Title: A METHOD FOR IMPROVING DEVELOPMENT POTENTIAL OF AN EMBRYO AND EMBRYOS DEVELOPED THEREFROM

Abstract: The present invention relates to a method of improving development potential of an embryo, embryos developed therefrom and organisms resulting from embryos developed from the method. In a first aspect of the present invention, there is provided a method of culturing an embryo to improve development potential, said method comprising: obtaining an embryo; and culturing the embryo to enhance trophoderm development of the embryo. The method relates to improving the chances of an embryo implanting to result in a successful pregnancy. The embryos desirably become implantation competent favouring foetal-maternal interaction and development to term of an embryo. The trophoderm development stimulating agent may be any compound which is proven to stimulate normal trophoderm development. Preferably the agent is fibroblast growth factor-4 (FGF4) protein.
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INTRODUCTION

The implantation and future survival of embryos in utero is dependent upon
many factors which contribute to proper development of the embryo to
blastocyst stage. After blastocyst development, favourable foetal-maternal
interactions contribute to long term survival and maintenance of the
embryo/foetus for further development into a successful pregnancy and finally
birth of an animal.

However, the environment in which the embryo develops is critical for both
successful implantation and further development. An understanding of the
requirements would enable a higher success rate for pregnancies since natural
and artificial implantation techniques are not always 100% possibly due to
defective embryos.

In nuclear transfer programs, the rate of success in inducing and maintaining
pregnancy drops dramatically from that seen in healthy embryos (eg. embryos
produced in vivo or in vitro). A large proportion of abnormal pregnancies
derived from nuclear transfer embryos are noted and caused by abnormal
placental development. This leads to defective foetal-maternal interactions
resulting in early in vivo death of the embryo or post natal mortality. This
difference in success rate between normal embryos and nuclear transfer
embryos, specifically somatic cell cloned nuclear transfer embryos indicates an
abnormality in the nuclear transfer embryos or the environment in which the
embryos implant and develop.
In normal embryos, it has been found that factors such as fibroblast growth factor 4 (FGF4) are crucial in proper embryo development. Homozygous deletion of the FGF4 gene results in a lethal embryonic phenotype similar to that observed for FGFr2 null mutants. Embryos develop normally to the blastocyst stage but degenerate soon after implantation, apparently due to an inability of the inner cell mass to thrive. *In vitro* culture of blastocysts demonstrated the absence of any extraembryonic endoderm formation in FGF4 null mutants and, that the mutant phenotype could be rescued by addition of recombinant human FGF4 in the culture medium.

An absence of FGF4 expression and the possibility that other genes from other cell lines involved in embryo development, are also aberrantly expressed in nuclear transfer embryos, may contribute to the low frequency of pregnancy and survival following transfer of cloned blastocysts to recipient animals.

Accordingly, it is an object of the present invention to improve the implantation and development of embryos, particular nuclear transfer embryos.

**SUMMARY OF THE INVENTION**

In a first aspect of the present invention, there is provided a method of culturing an embryo to improve development potential, said method comprising:

- obtaining an embryo; and
- culturing the embryo to enhance trophectoderm development of the embryo.

The method relates to improving the chances of an embryo implanting to result in a successful pregnancy. The embryos desirably become implantation competent favouring foetal-maternal interaction and development to term of an embryo.

In yet another aspect of the present invention, there is provided a method of developing an animal, said method comprising:
obtaining an embryo with improved development potential and prepared
by the methods described above;

obtaining a receptive animal capable of incubating an embryo to term;
implanting the embryo into the receptive animal; and

allowing the receptive animal to incubate the embryo to term.

In another aspect of the present invention, there is provided an animal obtained
by the methods described.

IN THE FIGURES

Figure 1 shows trophectoderm cell line resides with underlying monolayer.

Figure 2 shows RT-PCR results of fibroblast (F), term placenta (Pl) and TE
cells.

DESCRIPTION OF THE INVENTION

In a first aspect of the present invention, there is provided a method of culturing
an embryo to improve development potential, said method comprising:

obtaining an embryo; and

culturing the embryo to enhance trophectoderm development of the
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The method relates to improving the chances of an embryo implanting to result
in a successful pregnancy. The embryos desirably become implantation
competent favouring foetal-maternal interaction and development to term of an
embryo.

Applicants have found that FGF4 expression is aberrant in a high proportion of
embryos derived from somatic cell nuclear transfer techniques. This coincides
with the absence of viable trophectoderm cell lineages from blastocyst stage
mouse embryos lacking the FGF4 gene. These deficiencies correlated with an
observed higher proportion of abnormal pregnancies from nuclear transfer
embryos generally caused by an absence of successful implantation or abnormal placental development. Without being limited by theory, it is postulated that abnormal pregnancies may be associated with reprogramming failure in trophoderm lineages.

