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Studies reporting term pregnancy and the production of genetically identical offspring from isolated blastomeres of early stage embryos have been carried out in small and large animals. However, very little is known about the effects of embryo splitting on the development and reproductive competency of human embryos. Here, we investigated the effects of embryo splitting on profile of miRNAs detected in their spent blastocyst medium (SBM) by comparative analysis of miRNA profiles in SBM of human twin embryos created by blastomere biopsy and SBM of blastocysts that resulted in a healthy pregnancy and live birth following embryo transfer. The profile of miRNA secretion in in vitro culture media consistently distinguishes twin from control embryos. We found that six miRNAs are significantly more abundant in SBM from twin embryos, while nine are significantly more abundant in SBM from euploid implanted blastocysts. These nine include miRNA-30c, a previously reported marker of blastocyst implantation potential. Furthermore, 22.9% of miRNAs secreted by twin embryos were never detected in SBM from normal, reproductively competent blastocysts, or from TE samples from normal blastocysts donated for the research. The miRNA profile, unique to twin blastocysts, might be a result of differential lineage commitment in these embryos.
Human embryos created by embryo splitting secrete significantly lower levels of miRNA-30c

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Short title: miRNOme in human split embryos

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Abstract

Studies reporting term pregnancy and the production of genetically identical offspring from isolated blastomeres of early stage embryos have been carried out in small and large animals. However, very little is known about the effects of embryo splitting on the development and reproductive competency of human embryos. Here, we investigated the effects of embryo splitting on profile of miRNAs detected in their spent blastocyst medium (SBM) by comparative analysis of miRNA profiles in SBM of human twin embryos created by blastomere biopsy and SBM of blastocysts that resulted in a healthy pregnancy and live birth following embryo transfer. The profile of miRNA secretion in in vitro culture media consistently distinguishes twin from control embryos. We found that six miRNAs are significantly more abundant in SBM from twin embryos, while nine are significantly more abundant in SBM from euploid implanted blastocysts. These nine include miRNA-30c, a previously reported marker of blastocyst implantation potential. Furthermore, 22.9% of miRNAs secreted by twin embryos were never detected in SBM from normal, reproductively competent blastocysts, or from TE samples from normal blastocysts donated for the research. The miRNA profile, unique to twin blastocysts, might be a result of differential lineage commitment in these embryos.

KEYWORDS: human preimplantation embryos; embryo splitting; spent blastocysts medium; miRNA; miR-30c
Introduction

miRNAs are evolutionarily conserved, single-stranded, non-coding RNA molecules of ~22 nucleotides in length and are considered to be major transcriptional/post-transcriptional regulators of gene expression. In the human genome over 1,000 miRNAs have been identified (miRBase, www.mirbase.org). Human blastocysts express large number of miRNAs [1]. They are secreted and can be found in embryo culture media. Attempts have been made to link specific miRNAs detected in the medium with embryo ploidy status and reproductive competence in order to predict in vitro fertilization outcome, and several candidates have been identified [2-5]. Although further studies are warranted, a comprehensive profile of miRNAs secreted by human preimplantation embryos in spent culture media is taking shape [6].

Several studies have suggested that the splitting of human embryos might result in morphologically adequate, viable blastocysts [7, 8]. However, qualitative analyses of the embryos created in such a way have been relatively limited. We recently analysed 176 twin embryos created by blastomere separation of 88 human embryos from either early (2-5 blastomeres, n=43) or late (6-10 blastomeres, n=45) cleavage stage. Half of the blastomeres in each embryo were biopsied and placed into an empty zona pellucida (ZP) prepared in advance by removing the cellular content from immature oocytes or degraded, clinically unsuitable embryos. The blastomere donor embryos were dubbed twin A and the blastomere recipients twin B [9]. We found that they have distinctive features in comparison with normal blastocysts obtained through fertilization. Twin split embryo blastocysts were smaller and less cellular, the first fate decision was somewhat delayed, inner cell mass (ICM) was either absent or small and poor quality and the majority of cells were expressing ICM and trophectoderm (TE) markers simultaneously (Fig. 1).

