 1

Gene	Primer sequences (5'-3')	
Cdx2	CAAGGACGTGAGCATGTATCC	
	GTAACCACCGTAGTCCGGGTA	
Gata3	CTCGGCCATTCGTACATGGAA	
	GGATACCTCTGCACCGTAGC	
Eomes	GCGCATGTTTCCTTTCTTGAG	
	GGTCGGCCAGAACCACTTC	
Hand1	CTACCAGTTACATCGCCTACTTG	
	ACCACCATCCGTCTTTTTGAG	
Catal	CACCCCGCCGTATTGAATG	
Galaz	CCTGCGAGTCGAGATGGTTG	
Hes1	CCAGCCAGTGTCAACACGA	
11031	AATGCCGGGAGCTATCTTTCT	
Gandh	AGGTCGGTGTGAACGGATTTG	
Gapun	GGGGTCGTTGATGGCAACA	
Vwbaz	TTGATCCCCAATGCTTCGCAA	
TWHAZ	CAGCAACCTCGGCCAAGTAA	



Figure 1 A Schematic to illustrate the criteria for the 4-point grading of the stability of mouse blastocyst attachment to Ishikawa cells; [1] = not attached, [2] = weakly attached, [3] = intermediately attached, [4] = stably attached. B Hatched E4.5 mouse blastocysts were co-cultured with Ishikawa cells for 48h and the stability of attachment was scored (1-4) every 4h from 0-12h, 24-36h, and at 48h. The mean attachment stability scores, from 4 independent experiments using 102 blastocysts, are presented. Error bars +/- standard error of the mean (SEM). C The mean attachment stability score was calculated from the data in B. Error bars +/- SEM. D Phase contrast images of the mouse blastocyst-Ishikawa cell co-culture at 4h, 8h, 12h, 28h, 32h and 36h. Blastocysts are indicated by arrows and the attachment score for each blastocyst is displayed. Scale bars 200μm. E The cumulative frequency of the time at which the final position of attachment was reached was plotted alongside the percentage of blastocysts. F Ishikawa cells were pre-treated with 10nM 17β-estradiol 48h prior to co-culture, then with 10nM 17β-estradiol and 1μM progesterone 24h prior to co-culture and during co-culture with hatched E4.5 mouse blastocysts. The

attachment stability of mouse blastocysts was scored every 4h from 0-12h, 24-36h, and at 48h. Blastocysts scored as intermediately and stably attached were added together and the mean from 3 independent experiments using 72 blastocysts was plotted +/- SEM.

201x296mm (300 x 300 DPI)



Figure 2 Attached embryos not breaching (A) and breaching (B) Ishikawa cell layers were optically sectioned after PFA fixation and incubation with phalloidin to label actin fibres (red) and DAPI to label nuclei (blue). Invasive cells of breaching embryos exhibit binuclearity (closed arrow), prominent nucleosomes (arrowhead), and actin stress fibres (open arrow). Upper/lower section refers to depth of the optical section in the Z-axis. Scale bars 20µm. C The size of invasive embryonic nuclei in the lower optical section was measured and compared to the size of outer, peripheral embryonic nuclei, presumed to be TE and not inner cell mass, in the corresponding upper section. 309 nuclei were measured in 12 embryos and mean nuclear size was plotted +/- SEM. \*\*\* t-test p<0.001. Breaching embryos were fixed and labelled with anti-CDX2 antibody (D), rabbit isotype control antibody (D inset), anti-GATA3 antibody (E), or mouse isotype control antibody (E inset), as well as phalloidin and DAPI. Upper/lower section refers to depth of the optical section in the Z-axis. Scale bars 20µm. F The intensity of CDX2 and GATA3 fluorescence labelling was measured in invasive embryonic nuclei in the lower optical section and compared to that of outer embryonic nuclei, presumed to be TE and not inner cell mass, in the corresponding upper section. 158 nuclei were measured in 5 embryos labelled with anti-CDX2 and 226 nuclei were measured in 5 embryos labelled with anti-GATA3. The mean ratio of invasive nuclei to upper TE nuclei fluorescence intensity was plotted +/- SEM. \*\* t-test p<0.01, NS not significant.

208x239mm (300 x 300 DPI)



Figure 3 A Schematic representing embryo-Ishikawa cell co-culture experimental design. B Hatched E4.5 mouse blastocysts were either co-cultured with Ishikawa cells to E6.5 or cultured alone in co-culture media to E5.5 before co-culture with Ishikawa cells to E6.5 or E7.5. The stability of attachment was scored every 4h from E5.5-6 and at E6.5. Mean percentage blastocysts scored as intermediately or stably attached from 6 independent experiments using 149 blastocysts was plotted +/- SEM. C Attached embryos co-cultured from E4.5-6.5, E5.5-6.5 and E5.5-7.5, were fixed, labelled with phalloidin and DAPI and optically sectioned to determine whether embryonic cells had breached the Ishikawa cells. The mean proportion of attached embryos to have breached the Ishikawa cells was plotted +/- SEM. \*\* ANOVA p<0.01, \*\*\* ANOVA p<0.001, demonstrating significant difference from co-culture E4.5-6.5.