The embryo may be obtained from any source including naturally conceived embryos, artificially fertilised embryos, or they may be nuclear transfer embryos including those derived from somatic cell nuclear transfer techniques or they may be cloned nuclear transfer embryos or genetically modified embryos.

The term "embryo" as used herein is any young organism in the first stages of development. The embryo may be taken from the moment of conception or reconstruction or from the blastocyst stage or any stage between. The embryo may have an intact zona pellucida or the zona pellucida may be removed.

The term "blastocyst" as used herein is any embryo at any of the stages of blastocyst development including but not limited to "early blastocyst", "blastocyst", "expanding blastocyst" and "hatching or hatched blastocyst."

The embryo may be from any source selected from the group including bovine, ovine, porcine, caprine, murine or any animal that produces an embryo including humans.

The embryo may be any mammalian embryo but preferably a nuclear transfer embryo derived by any nuclear transfer method available to the addressee, using any cell type as the source of the donor nucleus. The embryo culture system used could be any culture system capable of supporting the successful development of nuclear transfer embryos to the blastocyst stage or further.

The embryo may be cultured in a medium capable of supporting development of the embryo to the blastocyst stage, for example, including but not limited to Synthetic Oviductal Fluid (SOF).
The trophectoderm cell lineage is considered to be important for successful implantation and further survival of the mammalian embryo *in utero*. However as recently found by the applicants, some embryos which result in abnormal pregnancy and/or abnormal placental development have a tendency to have deficient FGF4 expression and possibly aberrantly expressed genes involved in trophectoderm development.

Targeting the trophectoderm or enhancing its development may be achieved by exposure of the embryo to normal trophectoderm either directly, or indirectly, or through exposure of the embryo to a trophectoderm stimulating agent.

Alternatively, the embryo may be exposed to supernatant of a trophectoderm cell culture.

In a preferred aspect, the method includes the steps of:
- obtaining a source of trophectoderm cells; and
- culturing the embryo in the presence of the trophectoderm cells.

The trophectoderm cells may be derived from any source but preferably the source is compatible to the embryos that are being cultured.

The trophectoderm cells may be a cell line derived from trophectoderm cells of any species. Preferably such trophectoderm cells will be derived from the same species as the embryo. The term “trophectoderm cells” as used herein is intended to include all types of trophectoderm cells including “mature” trophectoderm cells, trophectoderm stem cells, trophectoderm vesicles or trophectoderm like cells identifiable by the expression of growth factors selected from the group including but not limited to TP, FGFr-2, LIF, EGF, HB-EGF or EGFR.

The trophectoderm cells may be derived from the embryo itself to create a trophectoderm cell monolayer. Preferably, the trophectoderm cells are a normal trophectoderm cell derived from a healthy source.
Without being limited by theory, it is possible that the aberrant development of the trophectoderm lineages in embryos, particularly nuclear transfer embryos may be corrected if the nuclear transfer embryos were cultured in the presence of normal trophectoderm cells preferably prior to transfer to a recipient animal.

The trophectoderm cell lines may provide factors in the media that would support the normal foetal placental development. It is postulated that the trophectoderm cells may be male or female and derived from in vitro or in vivo produced embryos. They may be bovine, for bovine nuclear transfer embryos, but the trophectoderm cells from any species could be matched with the nuclear transfer embryos from another species. The trophectoderm lineages may be isolated as previously described (Tanaka et al., 1998; Flechon et al., 1995) or using alternative methods known to the addressee.

The trophectoderm cells may be present as a cell culture, preferably a monolayer or as a cell suspension or they may be trophoblast vesicles from in vitro or in vivo produced embryos. In this preferred aspect, the presence of the trophectoderm cells enhances the development of the trophectoderm cells or the embryo. The trophectoderm cells may be placed in close proximity to the embryo or be aggregated with the embryo either by placement of trophectoderm cells on the embryo, such as in the absence of the zona pellucida or they may be placed under the zona pellucida when the zona pellucida is present.

The embryo may be cultured to the blastocyst stage or to any stage where trophectoderm development of the embryo is enhanced for favourable implantation and placenta development. The embryo may be cultured to any stage of development. Preferably, the embryo is transferred preferably onto a monolayer of trophectoderm cells at day 5 of preimplantation development or the morula stage equivalent for any species, and cultured further to the blastocyst stage. However, the embryo may be transferred preferably onto the trophectoderm monolayer at any stage of preimplantation development.