To address whether such changes in the phenotype of embryos generated by embryo splitting affect miRNA secretion, we compared miRNA profile in spent blastocyst medium (SBM) of seven pair of twins and seven blastocysts generated by in vitro fertilization. All control blastocysts resulted in live births upon single embryo transfer.
Material and methods

The work described here is done under licence from the UK Human Fertilisation and Embryology Authority (research licence numbers: R0075 and R0133) and also has local ethical approval (UK National Health Service Research Ethics Committee Reference: 06/Q0702/90). Informed consent was obtained from all embryo donors and the experiments conformed to the principles set out in the WMA Declaration of Helsinki and the NIH Belmont Report. No financial inducements are offered for donation. All the embryos used in this project were cryopreserved with slow freezing method.

Embryo culture

Embryo culture and splitting using the blastomere biopsy technique has been described in detail previously \[9\]. All the embryos were split at the late cleavage stage (6-9 cells). Half of the blastomeres were aspirated and removed from a donor embryo and placed in an empty human zona pellucida, previously prepared from clinically unsuitable oocytes or embryos. After blastomere transfer, the donor (Twins A, n=7) and recipient embryos (Twins B, n=7) were cultured up to the blastocyst stage (day 5/6). SBM was collected as described \[2\]. Briefly, the embryos were cultured from Day 3 post-fertilization in a 35-µl drop of Quinn's Advantage Blastocyst Medium (Cooper Surgical) with 5% Quinn's Advantage Human Serum Albumin (Cooper Surgical) in a humidified atmosphere containing 5% O2 and 6% CO2. At the blastocyst stage, prior to transfer, 25 µl of SBM was taken for analyses. As a control, we used SBM of embryos that resulted in a healthy pregnancy and live birth following embryo transfer (n=7) (Fig. 2). “No Template” controls and unconditioned blank culture media were also run in parallel in order to exclude any possible false positive amplification from the subsequent analyses.
**miRNA isolation, profiling and screening**

miRNA isolation and profiling were described in details previously [2]. Briefly, miRNAs were isolated through a magnetic beads-based extraction (Anti-miRNA Bead Capture purification kit human panel A, Life Technologies, USA), and after targeted retrotranscription and pre-amplification steps using the Megaplex RT and Preamp human panel A primers pool (Life Technologies), samples were analysed in qPCR through the Taqman Low Density Array (TLDA) cards on a Viia7 instrument (Life Technologies). Data processing and statistical analyses were performed through the RealTime StatMiner software (Integromics, Spain) and the SPSS software (IBM, USA). Data were compared also with miRNAs profiled from 5 ICM-free TE samples obtained from normal blastocysts donated for the research, as previously described [2]. The primer sequences for targets within the human panel A+B are listed on the Thermo Fisher’s website

[http://tools.thermofisher.com/content/sfs/manuals/4473439C.pdf](http://tools.thermofisher.com/content/sfs/manuals/4473439C.pdf)

GenoSplice Technology performed quality control (QC), processing, and further analyses of the data using endogenous control for normalization. For each miRNA X of each sample S, the samples were normalized according to the formula $2^{-\Delta Ct_{X,S}} = 2^{(Ct_{X,S} - \text{mean}(Ct_{U6,S}))}$; samples 6A and 6B were omitted.

**Single assays**

Eluted left-over SBMs samples from twins A, twins B and randomly selected historical controls of euploid implanted blastocysts, respectively, were processed for single assays’ analyses through qPCR. Blank (molecular biology grade water) and media samples never exposed to embryos were processed in parallel as negative controls. We tested single assays for the U6-snRNA, miR-30c and miR-203. The first was chosen in order to confirm its weak reliability as detected through the miRNome panel, the second in order to confirm the differential expression in the comparison of twins A versus twins B, while the last miRNA was chosen as potential
normalizer since it showed a stable trend in the panels’ analysis among all these samples (Twins A Ct 26.3±1.1, 24.1-27.4; Twins B Ct 25.8±1.2, 23.5-27.3; euploid implanted blastocysts 24.1±1.1, 22.9-25.5). The minimum variance of the median method performed through the StatMiner software (Integromics) confirmed its eligibility to this aim.