209x234mm (300 x 300 DPI)



Figure 4 A Schematic representing embryo-Ishikawa cell co-culture experimental design. B Hatched E4.5 mouse blastocysts were co-cultured directly with Ishikawa cells from E4.5-6.5, co-cultured directly with Ishikawa cells to E5.5 before transfer to fresh Ishikawa cells and co-cultured to E6.5, or co-cultured in a transwell above Ishikawa cells to E5.5 before transfer to fresh Ishikawa cells and co-cultured to E6.5. The stability of attachment was scored every 4h from E5.5-6 and at E6.5. Mean percentage blastocysts scored as intermediately or stably attached from 5 independent experiments using 157 blastocysts was plotted +/- SEM. C Attached embryos were fixed, labelled with phalloidin and DAPI and optically sectioned to determine whether embryonic cells have breached the Ishikawa cells. The mean proportion of attached embryos to have breached the Ishikawa cells +/- SEM. \*\* ANOVA p<0.01, demonstrating significant difference from direct co-culture E4.5-6.5.

254x250mm (300 x 300 DPI)



Figure 5 Hatched E4.5 mouse blastocysts were cultured alone, co-cultured directly with Ishikawa cells, cocultured in a transwell above Ishikawa cells or co-cultured directly with human fibroblasts as a control cell type, to E5.5 before retrieval and lysis for RNA isolation and gene expression analysis by qPCR. Data is presented as expression relative to reference genes (2-ct, relative to geometric mean of Gapdh and Ywhaz) of A Cdx2, B Gata3, C Eomes, D Hand1, E Gata2, and F Hes1 in embryos cultured in indicated conditions (mean +/- SEM of 5 independent experiments using 200 embryos). \* ANOVA p<0.05, demonstrating significant difference from no co-culture E4.5-5.5.

207x198mm (300 x 300 DPI)

- 1 Apposition to endometrial epithelial cells activates mouse blastocysts for
- 2 implantation
- 3 **Running title: Blastocyst apposition to endometrial cells**
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## 20 Abstract

21	Study question: How do interactions between blastocyst-stage embryos and endometrial
22	epithelial cells regulate the early stages of implantation in an in-vitro model?
23	Summary Answer: Mouse blastocyst apposition with human endometrial epithelial cells
24	initiates trophectoderm differentiation to trophoblast, which goes on to breach the
25	endometrial epithelium.
26	What is known already: In-vitro models using mouse blastocysts and human endometrial
27	cell lines have proven invaluable in the molecular characterisation of embryo attachment to
28	endometrial epithelium at the onset of implantation. Genes involved in embryonic breaching
29	of the endometrial epithelium have not been investigated in such in-vitro models.
30	Study design, size, duration: This study used an established in-vitro model of implantation
31	to examine cellular and molecular interactions during blastocyst attachment to endometrial
32	epithelial cells.

33 Participants/materials, setting, methods: Mouse blastocysts developed from embryonic 34 day (E) 1.5 in vitro were hatched and co-cultured with confluent human endometrial 35 adenocarcinoma-derived Ishikawa cells in serum-free medium. A scale of attachment 36 stability based on blastocyst oscillation upon agitation was devised. Blastocysts were 37 monitored for 48h to establish the kinetics of implantation, and optical sectioning using 38 fluorescence microscopy revealed attachment and invasion interfaces. Quantitative PCR 39 was used to determine blastocyst gene expression. Data from a total of 680 mouse 40 blastocysts are reported, with 3-6 experimental replicates. T-test and ANOVA analyses 41 established statistical significance at p<0.05, p<0.01 and p<0.001.

Main results and the role of chance: Hatched E4.5 mouse blastocysts exhibited weak
attachment to confluent Ishikawa cells over the first 24h of co-culture, with intermediate and
stable attachment occurring from 28h (E5.5+4h) in a hormone-independent manner.

45 Attached embryos fixed after 48h (E6.5) frequently exhibited outgrowths, characterised 46 morphologically and with antibody markers as trophoblast giant cells (TGCs), which had 47 breached the Ishikawa cell layer. Beginning co-culture at E5.5 also resulted in intermediate 48 and stable attachment from E5.5+4h, however these embryos did not go on to breach the 49 Ishikawa cell layer, even when co-culture was extended to E7.5 (p<0.01). Blastocysts 50 cultured from E4.5 in permeable transwell inserts above Ishikawa cells before transfer to 51 direct co-culture at E5.5 went on to attach but failed to breach the Ishikawa cell layer by E6.5 52 (p<0.01). Gene expression analysis at E5.5 demonstrated that direct co-culture with 53 Ishikawa cells from E4.5 resulted in downregulation of trophectoderm transcription factors 54 Cdx2 (p<0.05) and Gata3 (p<0.05) and upregulation of the trophoblast giant cell transcription 55 factor Hand1 (p<0.05). Co-culture with non-endometrial human fibroblasts did not alter the 56 expression of these genes.

57 Large scale data: N/A

68

Limitations, reasons for caution: The in-vitro model used here combines human carcinoma-derived endometrial cells with mouse embryos, in which the cellular interactions observed may not fully recapitulate those in-vivo. The data gleaned from such models can be regarded as hypothesis-generating, and research is now needed to develop more sophisticated models of human implantation combining multiple primary endometrial cell types with surrogate and real human embryos.

Wider implications of the findings: This study implicates blastocyst apposition to endometrial epithelial cells as a critical step in trophoblast differentiation required for implantation. Understanding this maternal regulation of the embryonic developmental programme may lead to novel treatments for infertility.

