Let-7 derived from endometrial extracellular vesicles is an important inducer of embryonic diapause in mice

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Embryonic diapause is a maternally controlled phenomenon. The molecule controlling the onset of the phenomenon is unknown. We demonstrated that overexpression of microRNA let-7a or incubation with let-7g–enriched extracellular vesicles from endometrial epithelial cells prolonged the in vivo survival of mouse blastocysts, which developed into live pups after having been transferred to foster mothers. Similar to in vivo dormant blastocysts, let-7–induced dormant blastocysts exhibited low level of proliferation, apoptosis, and nutrient metabolism. Let-7 suppressed c-myc/mTORC1 and mTORC2 signaling to induce embryonic diapause. It also inhibited ODC1 expression reducing biosynthesis of polyamines, which are known to reactivate dormant embryos. Furthermore, the overexpression of let-7 blocked trophoblast differentiation and implantation potential of human embryo surrogates, and prolonged survival of human blastocysts in vitro, supporting the idea that embryonic diapause was an evolutionary conserved phenomenon. In conclusion, let-7 is the main factor inducing embryonic diapause.

INTRODUCTION
Embryonic diapause refers to a reversible arrest of the development of blastocysts. In wildlife, the phenomenon ensures that the young are born in a favorable environment. Embryonic diapause can be induced in a laboratory condition and is best studied in rodents. In mice, ovariectomy in the early morning of day 4 post-fertilization followed by progesterone administration induces embryonic diapause, during which the blastocysts become dormant and do not implant in the uterus, but can implant and develop normally when reactivated by a single dose of estradiol (1). A recent study showed that sheep blastocysts, which do not normally undergo embryonic diapause, could be induced to become dormant in the uterus of delayed implanting mice and could be reactivated when transferred back to the uterus of the ewe (2), suggesting that embryonic diapause is an evolutionary conserved phenomenon.

The onset, maintenance, and termination of embryonic diapause are under maternal control (3). Various factors regulate embryonic diapause. For instance, photoperiod influences embryonic diapause in minks and lactation affects that in rodents (3). High levels of uterine anandamide maintain embryonic diapause in mice (4), while the metabolite of estrogen, catechol estrogen (5), and polyamines (6) reactivate the dormant embryos in mice and mink, respectively. However, the factors that induce the onset of embryonic diapause in vivo remain unknown.

Let-7 is a family of microRNAs up-regulated in the dormant mouse blastocysts (7). Their expressions return to a low level after estrogen-induced reactivation (8). Although an overexpression of let-7a suppresses implantation (7), the origin and role of let-7 in the dormant blastocysts are unknown. As embryonic diapause is mainly a maternally controlled event and endometrial cells produce extracellular vesicles (EVs) containing microRNAs (9), we tested the hypothesis that endometrial cells produced let-7 containing EVs to induce embryonic diapause in this report. The mechanisms of the action of let-7 on embryonic diapause were investigated. In addition, we provided evidence suggesting that the actions of let-7 on embryos were conserved in humans.

RESULTS
Let-7 induces embryonic diapause
To test the hypothesis, the precursor of let-7a (pre-let-7a) or scrambled RNA (control) was electroporated into mouse blastocysts on day 4 of pregnancy. The level of mature let-7a was >40-fold higher in the pre-let-7a blastocysts than in the control blastocysts (Fig. S1A). After 3 days of culture (day 7), the level of let-7a remained threefold higher in the pre-let-7a group (Fig. S1A). The majority of the pre-let-7a blastocysts (95 ± 3.2%) were morphologically viable, with a large blastocoel (Fig. 1A), whereas 31 ± 2.2% of the control embryos had shrunken in size and some degenerated by that time. On day 12, over 50% of the embryos with a pre-let-7a overexpression remained morphologically viable, while all the control embryos had degenerated (Fig. 1A).

Consistent with the above observation, the percentage of apoptotic [terminal deoxynucleotidyl transferase–mediated deoxyuridine triphosphate nick end labeling–positive (TUNEL⁺) cells] in the day 7 blastocysts (N = 6 embryos per group) with electroporation of pre-let-7a on day 4 (D7-let7) was significantly lower than that of the control embryos (D7-scr, P < 0.05; Fig. 1B). Blastocysts collected in the afternoon of day 4 of pregnancy (in vivo activated blastocysts, D4-act) had a low percentage of TUNEL⁺ cells comparable to that of D7-let7 blastocysts (Fig. 1B).

It is well established that metabolism is suppressed in the dormant embryos (10). Therefore, we tested whether an overexpression of let-7a would change the energy metabolism of embryos. As expected, D7-let7 and in vivo dormant (Dor) blastocysts had low glucose.
metabolism; their glucose, pyruvate uptakes, and adenosine triphosphate (ATP) levels were significantly lower than those of the D4-act and estrogen-induced reactivated (E2-act) blastocysts, respectively (Fig. 1C; \( N = 5 \)). The reduced glucose metabolism was likely a result of the inhibitory action of let-7 on its target genes related to glucose metabolism (predicted by TargetScan or RNA22), \( Hk2, \ Fbp1, \ Dld, \) and \( Dlst \) (Fig. 1D). Lactate production was unexpectedly significantly higher in the D7-let7 blastocysts than in the D4-act blastocysts (Fig. 1C), which might be due to a high level of \( Ldha \) (lactate dehydrogenase A) mRNA in the former (Fig. 1D).

The effects of an overexpression of let-7a on the proliferation and extent of the active DNA synthesis of blastomeres were determined by immunostaining for Ki67 and 5-ethynyl-2’-deoxyuridine (EdU) incorporation assay, respectively. The percentage of Ki67+ cells in the D7-let7 blastocysts (\( N = 12 \)) was significantly lower than that of the D4-act blastocysts (\( N = 10, P < 0.05 \); Fig. 1E). The Ki67+ cells in the D7-let7 blastocysts were concentrated in the inner cell mass; this is consistent with the observation that the mouse trophectoderm cells entered dormancy earlier than the inner cell mass cells (11, 12). Consistent with cell proliferation, the percentage of EdU+ cells was about fourfold higher (\( P < 0.05 \)) in the D4-act blastocysts (\( N = 5 \)) than in the D7-let7 blastocysts (\( N = 7 \)) (Fig. 1F). The observations were in line with the report that cell cycle arrest in the delayed implanting blastocysts occurs before the S phase (13).

An increase in epidermal growth factor (EGF) binding is a marker of reactivation of dormant mouse blastocysts (5). Dormant blastocysts did not bind a significant amount of fluorescein isothiocyanate (FITC)–labeled EGF (Dor; Fig. 1G). EGF binding was up-regulated in blastocysts from delayed implanting mice at 6 hours after estradiol-induced reactivation (E2-act). However, D7-let7 and D7-scr blastocysts exhibited a low EGF binding. The similarly low EGF binding of D7-let7 and D7-scr blastocysts could be attributed to different causes; the former reflected a diapause state, while the latter reflected a deteriorating state with many apoptotic cells (Fig. 1B).