The primers for retrotranscription reaction of the three assays were pooled together (5µl each) and diluted through 485µl of nuclease-free water. The retrotranscription mix was composed as follows: 6µl of primers pool, 0.3µl of 100mM dNTPs (with dTTP), 3µl of MultiScribe™ Reverse Transcriptase 50 U/µL, 1.5µl of 10X Reverse Transcription Buffer, 0.2µl of RNase Inhibitor 20 U/µL, 1.5µl of nuclease-free water (TaqMan® Small RNA Assays protocol, Life Technologies) and 2.5µl of eluted left-over sample. The thermal protocol was conducted on a 2720 Thermal Cycler (Life Technologies) as follows: 16°C for 30’, 42°C for 30’, 85°C for 5’ and 4°C ∞. No preamplification step is entailed by this protocol and thus we raised the threshold Ct level for detection to 37 cycles. qPCR reaction mix was composed as follows: 0.5µl of TaqMan® Small RNA Assay (20X), 5µl of TaqMan® Universal PCR Master Mix II (2X) no UNG, 2µl of nuclease-free water and 2.5µl of sample. qPCR was performed on a ViiA7 machine (Life Technologies) according to manufacturer’s protocol. Each assay was run in triplicate in order to exclude the technical variability due to pipetting error.

The primer sequences for single essays are: hsa-miR-30c-5p (cat # 000419) target sequence UGUAACAUCCUACACUCUCAGC, u6 snRNA (cat # 001973) target sequence GUGCUCGCUUCGGCAGCACAUAUCUAAAUUGGAACGATACAGAGAAGAUUAGCAUGGCC CCUGCGCAAGGAUGACACGCAAAUUGCUGAAGCGUUCCAUAUUUU, and hsa-miR-203 (cat # 000507) target sequence: GUGAAUGUUAGGACCACUAG.

The Ct values for miR-30c were normalized on miR-203 ones and the differential expression analyses was conducted through the ΔΔCt method.
Results

Using a medium throughput analysis, we compared the profile of 377 miRNA sequences from the SBM of twin embryos (n=7 pairs; Supplemental figures S1 and S2) with those from control embryos that resulted in live birth after single embryo transfer (n=7). Correlation between samples of each group was high for all conditions (Fig. 3A). Correlation between samples in the control group was somewhat lower than between the samples in Twin A and Twin B groups (Fig. 3B). Pearson’s correlation matrix of the raw Ct values consistently distinguished control embryo from twin embryos (Fig. 3C). Heat map revealed that there were no significant clusters by experimental group; however, there were high numbers of missing Ct values (Fig. 3D).

To minimize experimental error, we analysed only miRNA with at least 60% of Ct values above the threshold that was set as 35 in each compared experimental group (namely 5/7 valid values for control SBMs and 9/14 valid values for twin SBM).

Data analysis using common intracellular normalizers

Commonly used intracellular normalizers, such as small nuclear RNA U6, often show large fluctuations in samples measuring secreted miRNAs [10]. In our case, after quality control analyses, only one out of three endogenous controls was available (U6). There were no Ct values for endogenous control small nucleolar RNA RNU44 and only eight values for RNU48. Despite Ct of U6 endogenous control varying across the samples, Ct values of U6 were homogenous within each sample (Fig. 4A). Since the four different U6 reporters provided approximately the same Ct values for the same sample, U6 can be used for computing - ΔCt normalization of the other miRNAs. However, even though reproducibility across the samples was good, due to very low U6 Ct values in the sample 6B, we excluded the twins 6A and 6B from the analyses (Fig. 4B, C).

We found that miR-515-5p and miR-490 were detected at significantly higher levels, whereas miR-486-3p, miR-30c and miR-509-3-5p had significantly lower concentrations in twins in
comparison with control embryos (FC ≥ 1.5, p-value ≤ 0.05) (Fig. 4D). However, a lack of two internal reference genes (RNU44 and RNU 48) and large fluctuations in the third one (U6) for secreted miRNAs prompted us to approach data analysis using a different type of normalization: the global normalization strategy [11].