69 charities Wellbeing of Women (RG1442) and Diabetes UK (15/0005207), and studentship

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#### 71 Keywords: implantation, endometrium, embryo development, trophoblast,

#### 72 transcription factors

#### 73 Introduction

74 Implantation begins with attachment of the trophectoderm (TE) of a blastocyst-stage embryo 75 to the hormonally-responsive receptive endometrial luminal epithelium (LE), followed by 76 breaching of this barrier and invasion of TE-derived trophoblast into the underlying decidua 77 (Aplin & Kimber 2004, Aplin & Ruane 2017). Implantation failure remains a bottleneck in 78 human assisted reproduction treatment, with only ~25% of treatment cycles resulting in a 79 live birth (Ferraretti et al. 2012), despite efforts to select developmentally-competent 80 embryos and receptive endometrium (Brison et al. 2004, Aplin 2006, Dominguez et al. 2008, 81 Glujovsky et al. 2012, Ruiz-Alonso et al. 2013, Salamonsen et al. 2013, Armstrong et al. 82 2015, Harbottle et al. 2015, Simon et al. 2015). A comprehensive understanding of this 83 founding stage of pregnancy is necessary to improve treatments for infertility. 84 Initial maternal-embryo interactions are mediated through the trophic and paracrine action of 85 uterine fluid, secreted by uterine glands, which promotes embryo development to blastocyst 86 and subsequent implantation (Filant & Spencer 2014). There is now considerable evidence 87 that bidirectional signalling between human blastocysts and endometrium elicits responses 88 which modulate implantation receptivity (Evans et al. 2016). For example, blastocyst-89 dependent local reduction in the anti-adhesive glycocalyx of the LE (Meseguer et al. 2001), 90 and LE microRNA regulation of TE gene expression (Vilella et al. 2015), are suggested to 91 promote embryo attachment. Molecular dialogue has been observed upon blastocyst 92 apposition to the endometrium in mice, with both paracrine and juxtacrine activation of 93 ErbB4 by soluble and membrane-bound HB-EGF, respectively, promoting expression of 94 integrins at the surface of LE and TE, leading to blastocyst attachment and invasion (Wang 95 et al. 2000, Wang et al. 2002). In humans, the homophilic receptor, trophinin, may feed into 96 the HB-EGF axis at apposition to promote TE proliferation in readiness for invasion while

triggering LE apoptosis to allow epithelial breaching (Sugihara *et al.* 2007, Tamura *et al.*2011).

99 After blastocyst attachment to the LE in humans and rodents, embryonic invasion results in 100 haemochorial placentation (Rossant & Cross 2001). The initial invasive cell type in the 101 mouse embryo is the trophoblast giant cell (TGC) (Dickson 1963), whereas in human and 102 macaque this is thought to be primary syncytium (Hertig et al. 1956, Enders 2007). 103 Significant progress has been made in understanding the regulatory networks governing the 104 formation of these cell types (Knott & Paul 2014). Recently trophoblast lineages have been 105 shown to arise from embryos attaching to culture surfaces in the absence of maternal cells 106 (Bedzhov & Zernicka-Goetz 2014, Deglincerti et al. 2016, Shahbazi et al. 2016), however the 107 contribution of endometrial-embryo interactions to the development of the pioneering 108 invasive embryonic cells remains unknown.

109 Characterisation of this early stage of implantation is particularly tractable in vitro. The 110 human endometrial adenocarcinoma Ishikawa cell line (Nishida et al. 1985), offers a model 111 epithelial system for investigating interactions with rodent and human embryos (Singh et al. 112 2010, Kaneko et al. 2011, Kang et al. 2014). Ishikawa cells exhibit moderate epithelial 113 polarisation and surface glycoprotein composition comparable with LE in vivo (Heneweer et 114 al. 2005, Singh & Aplin 2014, Buck et al. 2015). They also mount transcriptional responses 115 to estrogen and progesterone, but do not require steroid hormones for receptivity to mouse 116 embryos (Lessey et al. 1996, Castelbaum et al. 1997, Singh et al. 2010, Tamm-Rosenstein 117 et al. 2013). Ishikawa cell-rodent embryo co-cultures are recognised as a useful model to 118 investigate molecular pathways of attachment (Kaneko et al. 2011, Kaneko et al. 2012, Kang 119 et al. 2014, Green et al. 2015, Kang et al. 2015), however a thorough dissection of 120 attachment and invasion in this model system has not been performed. 121 Here, we have characterised the kinetics of attachment and invasion of mouse embryos on

122 Ishikawa cells. We show that prior to stable attachment, apposition is required for

- subsequent TGC invasion. Moreover, apposition leads to changes in embryonic gene
- 124 expression consistent with TE differentiation to invasive TGCs. Our data suggests that the
- 125 differentiation of trophoblast required for implantation of mouse blastocysts is maternally
- regulated, and implicates a conserved system in human LE.

#### 127 Materials and Methods

#### 128 Cell culture

- 129 Ishikawa cells (ECACC 99040201) and primary human foreskin fibroblasts were maintained
- at 37°C, 95% air and 5% CO<sub>2</sub> in growth medium (1:1 Dulbecco's modified Eagle's
- 131 medium:Ham's-F12 (Sigma) containing 10% fetal bovine serum (Sigma) supplemented with
- 132 2mM L-glutamine, 100µg/ml streptomycin and 100IU/ml penicillin (Sigma)). Cells were
- cultured to confluency in 24-well plates (Greiner) on 13mm glass coverslips coated with 2%
- 134 growth factor-reduced Matrigel (Sigma).