**Let-7–induced embryonic diapause is reversible**

To determine whether the overexpression of let-7 in embryos had been in a reversible dormant state in vivo, a 2-fluoroestradiol-17β (2-Fl-E2)–treated mouse model was used. 2-Fl-E2 treatment induces a receptive state of the endometrium (5). It also inhibits the uterine estrogen-2/4-hydroxylase activity, thereby suppressing the synthesis of 4-hydroxy-E2, a catechol metabolite of estradiol-17β (E2) required for the activation of dormant mouse blastocysts (5). Thus, 2-Fl-E2 injection induces uterine receptivity but fails to reactivate the dormant blastocysts in the delayed implanting mice (5). We transferred D7-let7 blastocysts into the 2-Fl-E2–treated pseudo-pregnant uteri (\( N = 6 \) mice per group); the number of implantation sites at 28 hours after embryo transfer was significantly lower than that after the transfer of the D4-act blastocysts (\( P < 0.05 \); Fig. 2A).

The reversible nature of let-7–induced embryonic diapause was tested by transferring D7-let7 blastocysts into D3 pseudo-pregnant uteri. Live births were obtained from blastocysts with an overexpression of let-7a for 3 days (D7-let7) and 4 days (D8-let7; Fig. 2B). All the pups had normal birth weight (Fig. S1B). No pup was born after the transfer of the D12-let7 and D7-scr blastocysts (data not shown).

In the above experiments, electroporation induced only a transient rise in the let-7a level for a few days. To determine the prolonged action of a high level of let-7 on embryo survival in vitro, we produced transgenic mice from embryonic stem cells with doxycycline (DOX)–inducible let-7g (a gift from G. Q. Daley, Harvard Stem Cell Institute, Boston, MA, USA). DOX treatment induces let-7g expression in the transgenic mice. The transgene is unique in that the loop region of the precursor let-7g (pre-let-7g) in the transgene is replaced by that of microRNA-21. Therefore, endogenous Lin28 cannot bind to pre-let-7g and block let-7g biogenesis in the transgenic mice (14). Let-7 family members contain a similar “seed sequence” that spans from nucleotide 2 to 8 in mice (15). This conserved feature suggests that the let-7 family members have similar target mRNAs and functions. Identical changes in the expression of 14 genes were observed in the blastocysts after an overexpression of let-7a and let-7g (Fig. S2). The above embryo transfer experiment was repeated with the transgenic mice.

**Fig. 1. Overexpression of let-7a prolongs survival of mouse blastocysts in vitro.** (A) Overexpression of let-7a extended the survival of blastocysts in culture. Survived blastocysts were defined as those with a well-defined blastocoel. (B) The percentage of apoptotic cells was low in blastocysts with overexpression of let-7a on day 7 (D7-let7) and was comparable to that in day 4 activated (D4-act) blastocysts. Blastocysts electroporated with scrambled RNA (D7-scr) had significantly higher percentage of TUNEL+ cells. *\( P < 0.05. \) (C) Glucose metabolism of the D7-let7 and in vivo dormant (Dor) blastocysts was low with significant reduction in glucose, pyruvate uptake, and ATP levels when compared with the D4-act blastocysts and E2-induced reactivated blastocysts (E2-act), respectively. D7-let7 had a high lactate production, which was low in the dormant blastocysts. *\( P < 0.05. \) (D) Expression of genes related to glucose metabolism. (E and F) The percentages of proliferating (Ki67+, green) cells (E) and cells with DNA synthesis (EdU+, green) (F) were lower in blastocysts of D7-let7 when compared to that in D4-act blastocysts. Numbers in parenthesis are number of embryos analyzed. *\( P < 0.05. \) (G) Dor, D7-let7, and D7-scr blastocysts did not bind significant amount of FITC-labeled EGF. The binding of EGF was high in the D4-act blastocysts and the delayed implanting mice 6 hours after E2-act.
When blastocysts from the transgenic mice were treated with DOX during culture, the let-7g level in the embryos increased 32-fold, and 50% of the blastocysts survived for 14 days in vitro (Fig. 2C). Pups were obtained after transfer of DOX-treated D7 blastocysts and D9 blastocysts, although the live birth rate was lower than that from D4-act blastocysts. Numbers in parenthesis are number of live birth/total number of embryos transferred. (C) Survival curve of blastocysts carrying an inducible let-7g transgene with and without DOX treatment cultured in KSOM + AA medium. Most of the control embryos survived until day 6, whereas 50% of the DOX-treated embryos maintained good morphology even on day 13. (D) Live births were obtained after transfer of let-7g blastocysts treated with DOX treatment for 3 days (D7+DOX) and 5 days (D9+DOX). No live birth was obtained from transfer of day 7 WT blastocysts. Numbers in parenthesis are number of live birth/total number of embryos transferred. (E) Microarray analyses of the mRNA expression of D4-act (D4-act-7g), in vitro (DOX treatment in culture, DOX-7g), and in vivo induced dormant (Dor-7g) let-7g blastocysts. Heatmap of the top 500 differentially expressed genes between the dormant and D4-act blastocysts was shown. (F) Venn diagram showing the number of differentially expressed genes between in vitro (DOX treatment in culture) or in vivo induced dormant blastocysts against D4-act blastocysts. Photo credit: W. M. Liu, The University of Hong Kong.

Transcriptomic analysis of let-7-induced embryos
To understand the molecular actions of let-7 on embryonic diapause, the GeneChip™ Mouse Gene 2.0 ST Array was used to determine the transcriptomes of D4-act, in vivo dormant, and let-7–induced dormant blastocysts from let-7g–transgenic mice. Unsupervised hierarchy clustering (Fig. 2E) and principal component analysis (fig. S3A) showed that the mRNA profiles (table S1) of the in vitro induced dormant blastocysts (DOX treatment in culture, DOX-7g) and the in vivo dormant blastocysts (Dor-7g) were similar but distinct from those of the D4-act let-7g blastocysts (D4-act-7g). To confirm the data obtained from the mRNA profiles, the total RNAs isolated from five pools of D4-act-7g, Dor-7g, and DOX-7g blastocysts were subjected to direct quantitative polymerase chain reaction (qPCR) analyses for the transcript levels of six genes, namely, Ccn1, Btg1, Pkm, Oxtct1, Fbp1, and Sap1 (fig. S3B). These genes were involved in cell cycle (Ccn1 and Btg1), carbohydrate metabolism, energy pathway (Pkm and Oxtct1), and chromatin remodeling (Sap1). The expression patterns were consistent with the results of the array. Among them, two genes (Btg1 and Oxtct1) were significantly higher, while the rest were significantly lower in the Dor-7g and DOX-7g blastocysts than in the D4-act-7g blastocysts (P < 0.05).