In a further preferred aspect, there is provided a method of culturing an embryo to improve development potential, said method comprising:
obtaining an embryo at the blastocyst stage;
obtaining a source of trophectoderm cells; and
introducing the trophectoderm cells into the blastocyst to provide an
embryo suitable for culturing or implantation.

The embryo may be as described above and cultured to a blastocyst stage by
any methods known to the skilled addressee. Preferably, the blastocyst stage
is a stage where the blastocyst cavity has developed.

The embryo may be any mammalian embryo but preferably it is a nuclear
transfer embryo derived by any nuclear transfer method available to the
addressee, using any cell type as the source of the donor nucleus. The embryo
culture system used may be any culture system capable of supporting the
successful development of nuclear transfer embryos to the blastocyst stage.

Similarly, the trophectoderm cells may be as described above and cultured by
any methods known to the skilled addressee. Specifically, such trophectoderm
cells may be derived from a trophectoderm cell line or isolated as trophoblast
vesicles from *in vitro* or *in vivo* produced embryos. Such trophectoderm cells
may be derived from any species. However, it is preferred that the
trophectoderm cells will be derived from the same species as the embryo. For
instance, bovine trophectoderm cells will be used for bovine embryos, but it is
also within the scope of the invention to use trophectoderm cells from any
species to inject into embryos of other compatible species.

The trophectoderm cells may be injected into the cavity of blastocyst stage
embryos. The injected trophectoderm cells may contribute to the
extraembryonic cell lineages and may help support the development of
embryos, particularly nuclear transfer embryos, specifically the extraembryonic
cell lineages.

The trophectoderm cells may be injected into the blastocyst cavity by any of the
methods available which do not harm the embryo. Micromanipulation is
preferred.
The number of trophectoderm cells may be varied. However, it is preferred to inject from 1 to 100 trophectoderm cells into the blastocyst cavity.

Alternatively, the trophectoderm cells may be introduced into the blastocyst by aggregating trophectoderm cells with the embryo by either placing trophectoderm cells on the embryo (in the absence of zona pellucida) or inserting trophectoderm under the zona pellucida when the zona pellucida is present. This allows the trophectoderm cells to integrate with the embryo.

In a further preferred aspect, the method further includes the step of:
   culturing the embryo preferably to the hatching blastocyst stage or any stage of blastocyst development.

The further culturing period will depend on the preferred stage of development of the blastocyst and also of the species of embryo cultured. However, any period of 24 to 48 hours is preferable. After this period, the injected embryo may be transferred to a recipient animal.

In yet another preferred aspect of the invention, the method further includes the step of:
   transferring the embryo after introduction of the trophectoderm cells to a recipient animal.

Any methods of transfer are available to the skilled addressee. However, general IVF techniques are suitable.

In a further preferred aspect of the present invention, there is provided a method of culturing an embryo to improve development potential, said method comprising:
   obtaining an embryo; and
   culturing the embryo in the presence of a trophectoderm stimulating agent.
The trophectoderm stimulating agent may be any compound which is proven to stimulate normal trophectoderm development. Preferably the agent is fibroblast growth factor-4 protein (FGF4) either in its natural or recombinant form, wherein the recombinant form is added extrinsically or produced in-situ. The FGF-4 may also be derived from cell cultures. Preferably, FGF-4 is provided in the supernatant of an embryonic carcinoma cell (ECC) culture.

Fibroblast growth factor 4 (FGF4) has previously been shown to be essential for the isolation of trophectoderm cell lines from mice and pigs. In addition, in the mouse, the aberrant developmental phenotype of FGF4 homozygous mutant embryos in vitro has been reversed by the addition of FGF4 to the culture media (Feldman 1995). However, despite the high number of reports of placental abnormalities and low implantation rates for embryos produced by nuclear transfer techniques and, the applicants recent finding that a large percentage of nuclear transfer embryos aberrantly express FGF4 at the blastocyst stage, there are no reports in the literature regarding the use of FGF4, or any other growth factors, in the culture media of nuclear transfer-derived embryos in an attempt to correct the apparently abnormal development of the extraembryonic cell lineages.

The embryo is as described above. Preferably, the embryo has been cultured to the morula stage or the blastocyst stage prior to addition of the trophectoderm stimulating agent. Preferably, the embryo is at the morula stage.