#### 135 Mouse embryos

136 All experiments were conducted and licensed under the Animal Act, 1986, and had local 137 ethical approval for care and use of laboratory animals and standards of humane animal 138 care. CD1 strain mice (Charles River) were maintained by the Biological Services Unit at the 139 University of Manchester and kept under standard environmental conditions of 12h light and 140 12h dark at 20–22°C and 40–60% humidity with food and water provided ad libitum. 8-10-141 week old female mice were superovulated by intraperitoneal injection with 5IU pregnant 142 mare serum gonadotrophin (Intervet), followed by 5IU human chorionic gonadotrophin 143 (Intervet) 46h later, then housed overnight with  $\leq 9$ -month-old stud males for mating. Midday 144 the following day was designated embryonic day (E) 0.5. Embryos were collected at E1.5 by 145 flushing dissected oviducts with M2 medium (Millipore) containing 0.4% w/v BSA (Sigma). All 146 embryo manipulation was performed using a Flexipet with 140µm and 300µm pipettes 147 (Cook). E1.5 embryos were cultured for 72h in KSOM medium (Millipore) containing 0.4%

BSA at 37°C, 95% air and 5% CO<sub>2</sub>. E4.5 blastocysts were hatched from the zona pellucida
(ZP) using acid Tyrode's (pH 2.5) (Sigma).

#### 150 In-vitro implantation model

151 Ishikawa cells were grown to full confluence in 24-well plates, washed and replenished with 152 serum-free co-culture medium (1:1 DMEM:F12 supplemented with 2mM L-glutamine, 100 153 µg/ml streptomycin and 100IU/ml penicillin) 24h before transfer of three hatched mouse 154 blastocysts per well and co-culture at 37°C, 95% air and 5% CO<sub>2</sub>. The stability of mouse 155 blastocyst attachment to Ishikawa cells was assessed using a four-point scale of blastocyst 156 behaviour upon agitation of the sample; translocation (not attached), major oscillation about 157 an attachment point (weakly attached), minor oscillation (intermediately attached), and no 158 oscillation (stably attached) (Figure 1A, Movie 1) (Kang et al. 2014, Kang et al. 2015). The 159 stability of blastocyst attachment was assessed at 4h intervals from 0-12h, 24-36h and 48h 160 using an inverted phase contrast microscope (Evos). At termination, co-cultures were 161 washed in phosphate-buffered saline (PBS) and fixed with 4% paraformaldehyde (PFA) in 162 PBS for 20 minutes. For hormone treatments, Ishikawa cells were maintained in growth 163 medium containing 10nM 17β-estradiol (Sigma) for 24h, then co-culture medium containing 164 10nM 17 $\beta$ -estradiol and 1 $\mu$ M progesterone (Sigma) for a further 24h, before addition of 165 embryos. To separate embryos from Ishikawa cells during co-culture, 3µm transwell filters 166 (Costar) were used. Blastocysts were collected from co-cultures and transwells using a 167 300µm Flexipet.

#### 168 Fluorescence staining and microscopy

Fixed attached embryo samples were washed with PBS and quenched with 50mM
ammonium chloride solution before permeabilisation with 0.5% Triton-X100 PBS. Samples
were incubated with primary antibody (rabbit anti-CDX2 or mouse anti-GATA3, both Cell
Signalling Technologies) in PBS for 2h or overnight, followed by alexa568-phalloidin (Life
Technologies), 4',6-diamidino-2-phenylindole (DAPI) (Sigma), and alexa488 secondary

- antibodies (Life Technologies) for 1h before mounting in a chamber of 3% 1,4-
- diazabicyclo[2.2.2]octane (Sigma) in PBS. Fluorescence microscopy was carried out using a
- 176 Zeiss Axiophot microscope equipped with an Apotome module for optical sectioning. Images
- 177 were analysed and processed using Zeiss Zen software and ImageJ.

#### 178 Blastocyst RNA extraction and quantitative polymerase chain reaction

- 179 RNA from 10 blastocysts per treatment was isolated using the RNeasy Micro Kit (Qiagen),
- according to the manufacturer's instructions. Samples of 25ng RNA were added to reverse
- transcription (RT) reactions with random 9mer primers (Agilent) using the Sensiscript RT kit
- 182 (Qiagen), according to the manufacturer's instructions. Quantitative polymerase chain
- reactions (qPCR) were carried out using the RT reactions along with 0.25µM primers (Table
- 184 1) and QuantiTect SYBR green PCR kit (Qiagen). qPCRs were run on a Stratagene
- 185 Mx3000p machine with thermocycle parameters according to QuantiTect instructions (35
- 186 cycles using 58°C annealing temperature for all primers), and analysed with Stratgene
- 187 MxPro to yield cycle threshold (Ct) values. RT reactions without sample RNA and without
- 188 reverse transcriptase enzyme were used as controls in qPCR reactions with all primer pairs,
- and all yielded no Ct value. Dissociation curves were run with each sample to rule out the
- 190 presence of non-specific PCR products.