Compared with the D4-act-7g blastocysts, the in vitro and the in vivo induced dormant blastocysts exhibited 3444 and 2452 differentially expressed genes, respectively (Fig. 2F). Among the differentially expressed genes, 1006 of them were common in the two comparisons (Fig. 2F). Gene ontology analysis of these common genes using Database for Annotation, Visualization, and Integrated Discovery (DAVID) showed that they were related to mitotic nuclear division, cell division, G1-S transition of mitotic cell cycle, DNA repair, DNA replication, and cell cycle (table S2).

Transfer of let-7 in uterine fluid EVs to embryos
Because embryonic diapause is a maternally regulated phenomenon, we tested the possibility of a maternal origin of let-7 in the delayed implanting blastocysts. This possibility is supported by three observations. First, transmission and scanning electron microscopy showed EV-like structures in the mouse uterine lumen (fig. S4A) and on the trophoderm of blastocysts collected in the uterine lumen (fig. S4B), respectively. Second, immunostaining detected the presence of CD63, a marker of EVs, in the uterine epithelium of mice (fig. S4C) and on the surface of EVs from uterine luminal fluid (ULF; fig. S4D). Nanoparticle tracking analysis showed that the EVs in ULF had a mean size of 82.3 nm (fig. S4E). Western blot analysis showed that these ULF-EVs were positive for HSP70, CD63, and TSG101 and negative for calnexin and GM130. Third, mouse ULF contains let-7 carrying CD63+ EVs, which can be internalized by blastocysts (16). To obtain further evidence, the expression patterns of let-7 in the endometrial epithelial cells, EVs from ULF (ULF-EVs), and in blastocysts from mice before embryonic diapause, during embryonic diapause, and after E2-induced reactivation were determined. The results showed that they had similar patterns with a high let-7a expression only in the dormancy period (Fig. 3A). The pattern was markedly different from that of the stromal cells (fig. S5), supporting the idea that the endometrial epithelial cells produced EVs containing let-7a during delayed implantation.

The inducible let-7g transgenic mice carry a unique chimeric let-7g Stem/mir-21-21 loop sequence (S7gL21), in which the loop of pre-let-7g is replaced by that of microRNA-21. qPCR assay was developed to detect the expression of the transgene; the forward primer targeted
on a sequence that crossed the stem and the loop of S7gL21 so that only the precursor of the transgene, but not pre-let-7g, was amplified. DOX treatment significantly induced the expression of the sequence in the liver (data not shown) and ULF-EVs (Fig. 3B, upper panel) of the transgenic mice, but not that of the ICR (Institute of Cancer Research) mice. Blastocysts from ICR mice were transferred into the pseudo-pregnant delayed implanting let-7g transgenic mice. Three days later, the expression of the S7gL21 sequence was significantly higher in the transferred embryos than in those that were not transferred (Fig. 3B, lower panel). The result confirmed that let-7 was transferred from the mother to the embryos in vivo.

To study the biological effect of let-7a and let-7g in EVs, we produced let-7–enriched EVs by two methods. First, let-7a–enriched EVs were isolated from the spent culture medium of human endometrial Ishikawa cells transfected with pre-let-7a. Transfection with scrambled RNA was used as control. The collected EVs at physiological concentration were then used to treat day 4 blastocysts or human trophoblast JEG-3 cells. Endometrial EVs containing let-7a, but not EV-free let-7a, were biologically active in suppressing the expression of let-7 targets, LIN28A [lin-28 homolog A (Caenorhabditis elegans); Fig. 3C], C-MYC (MYC proto-oncogene), and RICTOR (RPTOR independent companion of MTOR, complex 2; fig. S5B) in the JEG-3 cells. Incubation of the let-7a–enriched EVs for 24 hours significantly decreased the expression of c-myc in the treated blastocysts relative to the control embryos (Fig. 3D) and reduced their DNA synthesis (let-7 EVs; Fig. 3E) to a level comparable to that in the in vivo dormant blastocysts (Dor; Fig. 3E).

To better simulate the in vivo situation, let-7g–enriched EVs were obtained from endometrial epithelial cells of let-7g transgenic mice treated with DOX for 4 days in a medium supplemented with 10% EV-free fetal bovine serum (FBS) and were used at physiological concentration to treat WT day 4 blastocysts in KSOM + amino acid (AA) medium. After 3 days of culture, 82 ± 16.2% (N = 150) of the let-7g-EV–treated blastocysts still had the blastocoel, and 12 ± 5.4% of them developed to term after an embryo transfer. In contrast, 85 ± 10.1% (N = 100) of the blastocysts degraded in the absence of EVs (Fig. 3F), and the survived ones produced no pups after transfer.

**Let-7 is upstream of mTOR and c-myc signaling in delayed implanting blastocysts**

Inhibition of mTOR (mammalian target of rapamycin) (17) or MYC (18) induces a diapause-like state in mouse embryos. We consistently observed significant decreases in the transcript expression of c-myc and Akt1 (mTOR activator) and increases in that of Tsc1 and Tsc2 (mTOR inhibitors) in the let-7g–induced dormant blastocysts when compared with the untreated blastocysts (Fig. 4A). On the other hand, Pten of PIK3 (phosphatidylinositol-3-kinase) signaling, an upstream pathway of mTOR, was unaffected by the DOX treatment.

Next, we determined whether c-myc mediated the action of let-7g on the induction of embryonic diapause. In vitro transcription was used to generate c-myc mRNA. The mRNA was biologically active, and the level of C-MYC protein in the JEG-3 cells increased fourfold at 24 hours after transfection of the mRNA (Fig. 4B). Electro- poration of the c-myc mRNA enhanced C-MYC expression (fig. S6A) and Edu incorporation (Fig. 4C) in D4 blastocysts. Overexpression of c-myc also nullified the effects of let-7g overexpression on the prolongation of the embryo survival in vitro; in the presence of DOX, the let-7g transgenic blastocysts transfected with c-myc mRNA could only survive until day 6; the vast majority died the next day, whereas 50% of the DOX-treated transgenic blastocysts without the transfection survived up to day 15 (fig. S6B).

C-MYC is upstream of mTORC1 in rat fibroblasts (19). The incubation of JEG-3 cells with an inhibitor of mTORC1/mTORC2 pathways, INK-128, for 24 hours reduced phosphorylation of mTORC1 target and RpS6 phosphorylated at Ser235/236 (pRpS6) and had no effect on the expression of C-MYC (Fig. 4D). On the other hand, treatment with the C-MYC inhibitor 10058-F4 [MYC inhibitor (MI)] (20) significantly decreased the phosphorylation of mTORC1 targets, pRpS6, and p4EBP1 (4EBP1 phosphorylated at Thr37/46) in
identified that a component of mTORC2, Rictor, was a potential target of let-7. The prediction was supported by the down-regulation of RICTOR expression in the JEG-3 cells at 48 hours after transfection of the cells with pre-let-7a (Fig. 4G). Dual luciferase assay confirmed that Rictor was a direct target of let-7a. At 24 hours after transfection, let-7a mimics reduced the luciferase activity of Rictor 3′ untranslated region (3′UTR) reporter by about fivefold when compared to the scramble control (Fig. S7). Mutation of the let-7a binding sites on the reporter construct abolished the reduction in luciferase activity (fig. S7). As expected, the expression of RICTOR was lower in the dormant embryos (Dor) than in those at 6-hour post–E2-induced reactivation (A6h; Fig. 4H). The action of let-7 on mTORC2 signaling was further confirmed by a significant decrease in pAKT expression in the JEG-3 cells at 48 hours after transfection of pre-let-7a when compared with the scramble control (Fig. 4G).