Data analysis using global geometric mean of expression of all detected miRNAs

Global normalization strategy consisted of three successive steps. Any Ct value above the threshold set at 35 was considered as a noise and was discarded from further analysis. The arithmetic average Ct value for all detected miRNAs was then calculated for each individual sample and subsequently subtracted from each individual Ct value for that sample. The procedure results in normalized expression values in logE scale (E being the base of the exponential amplification function, with 2 being a good estimate); for individual miRNAs, values are inversely correlated with expression levels. Whole-genome RT-qPCR based miRNA profiling in combination with a global mean normalization strategy has proven to be the most sensitive and accurate approach for high throughput miRNA profiling, whose effectiveness when applied to low input samples as well as SBMs has been validated and reported previously [2].

MicroRNA profile in SBMs is similar among twins A and B, but significantly different from miRNA profile of control embryos

Seven pairs of SBMs from twin embryos were compared to SBMs from reproductively competent blastocysts. After all QC steps, 48 miRNAs were found to be consistently secreted in IVF culture media by twin A and B blastocysts, while 59 by control ones. The twins A and B shared 46 out of 48 miRNA detected in their SBM. miRNA detected only in SBM of twin A was miR-193b,
whereas miR-636 was detected only in SBM of twin B (Fig. 5A). Differences between miRNA profiles in SBM of twins A vs. control blastocyst were overlapping with differences of profiles in SBM of twins B vs. control blastocysts (Fig. 5B).

Twenty-two miRNAs were shared between SBM of control and twin embryos (Fig. 6A). Thirty-seven miRNAs were detected solely in SBM of controls, and 26 only from twin and never secreted by non-manipulated human blastocysts generated by in vitro fertilization. All 59 miRNAs detected in SBM of control blastocyst were also detected in ICM-free TE cells [2]. However, only 53.8% (14/26) of the twin-specific secreted miRNAs were co-expressed from normal blastocyst-derived TE cells, whereas 12 were not detected in TE cells previously. Among these 12, only one, miR-374-5p, was expressed in ICM (A. Capalbo, unpublished data). The detection of those novel 12 miRNAs may suggest abnormal blastocyst development and/or be an indicator of different lineage commitment stage in the split embryos.

A relative quantification analysis was conducted and a volcano plot was generated by comparing the detection levels in SBM from twin versus controls of the 22 common miRNAs (Fig. 6B). Six miRNAs were found to be significantly more abundant in the twin samples (miR-203, miR-136, miR-490, miR-758, miR-222 and miR-523), whereas nine miRNAs were more abundant in the control samples (miR-193b, miR-30b, miR-106b, miR-30c, miR-373, miR-24, miR-590-5p, miR-25 and miR-27b).

The results were further validated with single assays specific for miR-30c and for miR-203. Twin A mean Ct±SD were 34.9±1.9 and 31.3±1.1 for miR-30c and miR-203, respectively. Twin B mean Ct±SD were 35.3±2.2 and 30.8±1.2 for miR-30c and miR-203, respectively. The ΔΔCt analyses showed no difference in the levels of miR-30c normalized on miR-203 in twins A versus B (fold-change: 1.9; p-value: 0.7). When performing the same analysis in twins (A and B together) versus the historical control of SBMs from euploid implanted blastocysts, a statistically significant 93.2X lower level was detected (p<0.001) in the formers. No miRNA expression was found in negative controls.
Significantly lower expression levels of miR-30c in twins was detected with either of two normalization strategies

A statistically significant lower level of miRNA-30c detected in SBM from twin embryos culture than in SBM of control embryos was identified using either of two normalisation strategies. It captured our interest, because this miRNA has been identified as a putative biomarker of blastocyst implantation potential when secreted at high levels in the SBM during IVF cycles [2].