#### 191 Statistical Analysis

- 192 Independent t-test and ANOVA with Dunnett's post-hoc test were performed using SPSS
- 193 (IBM), with significance at p<0.05.
- 194 **Results**

#### 195 **E4.5 mouse blastocysts weakly and reversibly attach to lshikawa cells over 24h**

#### 196 before stably attaching in a hormone-independent manner

- 197 E4.5 mouse blastocysts hatched from the ZP barrier were introduced into co-culture with
- 198 confluent Ishikawa cells in serum-free medium and monitored at 4h intervals from 0-12h, 24-

199 36h, and at 48h, to characterise the kinetics of attachment (Figure 1B). Only weak 200 attachment was observed during the first 24h of co-culture (E4.5-5.5), with intermediate and 201 stable attachment increasing from 28-48h (E5.5+4h-E6.5), as demonstrated by plotting the 202 average stability of attachment at each time point (Figure 1C). Weak attachment was 203 reversible since blastocysts cycled between weakly attached and not attached, and 204 positional analysis demonstrated that weak attachment could occur at multiple successive 205 sites (Figure 1D). Later attached blastocysts did not change position over time (Figure 1D), 206 and cumulative frequency of the final position of blastocyst attachment correlated with 207 intermediate and stable attachment scores (Figure 1E). These scores of attachment stability 208 therefore represent irreversible attachment, with the lag between final position of attachment 209 and intermediate-stable attachment likely reflecting a weak attachment phase prior to more 210 stable attachment.

This process of prolonged, reversible weak attachment followed by initiation of stable attachment after 28h co-culture appears to mirror the apposition and attachment stages, respectively, of implantation in vivo (Enders & Schlafke 1969). Thus, the summation of intermediate and stable attachments was used for all subsequent analyses. Additionally, the kinetics of attachment were not altered by treating Ishikawa cells with estradiol and progesterone (Figure 1F), therefore further experiments were performed in the absence of steroid hormones.

#### 218 Mouse blastocysts produce trophoblast giant cells to breach Ishikawa cell layers

Blastocysts attached to Ishikawa cells were fixed after 48h co-culture, fluorescently labelled and optically sectioned to visualise the interface. Although embryos that had attached to the apical surface of the Ishikawa cells but not breached the cell layer were observed (Figure 2A), outgrowths from attached embryos that had breached the Ishikawa cell layer were frequently seen (Figure 2B).

224 These outgrowths contained spread cells with large nuclei, clear nucleoli (arrowheads) and 225 prominent actin stress fibres (thin arrows), and were occasionally bi- and tri-nucleate (thick 226 arrows), reminiscent of trophoblast giant cells (TGCs) (Figure 2B). When compared to the 227 TE nuclei above the plane of the Ishikawa cells (nuclei on the periphery of the embryo), the 228 nuclei of embryonic cells invading into the Ishikawa cell layer were significantly larger (Figure 229 2C). Moreover, antibodies to the TE transcription factors CDX2 and GATA3, which are 230 downregulated upon trophoblast differentiation to TGCs (Knott & Paul 2014), appeared to 231 preferentially label the upper, non-invasive TE nuclei (Figure 2D, E), although only CDX2 232 labelling exhibited significantly lower intensity in the invasive nuclei (Figure 2F). Altogether, 233 these observations demonstrate that mouse embryos can breach and invade Ishikawa cell 234 layers and implicate TGCs in this process.

# 235 **Co-culture from E4.5-5.5 is required for embryos to progress beyond attachment to**

#### 236 breach the Ishikawa cell layer

237 We asked whether early weak attachments influence later embryo-lshikawa interactions by 238 comparing the attachment kinetics of blastocysts cultured with or without Ishikawa cells 239 during E4.5-5.5 (Figure 3A). Embryos without prior co-culture were still able to stably attach 240 to Ishikawa cells during E5.5-E6.5 (Figure 3B), though such embryos exhibited very few 241 breaching events (Figure 3C). Notably, continued co-culture of these embryos to E7.5 still 242 did not yield the levels of breaching observed for embryos co-cultured from E4.5-6.5 (Figure 243 3C). This suggests that E4.5-5.5 blastocysts require a dialogue with Ishikawa cells in order 244 to breach the Ishikawa cell layer at a later embryonic stage.

#### 245 Physical apposition with Ishikawa cells from E4.5-5.5 promotes subsequent

#### 246 embryonic breaching

- To characterise the dialogue between embryos and Ishikawa cells, we employed a
- separated co-culture system. Blastocysts were incubated in permeable transwell inserts
- above Ishikawa cells to allow paracrine crosstalk but prevent direct contact during E4.5-5.5.

The capacity of these embryos to attach and breach after transfer to fresh Ishikawa cells at