**Let-7 inhibits production of polyamines**

The inhibition of polyamine biosynthesis delays the reactivation of dormant embryos in vitro (21). We consistently found an increased expression of polyamine biosynthetic enzymes, ornithine decarboxylase (ODC1) and spermine synthesis enzyme (SMS) in the reactivated embryos (A24h), relative to the dormant embryos (Dor; Fig. 5A).

Odc1 encodes a rate-limiting enzyme in polyamine biosynthesis. An analysis of a genome-wide embryonic C-MYC chromatin immunoprecipitation (ChIP) sequencing dataset (22) revealed recruitment of C-MYC at the promoter of Odc1 and identified five putative c-myc response elements on the promoter (Fig. 5B). Luciferase reporter assays showed that response elements 3, 4, and 5 were the major sites that conferred C-MYC responsiveness (Fig. 5B). Site-specific ChIP assays using the DNA from mouse primary uterine epithelial cells isolated from delayed implanting (Dor-ME) and day 4 activated mouse uteri (D4-act) confirmed the recruitment of C-MYC to these response elements (Fig. 5C). There were reductions in the recruitment of C-MYC to these elements of Odc1 promoter in the uterine epithelial cells during embryonic diapause (Dor-ME) when compared to those from D4-act mice (Fig. 5C).

Our data further showed that the inhibition of C-MYC by MI for 24 hours significantly decreased the protein level of ODC1 in the JEG-3 cells (Fig. 5D) and D4 blastocysts (Fig. 5E). Treatment with DOX reduced the protein expression of ODC1 in the let-7g blastocysts relative to the D4 blastocysts (Fig. 5F). The action was mediated by C-MYC as the transfection of c-myc mRNA abolished the let-7–induced down-regulation of ODC1 in the embryos (Fig. 5F). Together, endometrial epithelial cell–derived let-7 suppressed c-myc/mTORC1, mTORC2 signaling, and polyamine biosynthesis to induce embryonic diapause (Fig. 5G).

**Let-7–enriched EVs modulate human blastocyst development and implantation**

Non-diapause sheep blastocysts become dormant in the uteri of delayed implanting mice and can be reactivated after a transfer to the uterus of ewe (2). We tested whether let-7–enriched EVs would affect the differentiation of a human embryo surrogate model termed BAP-EB (23). BAP-EB was derived by BAP (BMP4, A83-01, and PD173074)–induced differentiation of embryoid bodies of human embryonic stem cells (hESCs) into trophoblast spheroids. BAP-EB spheroids resemble human blastocysts in size and morphology. They express markers of trophoderm and trophoblast and do not express those of other germ layers. BAP-EB selectively attached onto the JEG-3 cells (Fig. 4E) and D4 blastocysts (Fig. S6C). The phosphorylation level of mTORC2 target, AKT at Ser 473 (pAKT), was unexpectedly not affected (Fig. 4E and fig. S6C), suggesting that mTORC1, but not mTORC2, signaling was downstream of C-MYC in blastocysts and trophoblast cells. Overexpression of c-myc mRNA in day 4 let-7g blastocysts for 48 hours reduced the inhibitory effects of DOX-induced let-7g on mTORC1 targets (Fig. 4F).

Inhibition of mTORC1 alone was insufficient to induce embryonic diapause (17). Therefore, bioinformatics analysis was conducted to find whether let-7 also targeted the mTORC2 pathway. TargetScan

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**Fig. 4. Molecular mechanism of let-7–induced diapause.** (A) RT-PCR analysis showing the expression of c-myc and mTOR signaling components after let-7g overexpression. D4 blastocysts from let-7g transgenic mice were cultured in the presence or absence of DOX for 48 hours before determination of gene expression. *P < 0.05. (B) Expression of c-myc in JEG-3 trophoblast cells at 24 hours after transfection of c-myc mRNA. **P < 0.01. (C) Representative confocal images of EdU incorporation (green) in blastocysts at 24 hours after electroporation of c-myc mRNAs or scrambled RNA. Blue, nuclei. Scale bar, 50 μm. (D) Expression of c-Myc and pRpS6 in JEG-3 cells treated with mTOR inhibitor, INK-128, for 24 hours. ***P < 0.001. (E) Western blot analysis of the effect of c-myc inhibitor 10058-F4 (MI) on mTORC1 targets (pRPs6 and p4EBP1) and mTORC2 target (pAKT) in JEG-3 cells. All quantification measurements are normalized to β-actin. **P < 0.01. (F) Representative confocal images of blastocysts immunostained for p4EBP1 and pRpS6. D4 blastocysts from let-7g transgenic mice were electroporated with or without c-myc mRNA before culture in the presence or absence of DOX. ICR D4 blastocysts (D4) and dormant embryos (Dor) served as positive and negative controls, respectively. (G) Expression of Rictor and phosphorylated AKT protein in JEG-3 cells transfected with either let-7a mimics or scramble for 24 hours. Data are presented as means ± SE. *P < 0.05; **P < 0.01. (H) Representative confocal microscope images showing Rictor in embryos during diapause and at 6-hour post–estrogen-induced reactivation (A6h). Scale bar, 50 μm.
normalized to (D) and in blastocysts (E). All quantification measurements in Western blot were ** and ** depicted in the upper panel of (B). Data represent means ± SE. * diapause (Dor-ME). Amplified putative myc response elements (1, 2, 3, and 4-5) are isolated from uterine epithelial cells on day 4 of pregnancy (D4-ME) and during (**). Rictor embryo. Let-7 also suppresses mTORC2 directly by its action on mTORC2 component (pid). During differentiation, the expression of the marker of inner cell mass (OCT4) in the BAP-EB spheroids decreased rapidly at 48-hour pid and was undetectable by 96-hour pid. The expression pattern was different from that of the trophectoderm and trophoblast markers. The trophectoderm marker (CDX2) showed a transient increase at 48-hour pid, while those of tropoblast (CK7, CDH1, and GATA2), syncytiotrophoblast (ERVW-1 and CGB), and extravillous trophoblast (MMP2 and HLA-G) increased progressively with differentiation. Treatment with let-7g-EVs at 48-hour pid significantly affected the mRNA expression of these trophoblast markers at 96-hour pid relative to the control-EV–treated spheroids (Fig. 6A); let-7–treated BAP-EB at 96-hour pid exhibited significantly higher levels of trophoblastic markers (GATA2, CK7, CDH1, ERVW-1, CGB, MMP2, and HLA-G). The comparable expression levels of these markers in the let-7g–EV–treated spheroids at 96-hour pid with that of the spheroids at 48-hour pid were consistent with a high level of let-7–induced dormancy and cessation of differentiation.