Next, using a computational analysis, we investigated possible roles of miRNA-30c during early development. According to the miRNA-gene interactions data provided by the DIANA TarBase v7 (http://diana.cslab.ece.ntua.gr/tarbase), miRNA-30c has 1643 putative gene targets. To identify which of them are expressed in TE and/or ICM of human blastocyst, we filtered these genes against our blastocyst transcriptome database, which contains a list of the genes detected by the RNAseq method in isolated TE and ICM samples [12]. We found that 1061 of these genes are expressed in ICM, 1166 in TE and 1006 in both ICM and TE (Fig. 6B). Based on the pathway information provided by the Kyoto Encyclopedia of Genes and Genomes (KEGG; www.genome.jp/kegg), we found that the main pathways retrieved were ubiquitin mediated proteolysis and spliceosome (Fig. 6C).

Discussion

miRNAs act mostly as post-transcriptional repressors of their target genes. The importance of miRNAs in early embryo development has been identified in a range of species, though relatively little is known about the miRNA regulatory network in mammalian preimplantation embryos [1, 13-17]. Aberrant miRNA profiles in transferable blastocysts have been linked with male factor infertility and polycystic ovaries [5]. Since miRNAs are known to be secreted [18], attempts have been made to link specific miRNAs detected in the SBM with embryo ploidy status and reproductive competence [2-5].
miRNAs detected in the spent medium from early embryos might be the output of paracrine circuits, reflecting normal communication among cells of healthy embryos; alternatively, they might be a component of the communication pathway between the embryo and the uterine epithelium, influencing epithelial readiness for embryo apposition and implantation. However, it remains possible that the miRNAs detected in the SBM are products of degrading cells.

Significantly lower levels of miR-30c in twins perhaps indicate poor developmental prognosis for the twin embryos. miR-30c is known to interfere with epithelial-to-mesenchymal transition (EMT) in breast cancer through regulation of TWF1 and IL11 [19], and has been identified as a key indicator of reproductive competence [2]. EMT is an essential part of the first lineage fate decision when during asymmetric cell division one of the daughter cells is pushed inwards, losing polarity and forming ICM [20, 21]. Whether miR-30c is directly involved in the ICM formation or is an essential part of the embryo/maternal dialogue at the time of implantation remains to be investigated.

Most of the miRNAs that were detected at significantly lower level in SBM of twin embryos using mean normalization were involved in various developmental processes and differentiation. For example, miR-30b is involved in development of embryonic ectoderm [22], miR-373 promotes mesendoderm differentiation [23], and miR-24 is required for hematopoietic differentiation [24]. However, only miR-24 [25, 26] and miR-25 [3] were detected in pre-implantation embryo culture media.

Evidence is emerging that miRNA plays a role in embryo-endometrium cross talk during implantation [25, 27]. Microarray profiling revealed that six miRNAs were differentially expressed in the human endometrial epithelium during the implantation window, and secreted by the endometrial glands into endometrial fluid. miR-30d, the most differentially secreted miRNA in this study, has been reported to be internalized as an exosome-associated molecule by embryo TE, leading to an increase in the expression of genes involved in adhesion [28]. miRNAs detected in follicular fluid, such as miR-320, are thought to influence embryo quality [29]. However, embryo development is almost certainly affected not only by miRNAs from surrounding tissues, but also by
miRNAs in the early embryos themselves. For instance, miR-29b negatively regulates 
DNMTA3A/3B expression, altering DNA methylation levels in transition from morula to blastocyst 
stage [30], and miRNA Let-7a post-transcriptionally regulates the expression of ribonuclease type 
III Dicer1 altering microRNA profile and the implantation competency of the activated blastocysts [31].

Therefore, the most interesting finding was that 22.9% (11 out of 48) miRNAs secreted by 
twin embryos were never detected in normal, reproductively competent blastocysts (Fig. 6A). Since 
the TE cells of twin embryos express dual markers of both ICM and TE [9], such discrepancy in 
miRNA profile may not come as a surprise. miR-155, specifically detected in SBM of twin embryos, 
inhibits proliferation and migration of trophoblast-derived cell lines [32], whereas nothing is known 
about potential roles of the other ten “twin-specific” miRNAs in human preimplantation embryo 
development. Although some information concerning their potential roles in cancer diagnosis, 
prognosis and therapy has been reported, extrapolating this data to draw conclusions on their 
function in embryo development would currently be too speculative.