11

251 E5.5 was compared to embryos which were co-cultured directly with Ishikawa cells from 252 E4.5 before collection (all embryos, non-attached and attached, were collected) and transfer 253 to fresh Ishikawa cells at E5.5 (Figure 4A). The E5.5-6.5 attachment kinetics of the two 254 groups were indistinguishable and did not differ from a control group of embryos co-cultured 255 directly with Ishikawa cells from E4.5-6.5 without transfer to fresh Ishikawa cells at E5.5 256 (Figure 4B). Conversely, compared to embryos in direct co-culture during E4.5-5.5. 257 significantly fewer embryos co-cultured in transwells from E4.5-5.5 went on to breach 258 Ishikawa cells at E6.5 (Figure 4C). These data rule out priming of Ishikawa cells by embryos 259 from E4.5-5.5 as a mechanism contributing embryonic invasion, and suggest that physical apposition with Ishikawa cells from E4.5-5.5 activates blastocysts for invasion. 260 261 Apposition from E4.5-5.5 induces a TGC differentiation program in the embryo 262 The cells forming the embryonic outgrowths at Ishikawa cell breach sites bore hallmarks of 263 TGCs (Figure 2), leading us to hypothesise that apposition during E4.5-5.5 initiates a TGC 264 differentiation program in the blastocyst TE. Therefore, we assessed the expression of 265 transcription factors associated with TGC differentiation (Simmons & Cross 2005, Liu et al. 266 2009, Knott & Paul 2014), in E5.5 embryos that had not been co-cultured, had been directly 267 or indirectly co-cultured with Ishikawa cells for 24h, or had been directly co-cultured with 268 human foreskin fibroblasts as an alternative cell type. We found significant changes in the 269 expression of the TE transcription factors Cdx2 and Gata3 (0.36- and 0.55-fold 270 downregulation, respectively) and the TGC transcription factor Hand1 (upregulated 3.17-271 fold), but only in blastocysts that had been directly co-cultured with Ishikawa cells from E4.5-272 5.5 (Figure 5A, B, D, respectively). No change in the expression of other TE (*Eomes*) or 273 TGC transcription factors (Gata2 and Hes1) was detected (Figure 5C, E, F, respectively). 274 Other TGC markers, such as Bhlhe40, Mdfi and Prl3d1, were not detected at this stage of

embryonic development (data not shown).

#### 276 Discussion

Dialogue between the embryo and endometrium is necessary to orchestrate implantation in both mouse and human (Wang & Dey 2006). Here, observations in an established in-vitro model of implantation lead to the proposal that blastocyst apposition with LE initiates changes in embryonic gene expression which result in the differentiation of TE to invasive trophoblast. Such maternal regulation of embryonic differentiation constitutes a novel and potentially critical stage in embryo development; a detailed mechanistic understanding of this process could offer targets for the treatment of infertility.

284 Combining a categorical scale of mouse blastocyst attachment stability with microscopic 285 analysis to determine embryonic breaching of the Ishikawa cell layer has revealed 286 progressive stages of weak reversible attachment, stable irreversible attachment and 287 subsequent breaching of the epithelium. The rates of stable attachment and breaching in our 288 in-vitro model correlate with in-vitro-matured blastocyst implantation rates in CD1 mice invivo 289 (Schwarzer et al., 2012; Hemkemeyer et al., 2014). Kinetic analysis demonstrated that in-290 vitro-matured, chemically hatched mouse blastocysts are competent for stable attachment to 291 Ishikawa cells between E5.5 and E6.5, independent of prior co-culture. These findings are 292 consistent with those of a previous study which found that in-vivo-matured, naturally hatched 293 mouse blastocysts co-cultured with Ishikawa cells from E4.5-6.5 attached only after E5.5 294 and did not require prior co-culture (attachment was scored using a binary scale in the face 295 of medium aspiration by pipette, likely to identify only intermediate and stably attached 296 embryos) (Green et al. 2015). Our data also imply that Ishikawa cells are constitutively 297 receptive to mouse blastocyst attachment, as neither co-culture with blastocysts prior to E5.5 298 nor hormonal stimulation was required to induce an adhesive Ishikawa cell phenotype. 299 Increased mouse blastocyst stable attachment to Ishikawa cells in response to estrogen and 300 progesterone has been reported (Singh et al. 2010), however subsequent studies have not 301 used hormones to induce receptivity to rodent and human embryos (Kaneko et al. 2011, 302 Kang et al. 2014, Green et al. 2015). The steroid hormone-mediated shift from non-receptive

303 to receptive endometrium seen in vivo is not replicated by Ishikawa cells in these culture 304 conditions, despite expression of estrogen and progesterone receptors and responsiveness 305 to steroid hormones at the level of gene expression (Tamm-Rosenstein et al. 2013). 306 The reversible weak attachment we observed from E4.5-5.5 may mimic the apposition stage 307 of mouse implantation in vivo, which is thought to occur from E4-5 and entail embryo-308 maternal dialogue leading to locally adhesive LE (Cha & Dey 2014, Aplin & Ruane 2017). 309 Our finding that apposition is required for embryonic breaching of Ishikawa cells suggests 310 that juxtacrine signalling prior to attachment promotes trophoblast penetration of the LE. In 311 contrast, apposition from E4.5-5.5 is not necessary for stable attachment to be achieved in 312 the period from E5.5-6.5. This also suggests that initiating TE differentiation to invasive 313 trophoblast is not required for blastocyst attachment. It is possible that LE-derived signals, 314 such as HB-EGF, trophinin and microRNA hsa-mIR-30d (Wang et al. 2002, Sugihara et al. 315 2007, Vilella et al. 2015), promote stable attachment just prior to and during the process. 316 Importantly, this model shows that the attachment competence of mouse blastocysts is 317 regulated during the window of attachment but progress to invasion requires maternal input 318 during the developmental window prior to attachment. 319 We characterised the invasive embryonic cells in our model as TGCs, as they were 320 occasionally bi-/tri-nucleate, exhibited large nuclei with prominent nucleoli, contained 321 pronounced actin stress fibres, and had reduced CDX2 levels (Simmons et al. 2007). This 322 mimics the pioneering invasive trophoblast observed in vivo (Dickson 1963), indicating that 323 mouse embryos respond to human Ishikawa cells in ways that resemble implantation in vivo. 324 Moreover, the finding that Ishikawa cells, but not human fibroblasts, can induce specific 325 changes in embryo gene expression suggests that human and mouse LE may share some 326 phenotypic properties, and that Ishikawa cells retain receptive LE-like functions despite 327 being derived from an endometrial adenocarcinoma (Nishida et al. 1985).