BAP-EB spheroids attach specifically onto receptive endometrial cells resembling the early implantation event (23). Treatment with let-7g-EVs reduced the protein expression of c-myc (fig. S8) in the BAP-EB and significantly decreased their attachment onto the receptive endometrial epithelial Ishikawa cells (P < 0.05; Fig. 6B). Good quality human blastocysts can be obtained in 65% of the cultured blastocysts on day 5 and 30% on day 6, but only 5% on day 7 (24). In this study, day 5 human blastocysts (N = 21) were treated with let-7g–enriched EVs and their viability in terms of the presence of a blastocoel, morphology of the trophectoderm cells, and the inner cell mass cells were examined. The blastocoel and good morphology were maintained in 52% of let-7g–EV–treated blastocysts on day 7, whereas only 30% did so in the untreated group (N = 10). One of the EV–treated blastocysts remained viable until day 8 (Fig. 6C). The observations were consistent with a beneficial effect of let-7 on embryo survival in vitro.

**DISCUSSION**

There have long been efforts searching for the natural initiator of embryonic diapause but without much success. Our data show that let-7 of endometrial epithelial origin is a key inducer of embryonic diapause in vivo. Specifically, when the mice undergo diapause, the endometrium generates let-7–enriched EVs, which are taken up by blastocysts. Two observations support the role of let-7 in the induction of embryonic diapause. First, the overexpression of let-7a and incubation with let-7g–enriched EVs prolonged blastocyst survival in vitro. Second, the treated embryos developed to term after an embryo transfer. The study further demonstrates a conserved action of let-7 on the induction of diapause-like phenotype in a human embryo surrogate and the prolongation of survival of human embryos in vitro.

Let-7 is a key inducer of embryonic diapause because it can simultaneously regulate the two known pathways leading to embryonic diapause. Two recent studies showed that simultaneous inhibition of C-MYC and N-MYC (18) or inhibition of mTORC1 and mTORC2 (17) signaling is required for the induction of embryonic diapause. The present data showed that let-7 induced embryonic diapause via the inhibition of both the C-MYC/mTORC1 and mTORC2 signaling pathways. Although the action of let-7 on n-myc has not been studied, it is known that there are two let-7 binding sites in the 3′UTR of Mycn (25). Therefore, it is likely that let-7 also suppresses the expression of n-myc.

Let-7 inhibits Odc1 expression via the suppression of c-myc. This can be a mechanism reducing the potential of diapausining primary receptive endometrial epithelial cells and receptive endometrial epithelial cell lines, but not other nonendometrial cell lines, nonreceptive endometrial cell line, and primary prereceptive endometrial epithelial cells (23).

The time when BAP was added to induce trophectoblast differentiation was considered as time zero of post-induction of differentiation (pid). During differentiation, the expression of the marker of inner cell mass (OCT4) in the BAP-EB spheroids decreased rapidly at 48-hour pid and was undetectable by 96-hour pid. The expression pattern was different from that of the trophectoderm and...
Liu et al., Sci. Adv. 2020; 6 : eaaz7070     9 September 2020

Let-7 overexpression induced dormancy via the suppression of apoptosis, cell proliferation, DNA synthesis, and energy metabolism. The majority of these phenotypes can be explained by the actions of Let-7 on its targets, for instance, a high level of Let-7 targets on c-myc to inhibit proliferation (27), caspase-3 to suppress apoptosis (28), and HK2 to reduce glucose utilization (29). A high Let-7 expression is also associated with the quiescence of fibroblasts (30). Our results on Hk2 and Fbp1 mRNA expression are in contrast with those of a previous proteomic study (31). The discrepancies could possibly be due to different comparisons made in the two studies: dormant versus reactivated blastocysts at 12 to 14 hours after E2 injection in the previous studies but dormant versus D4 activated blastocysts in the present study.

Contrary to the low lactate production of dormant embryos in vivo, an overexpression of Let-7 up-regulated lactate production. Dld and Dlst are Let-7 targets. They are subunits of the 2-oxoglutarate complex and the α-ketoglutarate dehydrogenase complex of carbohydrate metabolism. Inhibition of DLD (Dihydrolipoamide dehydrogenase) activity in spermatozoa causes lactate accumulation (32), and infantile lactic acidosis is associated with severe deficiencies of the α-ketoglutarate dehydrogenase complex (33). These conditions may have been recapitulated in the present model by the suppressive action of Let-7 on Dld and Dlst. Alternatively, the observation could be due to an indirect action of Let-7 on Ldh a via its well-known target Lin28a. Overexpression of Lin28a decreases the expression of Ldh a in the human embryonic kidney cells (34). The molecular mechanism of Lin28a on the observation remains to be determined.

Transcriptomic analyses showed that the mRNA profile of Let-7–induced embryonic diapause is about 30 to 40% similar to that of in vivo dormant embryos. These common genes represent the Let-7–affected genes contributing to in vivo–induced embryonic diapause. Consistently, gene ontology analysis of the common genes showed that they were related to pathways expected to be involved in embryonic diapause. The present study demonstrates that ULF-EVs are important in the induction of embryonic diapause. EVs contain many other components. The lack of some of these components could explain the differential expression of genes between the Let-7–induced and in vivo dormant embryos. The differentially expressed genes may be responsible for the phenotypes that are different between the Let-7–induced and the in vivo embryonic diapause, such as high lactate production after Let-7–induced embryonic diapause. Their absence may also explain the inability of an overexpression of Let-7 alone in maintaining the survival of embryos for a very long term.

The endometrium produces ULF-EVs containing proteins, mRNAs, and microRNAs that are believed to be important means of communication between the blastocysts and the endometrium (9). Let-7 containing EVs inhibited the differentiation of human embryonic surrogates and reduced their attachment onto receptive endometrial epithelial cells. Dysregulation of endometrial microRNAs occurs in subfertile women (35). It is possible that an abnormal expression of embryonic diapause–related microRNAs in the endometrial-derived EVs would retard the differentiation of the implanting embryos, leading to asynchronous development between the embryos and the endometrium. Desynchronization in the development between the blastocyst and the endometrium is a cause of implantation failure (36). However, further studies are required to explore this possibility.

Blastocysts to be reactivated, as polyamine biosynthesis is required for reactivation, and inhibition of their biosynthesis in embryos delays reactivation in vitro (21). Our site-specific ChIP assay demonstrated a reduced recruitment of C-MYC to the Odc1 promoter during diapause. Whether the reduction is due to the suppression of C-MYC expression resulting from high expression of Let-7 in the cells during diapause remains to be determined.