Moreover, these findings support miRNAs analysis from SBM as an effective approach to 
capture biological variability between embryos of different quality. In this study the miRNA analysis 
from SBM was able to clearly distinguish between low quality manipulated and non-manipulated 
high quality embryos. Accordingly, miRNAs analysis from SBM holds the potential to have enough 
resolution to capture biological variation from reproductively competent and non-competent 
blastocysts and to be use as new non-invasive biomarker of embryo selection.

In conclusion, we are only just starting to get a glimpse of the role of miRNA in 
development of human pre-implantation embryos and therefore, although the overall significance 
of the data presented here is not yet clear, it may add important information to the very limited 
understanding of these mechanisms.
Acknowledgments

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Author Disclosure Statement

The authors have no commercial, proprietary, or financial in the products or companies described in this article.
References


Figure legends

Figure 1. Expression of the lineage markers in twin and control embryos at Day 5 post-fertilization. Twin embryos generated by embryo splitting are smaller, often with no distinctive inner cell mass (ICM). They are less cellular in comparison with non-manipulated control embryos and most of the cells have a dual expression of ICM and trophectoderm (TE) markers. Green: ICM marker NANOG. Red: TE marker CDX2. Blue: DNA dye Hoechst 33342. In control embryo, asterisk marks ICM cell expressing NANOG and arrow TE cell expressing CDX2. In twins A and B, the arrowheads point to cells with dual expression of NANOG and CDX2.

Figure 2. Study design. Control embryos were cultured undisturbed up to fully-expanded blastocyst stage and SBM were collected before thawing. All seven blastocysts were diagnosed as euploid and healthy pregnancies ensued after transfer. Twin embryos instead were produced after embryo splitting at the cleavage stage. Twins A were cultured up to the blastocyst stage in their own zona pellucidae, twins B were cultured in host zona pellucidae. SBM was collected at the blastocyst stage. MicroRNAs were extracted and analysed from 7 controls and 7 pairs of twins.

Figure 3. Estimation of the correlation between samples within each group. (A) Distribution of Ct values for twins A, twins B and control embryos. (B) Distribution of Spearman test rho values (Ct) indicates that median correlations are very high for all conditions. Correlation between twins A (0.929 ± 0.031) and twins B (0.928 ± 0.040) samples are similar, whereas between control samples is lower (0.893 ± 0.086). (C) Pearson’s correlation matrix comparing raw Ct data among all analysed samples. A clear discrimination between twin embryos and controls can be deduced from the low correlation of miRNA profiles from SBM after in vitro culture. The numbers show the actual correlation value among all analysed samples. (D) Heat map of Ct values from twins A,
twins B and controls indicates no significant cluster by experimental group and high number of missing values.

Figure 4. Quality of the endogenous control and effect of the normalization with the endogenous control. (A) Ct values of the endogenous control U6 are heterogeneous between the samples and homogenous within each sample. The sample 6B has very low Ct value. (B) Whereas distribution of −ΔCt values in samples from control embryo group is relatively homogenous, the values are much more heterogeneous in samples from both twin groups indicating that expressing or missing miRNA can be different for each sample. (C) Heat map of −ΔCt values from twins A, twins B and controls indicates no significant cluster by experimental group, high number of missing values and that 6B sample (yellow dot) does not cluster with others. (D) miRNAs regulated in at least one comparison (FC≥1.5 and p-value ≤0.05) where U6 was used for normalization.

Figure 5. Comparison between miRNAs in SBM from twin A and B. (A) More than 95% (46/48) of miRNAs detected are shared by SBM from both twins A and B. (B) The volcano plots comparing twins A or twins B versus controls are almost overlapping, in fact no significant difference can be detected when comparing instead twins A versus twins B.