328 The mechanism of breaching the Ishikawa cells is not yet clear, but our observations are 329 suggestive of an initial narrow penetration, possibly between cells, followed by trophoblast 330 outgrowth into the surrounding cell layer. LE apoptosis at the embryo attachment site has 331 been proposed as a mechanism of LE penetration in mouse and human (Parr et al. 1987, 332 Galan et al. 2000, Tu et al. 2015), and a recent in-vivo study in mice suggested entosis of LE 333 cells by TE as the mechanism of penetration (Li et al. 2015). We did not detect clear 334 morphological signs of either process at sites of embryonic breaching, however induction of 335 TE differentiation to TGC by apposition with LE is not incompatible with these mechanisms. 336 Furthermore, LE induction of TE differentiation to invasive trophoblast recalls human LE-TE 337 interactions from in-vitro studies, whereby homophilic trophinin engagement promotes TE 338 invasion by concomitantly driving LE apoptosis and TE proliferation (Sugihara et al. 2007, 339 Tamura *et al.* 2011).

340 Our investigation implicates the TE differentiation to TGC induced by apposition to LE as a 341 critical step in mouse embryo implantation. Rodent trophoblast differentiation has been 342 widely studied, particularly using blastocyst-derived trophoblast stem cells (TSCs), and 343 transcription factor hierarchies that lead to distinct trophoblast lineages have been 344 determined (Simmons & Cross 2005). Cdx2, Gata3 and Eomes are essential to the first 345 lineage allocation to TE, and are associated with TSC-like states, as their downregulation is 346 required for differentiation into all trophoblast lineages (Guzman-Ayala et al. 2004, Ralston et 347 al. 2010). TGC differentiation from TE/TSCs also requires upregulation of Hand1, Bhlhe40 348 and Mdfi (Cross et al. 1995, Kraut et al. 1998, Scott et al. 2000, Hughes et al. 2004). We 349 detected downregulation of both Cdx2 and Gata3, and upregulation of Hand1 in E5.5 mouse 350 blastocysts specifically after apposition with Ishikawa cells. In addition, anti-CDX2 antibody 351 staining was significantly reduced in embryonic cells breaching the Ishikawa cell layer. We 352 propose that a maternally-derived juxtacrine signal impinges on TE gene expression during 353 apposition to downregulate TSC-like transcription factors and upregulate Hand1. We could

354	not detect Bhlhe40 or Mdfi, suggesting that Hand1 is an early, maternally-regulated TGC
355	transcription factor.

- In humans, the pioneering invasive trophoblast at implantation is thought to be primary syncytium (Hertig *et al.* 1956, Aplin & Ruane 2017), the formation of which is controlled by the expression of the syncytium regulator, transcription factor GCM1 (Yu *et al.* 2002, Liang *et al.* 2010). The existence of an Ishikawa cell-derived signal that promotes mouse TE differentiation in this in-vitro model implicates maternally-regulated induction of trophoblast differentiation as an important mechanism in human embryo implantation. Understanding such events may lead to the development of novel treatments for implantation failure in
- human ART.

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368 Authors' roles

- 369 PTR, SJK, DRB, MW and JDA designed the study, and SJK, DRB, MW and JDA obtained
- funding. PTR, SCB, RK and JW carried out the experimental work. PTR wrote the paper,
- which was edited by SJK, DRB, MW and JDA.

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## 375 **Conflict of interest**

376 We declare no conflicts of interest.

#### 377

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#### 548 Table, movie and figure legends

- 549 **Table 1** Mouse gene names and corresponding forward and reverse primer sequences.
- 550 Primer sequences were obtained from PrimerBank (Spandidos et al. 2010), with the
- 551 following references: Cdx2 31560722a1; Gata3 6679951a1; Eomes 26354683a1; Hand1
- 552 118130896c3; Gata2 226530725c1; Hes1 6680205a1; Gapdh 126012538c1; Ywhaz
- 553 **359385697c3**.
- 554 **Movie 1** Differential interference contrast imaging at 14 frames per second of E5.5+4h
- 555 mouse embryos after 28h co-culture with Ishikawa cells exhibiting, upon agitation, weak
- attachment (major oscillation about an attachment point, left embryo), intermediate
- 557 attachment (minor oscillation, right embryo) and stable attachment (no oscillation, middle
- 558 embryo) to Ishikawa cells. Scale bar 100μm.
- 559 Figure 1 A Schematic to illustrate the criteria for the 4-point grading of the stability of mouse
- 560 blastocyst attachment to Ishikawa cells; [1] = not attached, [2] = weakly attached, [3] =
- 561 intermediately attached, [4] = stably attached. B Hatched E4.5 mouse blastocysts were co-
- 562 cultured with Ishikawa cells for 48h and the stability of attachment was scored (1-4) every 4h
- from 0-12h, 24-36h, and at 48h. The mean attachment stability scores, from four

564 independent experiments using 102 blastocysts, are presented. Error bars +/- standard error 565 of the mean (SEM). C The mean attachment stability score was calculated from the data in 566 B. Error bars +/- SEM. D Phase contrast images of the mouse blastocyst-Ishikawa cell co-567 culture at 4h, 8h, 12h, 28h, 32h and 36h. Blastocysts are indicated by arrows and the 568 attachment score for each blastocyst is displayed. Scale bars 200µm. E The cumulative 569 frequency of the time at which the final position of attachment was reached was plotted 570 alongside the percentage of blastocysts scored as intermediately or stably attached (mean 571 +/- SEM from three independent experiments using 84 blastocysts). F Ishikawa cells were 572 pre-treated with 10nM 17 $\beta$ -estradiol 48h prior to co-culture, then with 10nM 17 $\beta$ -estradiol 573 and 1µM progesterone 24h prior to co-culture and during co-culture with hatched E4.5 574 mouse blastocysts. The attachment stability of mouse blastocysts was scored every 4h from 575 0-12h, 24-36h, and at 48h. Blastocysts scored as intermediately and stably attached were 576 added together and the mean from three independent experiments using 72 blastocysts was 577 plotted +/- SEM.