We localized the C-MYC protein mainly to the cytoplasm of embryos. This differs from the expected nuclear localization of C-MYC in many cell types. The antibody used in this report localized the expression of C-MYC to the nuclei of embryonic stem cells (fig. S6D). Three other anti–C-MYC antibodies were tested and showed similar cytoplasmic C-MYC expression in embryos (data not shown). The expression of cytoplasmic MYC has been reported. MYC-nick is a cytoplasmic cleavage product of the full-length C-MYC, widely expressed in a large number of cell lines (26). It is expressed in differentiating cells and tissues and plays a significant role in the differentiation of myofibroblasts and the transdifferentiation of fibroblasts into muscle cells (26). Thus, a Let-7–induced decrease of cytoplasmic C-MYC may lead to a decrease in the differentiation potential of the diapauing blastocysts.
In conclusion, the study showed an important role of endometrial EVs in embryo dormancy, demonstrating that let-7 in EVs is a major player in the induction of embryonic diapause.

MATERIALS AND METHODS
A summary of the techniques and procedures for addressing the questions raised in the report can be found in the Supplementary Materials.

Use of animals
The study protocol was approved by the Committee on the Use of Live Animals in Teaching and Research, The University of Hong Kong (CULATR number: 3560-15). Females of the ICR mice were mainly used. Transgenic mice carrying a DOX-inducible let-7g gene (Slet-7gLmiR-21) were generated by the injection of embryonic stem cells carrying the let-7gStem/21loop sequence (a gift from G. Daley, Harvard Stem Cell Institute, Boston, MA, USA) into ICR blastocysts. The chimeric mice generated were then mated with CD-1 females to generate germline-transmitted pups. To match the genetic background of the embryonic stem cells carrying the transgene (V6.5 mESC), the mouse line was maintained on a C57/B6 background by backcrossing more than five times.

Collection of mouse embryos
ICR female mice aged 6 to 8 weeks were superovulated by successive intraperitoneal injections of 5 IU of pregnant mare serum gonadotropin (Sigma-Aldrich, St. Louis, USA) and 5 IU of human chorionic gonadotropin (hCG; Sigma-Aldrich) 47 to 48 hours apart and were mated with male mice. The vaginal plug was checked the day following mating. The day when a vaginal plug was seen was considered as day 1 of pregnancy. Blastocysts were collected from the uteri at 96 hours after hCG injection and cultured in M16 medium (Sigma-Aldrich) supplemented with 10% FBS (Thermo Fisher Scientific, CA, USA) or KSOM + AA medium (Millipore, MA, USA) into ICR blastocysts. The pure epithelial cells and stromal cells was over 90% as determined by immunostaining, using antibodies against mouse cytokeratin (Dako, Glostrup, Denmark) and mouse CD90 (BD Biosciences, MA, USA), respectively.

Manipulation of let-7 levels in embryos
Electroporation
Pre-let-7a or scrambled miRNA control (Thermo Fisher Scientific) was electroporated into day 4 blastocysts from ICR mice as described (7). Briefly, the uteri of day 4 pregnant mice were flushed with Hank’s solution to obtain blastocysts, which were then transferred to prewarmed droplets of M16 medium. Pre-let-7a or scramble control was electroporated into the embryos in a flat electrode chamber (1-mm gap between electrodes; BTX Inc., San Diego, USA) in 20 μl of Hepes-buffered saline (150 mM NaCl, 20 mM Hepes, Sigma-Aldrich), by two sets of three electric pulses of 1 ms at 30 V with 1-min interval between sets, and inverting polarity using the 830 Electro Square Porator (BTX Inc., San Diego, USA). Following electroporation, the embryos were cultured in KSOR + AA or M16 for experimentation. About 95% of the electroporated embryos survived the process; they showed no sign of cell lysis at 2 hours after electroporation.

Treatment with let-7–enriched EV
Ishikawa cells were transfected with pre-let-7a or pre-miR scramble using Lipofectamine 2000 (Thermo Fisher Scientific). After transfection, the transfection medium was replaced by fresh MEM medium supplemented with 1% penicillin/streptomycin, 1% L-glutamine, and 10% EV-depleted FBS (Thermo Fisher Scientific, CA, USA). The spent medium from 48 hours of culture was collected for EV isolation. EVs were isolated from the spent medium with the Total Exosome Isolation Kit (Thermo Fisher Scientific) according to the manufacturer’s instruction. The let-7a levels in the EVs were detected by reverse transcription qPCR (RT- qPCR) (let-7a primer, Thermo Fisher Scientific). Alternatively, let-7g–enriched EVs were isolated from the spent medium after culture of DOX-treated endometrial epithelial cells from the let-7g transgenic mice in medium supplemented with 10% EV-depleted FBS for 48 hours. The epithelial cells with a purity of more than 90% were isolated as above. The protein concentration of the EV preparation was determined with the BCA Protein Assay Kit (Thermo Fisher Scientific) with a working concentration range of 5 to 2000 μg/mL. The let-7–enriched EVs for coincubation with embryos.
**Isolation of EVs from ULF**

Pregnant mice or delayed implanting mice at 3, 6, and 24 hours after E2 reactivation were sacrificed by an overdose of pentobarbital (150 to 200 mg/kg, intraperitoneally). Their uteri were isolated, and ULF was collected by flushing the uterine lumen with 500 μl of PBS. The embryos in the flushing were removed under a dissection microscopy. EVs in the ULF were isolated with the Total Exosome Isolation Kit. Briefly, the ULF was centrifuged successively at 300g for 10 min to remove cells, at 2000g for 10 min to remove dead cells, and at 20,000g for 60 min to remove cell debris and large vesicles. The pellets that formed in each centrifugation were discarded. After the last centrifugation, the total exosome isolation reagent (250-μl volumes) was mixed with the supernatant overnight at 4°C on a roller mixer, before the samples were centrifuged at 10,000 g for 60 min at 4°C. The supernatant was discarded, and the ULF-EV pellet was gently washed once with 200 μl of PBS to remove residual extract before the samples were centrifuged at 10,000 g for 60 min to remove cell debris and large vesicles. The pellets in the ULF were isolated with the Total Exosome Isolation Kit. Briefly, the ULF was centrifuged successively at 300g for 10 min to remove cells, at 2000g for 10 min to remove dead cells, and at 20,000g for 60 min to remove cell debris and large vesicles. The pellets that formed in each centrifugation were discarded. After the last centrifugation, the total exosome isolation reagent (250-μl volumes) was mixed with the supernatant overnight at 4°C on a roller mixer, before the samples were centrifuged at 10,000 g for 60 min at 4°C. The supernatant was discarded, and the ULF-EV pellet was gently washed once with 200 μl of PBS to remove residual extract buffer and resuspended in 20 μl of PBS and stored at −80°C. The μl of PBS was added to remove residual extract gently washed once with 200 μl of PBS. The pellets in the supernatant were collected by flushing the uterine lumen with 500 μl of PBS. The embryos in the flushing were removed under a dissection microscopy.