Figure 6. Comparison between twin and control SBM miRNA expression profile. (A) Thirty-seven miRNAs (green) were found solely in SBM from control blastocysts and are all co-expressed from TE cells; 22 miRNAs (blue) were found in SBMs from both control and twin blastocysts and are all co-expressed from TE cells; 26 miRNAs (dark and light violet) were found exclusively in SBM from twin blastocysts, but only 53.8% (14/26, dark violet) are co-expressed from TE cells. (B) The volcano plot shows 9 miRNAs significantly less abundant in SBM from twins versus control reproductively competent blastocysts, with a fold-change variable between -167.6 and -2.8, and 6
miRNAs instead significantly more abundant, with a fold-change between 2.7 and 319148.

Interestingly, miR-30c, putative biomarker of reproductive potential was found in the former cluster with a fold-change -47.0.

Figure 7. Putative roles of miRNA-30c in human blastocysts. (A) From 1643 miRNA-30c target genes, 1061 were expressed in ICM, 1166 in TE, and 1006 in both ICM and TE of the human blastocysts. (B) The pathways involving miR-30c targeted genes in ICM and TE according to the KEGG database.
Figure 1

DAY 5

CONTROL

NANOG  CDX2  Hoechst

TWIN A

TWIN B

DAY 6

CONTROL

NANOG  CDX2  Hoechst

TWIN A

TWIN B

Nolifigure1.eps

Figure 1

238x396mm (300 x 300 DPI)
Figure 3

254x362mm (300 x 300 DPI)
Figure 4

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Nolifigure4.eps

Figure 4

208x244mm (300 x 300 DPI)
Figure 5

94x101mm (300 x 300 DPI)
Figure 7

Pathway Description (KEGG) | Nb Genes in Pathway | Nb Regulated Genes | P Value | Corrected P Value
--- | --- | --- | --- | ---
Trophoblast mediated proliferation | 137 | 32 | 7.20E-06 | 1.17E-04
Wound healing | 134 | 29 | 4.10E-06 | 1.60E-05
Cell cycle | 125 | 19 | 3.52E-04 | 1.70E-02
Apoptosis | 110 | 22 | 8.09E-02 | 1.60E-01
Wnt signaling pathway | 69 | 37 | 1.03E-02 | 5.77E-01
Gom signaling pathway | 49 | 6 | 1.32E-02 | 2.94E-01
Hedgehog signaling pathway | 62 | 7 | 4.56E-02 | 3.39E-01
Progression mediated oocyte maturation | 117 | 14 | 2.59E-02 | 3.12E-01
Oocyte repair | 47 | 2 | 3.50E-02 | 6.35E-01
Pathway Description (KEGG) | Nb Genes in Pathway | Nb Regulated Genes | P Value | Corrected P Value
--- | --- | --- | --- | ---
Oocyte mediated attachment | 125 | 19 | 7.20E-06 | 1.17E-04
Oocyte incidence | 110 | 22 | 8.09E-02 | 1.60E-01
Wnt signaling pathway | 69 | 37 | 1.03E-02 | 5.77E-01
Gom signaling pathway | 49 | 6 | 1.32E-02 | 2.94E-01
Hedgehog signaling pathway | 62 | 7 | 4.56E-02 | 3.39E-01
Progression mediated oocyte maturation | 117 | 14 | 2.59E-02 | 3.12E-01
Oocyte repair | 47 | 2 | 3.50E-02 | 6.35E-01

Nolifigure7.png

Figure 7

73x30mm (300 x 300 DPI)
Supplemental Figure legends

Supplemental Figure S1. Immunostaining of lineage markers in Twin embryos, pair 1-3, used in the experiment. Their expression in blastocysts derived from split embryos is frequently non-specific. In addition, the embryos have have poorly developed ICM (arrowheads).


Supplemental Figure S2. Twin embryos, pair 4-7, used in the experiment.
Figure S1

Pair 1

Pair 2

Pair 3

NolIFigureS1.eps

Figure S1

254x362mm (300 x 300 DPI)
Figure S2
152x261mm (300 x 300 DPI)