578 Figure 2 Attached embryos not breaching (A) and breaching (B) Ishikawa cell layers were 579 optically sectioned after PFA fixation and incubation with phalloidin to label actin fibres (red) 580 and DAPI to label nuclei (blue). Invasive cells of breaching embryos exhibit binuclearity 581 (thick arrow), prominent nucleosomes (arrowhead), and actin stress fibres (thin arrow). 582 Upper/lower section refers to depth of the optical section in the Z-axis. Scale bars 20µm. C 583 The size of invasive embryonic nuclei in the lower optical section was measured and 584 compared to the size of outer, peripheral embryonic nuclei, presumed to be TE and not inner 585 cell mass, in the corresponding upper section; 309 nuclei were measured in 12 embryos and 586 mean nuclear size was plotted +/- SEM. \*\*\* t-test p<0.001. Breaching embryos were fixed 587 and labelled with anti-CDX2 antibody (**D**), rabbit isotype control antibody (**D** inset), anti-588 GATA3 antibody (E), or mouse isotype control antibody (E inset), as well as phalloidin and 589 DAPI. Upper/lower section refers to depth of the optical section in the Z-axis. Scale bars 590 20µm. F The intensity of CDX2 and GATA3 fluorescence labelling was measured in invasive

embryonic nuclei in the lower optical section and compared to that of outer embryonic nuclei,
presumed to be TE and not inner cell mass, in the corresponding upper section; 158 nuclei
were measured in five embryos labelled with anti-CDX2 and 226 nuclei were measured in
five embryos labelled with anti-GATA3. The mean ratio of invasive nuclei to upper TE nuclei
fluorescence intensity was plotted +/- SEM. \*\* t-test p<0.01, NS not significant.</li>

596 Figure 3 A Schematic representing embryo-Ishikawa cell co-culture experimental design. B 597 Hatched E4.5 mouse blastocysts were either co-cultured with Ishikawa cells to E6.5 or 598 cultured alone in co-culture media to E5.5 before co-culture with Ishikawa cells to E6.5 or 599 E7.5. The stability of attachment was scored every 4h from E5.5-6 and at E6.5. Mean 600 percentage blastocysts scored as intermediately or stably attached from six independent 601 experiments using 149 blastocysts was plotted +/- SEM. C Attached embryos co-cultured 602 from E4.5-6.5, E5.5-6.5 and E5.5-7.5, were fixed, labelled with phalloidin and DAPI and 603 optically sectioned to determine whether embryonic cells had breached the Ishikawa cells. 604 The mean proportion of attached embryos to have breached the Ishikawa cells was plotted 605 +/- SEM. \*\* ANOVA p<0.01, \*\*\* ANOVA p<0.001, demonstrating significant difference from 606 co-culture E4.5-6.5.

607 Figure 4 A Schematic representing embryo-Ishikawa cell co-culture experimental design. B 608 Hatched E4.5 mouse blastocysts were co-cultured directly with Ishikawa cells from E4.5-6.5, 609 co-cultured directly with Ishikawa cells to E5.5 before transfer to fresh Ishikawa cells and co-610 culture to E6.5, or co-cultured in a transwell above Ishikawa cells to E5.5 before transfer to 611 fresh Ishikawa cells and co-culture to E6.5. The stability of attachment was scored every 4h 612 from E5.5-6 and at E6.5. Mean percentage blastocysts scored as intermediately or stably 613 attached from five independent experiments using 157 blastocysts was plotted +/- SEM. C 614 Attached embryos were fixed, labelled with phalloidin and DAPI and optically sectioned to 615 determine whether embryonic cells have breached the Ishikawa cells. The mean proportion 616 of attached embryos to have breached the Ishikawa cells +/- SEM. \*\* ANOVA p<0.01,

demonstrating significant difference from direct co-culture E4.5-6.5.

- 618 Figure 5 Hatched E4.5 mouse blastocysts were cultured alone, co-cultured directly with
- 619 Ishikawa cells, co-cultured in a transwell above Ishikawa cells or co-cultured directly with
- 620 human fibroblasts as a control cell type, to E5.5 before retrieval and lysis for RNA isolation
- 621 and gene expression analysis by qPCR. Data is presented as expression relative to
- 622 reference genes (2<sup>-ct</sup>, relative to geometric mean of Gapdh and Ywhaz) of A Cdx2, B Gata3,
- C Eomes, D Hand1, E Gata2, and F Hes1 in embryos cultured in the indicated conditions 623
- 624 (mean +/- SEM of five independent experiments using 200 embryos). \* ANOVA p<0.05,
- 625 demonstrating significant difference from no co-culture E4.5-5.5.

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