**EdU incorporation assay**

DNA synthesis was determined by the EdU incorporation assay (Thermo Fisher Scientific) according to the manufacturer’s instruction. Briefly, embryos were cultured in KSOM medium containing 10 μM EdU for 30 min before washing with PBS. After the removal of the zona pellucida, the embryos were fixed in methanol at −20°C for 20 min and permeated in PBS containing 1% bovine serum albumin (BSA) and 0.5% fluorescein azide for 30 min. The staining mixture was prepared fresh each time. The embryos were washed three times with PBS containing 0.05% Tween 20 before the fluorescence signal was visualized under a confocal microscope (Carl Zeiss LSM 700, Zeiss, Germany).

**Immunofluorescence staining of embryos**

The embryos were washed with M2 medium (Sigma-Aldrich) and fixed in 4% paraformaldehyde for 15 min at room temperature. They were permeabilized with 0.1% Triton X-100 in Dulbecco’s PBS (DPBS) for 30 min, and then incubated with TUNEL reaction mixture containing 5 μl of enzyme solution and 45 μl of label solution at 37°C for 60 min. The embryos were washed three times with PBS. Their nuclei were stained with propidium iodide (Sigma-Aldrich) for 5 min before the embryos were mounted on microscope slides for examination under a fluorescence-inverted microscope (TE300; Nikon, Japan).

**Metabolite analysis**

The method used was a miniaturized version of conventional enzymatic methods, which rely on the detection of ultraviolet (UV)–excited NADH (reduced form of nicotinamide adenine dinucleotide) and NADPH (reduced form of nicotinamide adenine dinucleotide phosphate) in enzyme coupled reactions (38). Instead of having the enzymatic reactions conducted in cuvettes, they were done in 20-μl droplets on a petri dish. The specific enzyme cocktails for the metabolite studied were as follows: glucose cocktail: 42 mM EPP S (4-(2-Hydroxyethyl)-1-piperazinepropanesulfonic acid) buffer (pH 8.0), 42 μM dithiothreitol, 3 mM MgSO4·7H2O, 0.42 mM ATP, 1.2 mM NADP, and hexokinase (14 U/ml)/glucose-6-phosphate dehydrogenase (7 U/ml; Roche Applied Science); pyruvate cocktail: 63 mM EPP buffer (pH 8.0), 0.1 mM NADH, and l-lactate dehydrogenase (75 U/ml; Roche Applied Science); lactate cocktail: 0.45 M glycine/0.73 M hydrazine buffer, 4.5 mM NAD, and l-lactate dehydrogenase (69 U/ml; Roche Applied Science). The cocktail droplet (2 μl) was mixed with 18 μl of spent culture media. Following a 3-min incubation at room temperature, 5 μl of the medium was transferred to a homemade chamber with a chamber depth of 1 mm, and its fluorescence intensity was determined under a fluorescence microscope. The fluorescence signal relative to the background was determined by the pixel intensities using the Image-Pro Plus 6.0 software (Media Cybernetics Inc., Silver Spring, MD, USA). The background signal was estimated using a method described previously (39). The changes in fluorescence were converted to changes in the concentration based on standard curves performed on the same day with known concentrations of the appropriate substrates.

**Western blot analysis**

Proteins were separated by SDS–polyacrylamide gel electrophoresis, transferred onto polyvinylidene difluoride membranes (Merck Millipore, Germany), and probed with antibodies against c-myc (Cell Signaling Technology, MA, USA), ODC1, SMS, pAKT, p4EBP1, pRpS6, and the total protein of AKT, 4EBP1, and RpS6 (Cell Signaling Technology). β-Actin (Sigma-Aldrich) was used as the internal control. The membranes were incubated with the WesternBright ECL Kit (Advansta, CA, USA) and exposed to x-ray films.
Extraction of total RNA from embryos

Embryos at the same developmental stage were randomly pooled into three groups with five embryos per group. Total RNA was extracted from each group in 0.5 μl of 2 M guanidine isothiocyanate (Sigma-Aldrich) buffer at room temperature for 5 min. Complete lysis of the embryos in the buffer was confirmed under a microscope. The samples were diluted to 5 μl with double-distilled water and were used directly for the multiplex microRNA assays.

Precursor-microRNA primer design for Slet-7gLmiR-21

To amplify the precursor-microRNA, the forward and reverse primers were designed to anneal to the stem portion of the hairpin. The forward primer was designed for a sequence that crossed the stem and the loop of the precursor of Slet-7gLmiR-21 so that only its stem, but not that of the pre-let-7g, was amplified. The sequences of the primers used for pre-S7gL21 were as follows: forward: 5′-GTAG-TAGTTGTGACATGCTG-3′; reverse: 5′-TAATCTGCAGCAAG-GC-3′; probe: 5′-CTGTACAGTCCATGAGATT-3′; Rt reverse: 5′-TAAATCTCTGGCAAGGCA-3′.

Structure of Slet-7gLmiR-21 in the transgenic mouse

![Structure of Slet-7gLmiR-21 in the transgenic mouse](image)

Precursor-microRNA detection in mouse embryos

Embryos were lysed in 5% NP-40 to release the total RNA. RT primer annealing was performed at 85°C for 5 min by adding 1 pmol of RT reverse primer to 1 μg of the above RNA. Then, the samples were placed immediately in ice to avoid the formation of stem-and-loop structure. RT was performed at 45°C for 60 min and at 85°C for 5 min and then kept at 4°C in a thermal cycler (T100 Thermal Cycler, Bio-Rad, CA, USA). RT-qPCR was performed with the 7500 Real-Time PCR System (Thermo Fisher Scientific). The program was a 10-min cycle at 95°C, followed by 45 cycles of 15 s at 95°C and 60 s at 60°C.

Microarray

Microarray (GeneChip™ Mouse Gene 2.0 ST) was used to study the effect of let-7 overexpression on the transcriptome of the treated blastocysts. For each sample, RNA was collected from 10 blastocysts. RNA extraction, amplification, and purification were performed according to Kurimoto et al. (40). The complementary DNAs (cDNAs) from day 7 dormant blastocysts, D4-act, and DOX-induced blastocysts were hybridized to the GeneChip™ Mouse Gene 2.0 ST array in duplicates (Affymetrix, CA, USA). All cDNA hybridizations were performed by the Centre for Genomic Sciences, The University of Hong Kong. The microarray data have been deposited to the Gene Expression Omnibus (GSE141900). The data were analyzed using Partek Genomics Suite 6.6 (St. Louis, MO, USA). The expression matrix was further subjected to the R package linear models for microarray data (41) for identifying the differentially expressed genes. Principal components were computed and plotted with the R packages FactoMineR and factoextra. Heatmaps were plotted with the R package gplots using z scores calculated for each gene across different samples. Biological process analysis was performed by DAVID (v6.8) (42).

Preparation of c-myc mRNA

To obtain the DNA template for in vitro transcription, the pcDNA 3.1_cMyc plasmid containing the coding DNA sequence was PCR-amplified using the following primers: 5′-TAATACGACTCACTATAGATGCCCTCAACGTGAAC-3′ (with T7 polymerase promoter) and 5′-TTATGCACAGAGTTTCTGGAAGC-3′. The product was purified using the GeneJET PCR Purification Kit (Thermo Fisher Scientific). The MEGAscript T7 ULTRA Transcription Kit (Thermo Fisher Scientific) was used for in vitro transcription from DNA to mRNA according to the manufacturer’s instructions. Briefly, DNA template was recovered at a final concentration of 1 μg/μl. Transcription reaction was performed by incubation at 37°C for 4 hours. TURBO DNase (1 μl) was added into the reaction and incubated for 15 min at 37°C before the addition of the tailing reagents for poly(A) tailing. Last, RNA was recovered using phenol:chloroform extraction and isopropanol precipitation. The recovered RNA was then quantified and stored at −80°C and was ready for transfection.

Dual luciferase assay

The mouse genomic DNA was extracted from the ICR mouse liver using a DNA extraction kit (Thermo Fisher Scientific) according to the manufacturer’s protocol. The 3′ UTR of Rictor was amplified with the Not I and Xho I digestion sites by the Phusion High Fidelity DNA Polymerase (New England Biolabs, Beverly, MA, USA). The PCR products were first purified with the GeneJET PCR Purification Kit (Thermo Fisher Scientific), digested with the Not I and Xho I enzymes (New England Biolabs), and purified with the GeneJET PCR Purification Kit again. The purified PCR products were ligated with the Not I- and Xho I-digested psiCHECK-2 vector (Promega, WI, USA). Lipofectamine 2000 transfection reagent (Thermo Fisher Scientific) was used to cotransfect 1 μg of the WT and mutant (Mut) reporter constructs with 5 nM let-7a mimic (Thermo Fisher Scientific) into a monolayer of JEG-3 cells at 70% confluence in Opti-MEM (Thermo Fisher Scientific). At 24 hours after transfection, the cells were lysed in 100 μl of 1× passive lysis buffer (Promega). The luciferase assays were performed using a luciferase assay kit (Promega) according to the manufacturer’s protocol and were measured using a luminometer (GloMax 96 Microplate Luminometer, Promega). Renilla luciferase was used for normalization.

ChIP assay

ChIP analysis was performed using the Pierce Chromatin Prep Module (Thermo Fisher Scientific, 26158) according to the manufacturer’s instruction. Uterine epithelium cells from delayed implanting and activated mice were isolated as described above. Formaldehyde was used to cross-link DNA and its interacting proteins in the cells. The cells were then lysed in the Lysis Buffer on ice for 10 min and centrifuged. The supernatant was discarded, while the nuclei were
resuspended in the MNase Digestion Buffer. Micrococcal nuclease was added to digest the chromatin. Immunoprecipitation was performed with the ChIP-grade c-Myc antibody (Cell Signaling Technology) at a dilution of 1:100. Proteinase K was used to disrupt the cross-links between the DNA and proteins. The DNA was then purified using the PCR Cleanup Extraction Kit (Thermo Fisher Scientific), after which quantitative RT-PCR was performed with the SYBR Green Master Mix (Thermo Fisher Scientific) using the following ChIP primer sequences (Table 1).

### EGF binding assay

Blastocysts were incubated for 10 min at 37°C in an atmosphere of 5% CO₂ in 40-μl microdrops of M16 containing Alexa Fluor 488–labeled EGF (2 μg/ml; Thermo Fisher Scientific). Unlabeled EGF peptide at a concentration of 20 μg/ml was used as control for nonspecific binding of the labeled peptide. After termination of the incubation, blastocysts were washed in medium and fixed in 4% paraformaldehyde in PBS for 15 min at 4°C. Z-stack images of fixed embryos were captured with a confocal fluorescence microscope (Carl Zeiss LSM700, Germany). For quantification of the EGF binding, the images of the embryos were analyzed with the ImageJ software (1.52p, USA). The average fluorescence intensity was calculated. The data presented were the averages of the fluorescence intensity from at least three embryos.

### Human trophoblastic spheroid (BAP-EB) formation and attachment assay

BAP-EB spheroids were generated from hESCs as described (23). Briefly, hESCs (VAL3, Spanish Stem Cell Bank, Spain) were digested to single cells with accutase (Thermo Fisher Scientific, Waltham, USA) and aggregated in AggreWellTM400 (STEMCELL Technologies Inc., Canada) in mTeSR™1 medium (STEMCELL Technologies Inc., Canada) for 24 hours before the induction of trophoblast differentiation in BAP medium (mouse embryonic fibroblast–conditioned medium supplemented with BMP4 (10 ng/ml; R&D Systems, Minneapolis, USA), 1 μM A83-01 (Stemgent, San Diego, USA), and 0.1 μM PD173074 (Stemgent). The medium was changed daily during a 96-hour differentiation.

Control EVs or let7–enriched EVs were added to the BAP-EB culture at 48-hour pid. For the attachment assay, BAP-EB at 72-hour pid was transferred onto a confluent monolayer of Ishikawa cells and cocultured for 3 hours. Nonadherent spheroids were removed, and the percentage of attached BAP-EB was determined. For the gene expression analyses, BAP-EB at 0 and 48 hours after EV treatments and BAP-EB at 96 hours after EV treatments were collected and subjected to total RNA extraction and real-time quantification of marker expressions, as described (23).

### Human embryos

Human embryos were obtained from infertile couples attending the assisted reproduction clinics at the Department of Obstetrics and Gynecology, General Hospital of Chinese People’s Liberation Army and the Center for Reproductive Medicine, The Third Affiliated Hospital, Sun Yat-Sen University. The Institutional Review Board of the Hospital approved the project (S2017-095-01), and written consent was obtained from each donor. The embryos were donated because the donor couples had completed their family (N = 17), or the embryos were chromosomal abnormal as determined by pre-implantation genetic testing for aneuploidy (N = 4). The donated embryos were cryopreserved on day 5 before experimentation.

EVs were obtained from endometrial cells of let-7g transgenic mice after treatment with DOX in DMEM/F12 medium supplemented with EV-free FBS for 4 days. Control EVs were obtained from cells without DOX treatment. On the day of experimentation, the donated blastocysts were thawed and cultured in G2 medium (Vitrolife, Sweden) supplemented with let-7g–enriched EVs or control EVs until they were morphologically not viable.

### Data analysis

All the results are shown as means ± SEM. All the data were analyzed using one-way analysis of variance (ANOVA). A P value of less than 0.05 was considered statistically significant.

### SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/6/37/eaa7070/DC1

View/request a protocol for this paper from Bio-protocol.

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