The trophectoderm stimulating agent or combination of agents may be added to an embryo culture at a suitable time of development of the embryo such as the morula or blastocyst stage, or the media may be changed to one already containing the trophectoderm stimulating agent. The time for changing the media or introducing the trophectoderm stimulating agent will vary. However, it is preferred to introduce the trophectoderm stimulating agent or combination of agents at approximately day 5 or at the morula stage equivalent depending on the species of animal.
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Where recombinant trophoderm stimulating agent is used, for instance recombinant FGF4 preferably FGF4 in the presence of heparin, the origin is preferably compatible with the species of embryo used. For instance for bovine embryos, bovine recombinant trophoderm stimulating agent or preferably bovine FGF4 is used. However, recombinant FGF4 protein derived from any species could be used with embryos from any other species dependent on cross species reactivity.

The amount of trophoderm stimulating agent used will depend on the species. However a concentration of 15 to 25 ng/ml preferably 20 ng/ml is used for addition to morula stage embryos.

In another aspect of the present invention, there is provided an embryo produced by the methods described. Preferably, the embryo is a blastocyst.

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The embryo, blastocyst or any stage of embryo development may be nuclear transfer derived. These may be further cultured to a stage of hatching demonstrating a level of implantation competency. Accordingly, in a preferred aspect, there is provided an embryo, blastocyst or any stage of embryo development ready for implantation.

It is also conceivable to use a genetically modified embryo, wherein the embryo is modified to express a trophoderm stimulating agent such as FGF-4. The embryo may be modified at any stage, preferably prior to fertilization at the oocyte and gamete stage. The oocyte or gamete may have introduced constructs which can express a trophoderm stimulating agent, preferably FGF-4. Enhanced expression may ensure improved development potential. Methods to enhance expression of trophoderm stimulating factor activity may be achieved by any recombinant means so as to achieve trophoderm development of the embryo. Suitable recombinant constructs incorporated into genetically modified embryos may allow the activation of expression of trophoderm stimulating agents at appropriate times to improve development potential.
In yet another aspect of the present invention, there is provided a method of developing an animal, said method comprising:

obtaining an embryo with improved development potential and prepared by the methods described above;

obtaining a receptive animal capable of incubating an embryo to term;
implanting the embryo into the receptive animal; and
allowing the receptive animal to incubate the embryo to term.

The embryo may be a blastocyst or be at any stage of embryo development providing it has been prepared by the methods described herein.

The receptive animal is an animal capable of carrying a foetus to term and may be a female animal in a breeding cycle or artificially induced to accept an embryo and to carry the foetus to term. By “artificially induced” it is meant that pharmaceutical grade synthetic hormones such as follicle stimulating hormone (FSH) in conjunction with luteinizing hormone (LH), using prescribed stimulation protocols for a given species, be injected in to the animal to prepare the womb for receiving the blastocyst.

In another aspect of the present invention, there is provided an animal obtained by the methods described.

The procedures described herein are designed to produce embryos, particularly nuclear transfer embryos with an improved capability of implantation in recipient animals and ultimately an improved efficiency of producing viable cloned animals. The procedures described have the advantage of producing embryos, particularly nuclear transfer embryos with an improved trophectoderm cell lineage with an increased chance of producing a viable extraembryonic cell lineage capable of normal implantation events, normal foetal / maternal interactions and capable of producing a placenta able to provide sufficient support to the developing foetus.
Throughout the description and claims of this specification, the word “comprise” and variations of the word, such as “comprising” and “comprises”, is not intended to exclude other additives, components, integers or steps.

Examples of the procedures used in the present invention will now be more fully described. It should be understood, however, that the following description is illustrative only and should not be taken in any way as a restriction on the generality of the invention described above.
EXAMPLES

Example 1: Analysis of FGF4 expression in bovine nuclear transfer embryos

(a) Collection of Bovine Oocytes
Bovine ovaries were obtained from a local slaughterhouse, transported at 25-30°C to the laboratory and washed in warmed phosphate buffered saline (PBS, Baxter, Australia). Ovarian antral follicles (2-8mm) were aspirated using an 18-gauge needle and collected into Hepses buffered Tissue Culture Medium 199 (TCM199, Gibco BRL/Life Technologies) with heparin (5000iu/ml, Sigma), 2% Foetal Calf Serum (FCS, Gibco/Life Technologies), and amphotericin B (250µg/ml, Sigma). Cumulus oocyte complexes (COC's) showing an even cytoplasm and surrounded by at least three layers of compact cumulus cells were collected from the follicular fluid. COC's were incubated and matured in groups of 25 in a TCM199 medium supplemented with gentamycin sulfate (10mg/ml), L-glutamine (29mg/ml, Sigma), human Chorionic Gonadotrophin (1500IU/ml, Lyppards, Australia) and 15% FCS at 39°C in 5%CO2 in air, for 20-24 hours.

(b) Preparation of Oocytes for Nuclear Transfer
In order to remove the surrounding cumulus, matured oocytes at 19-21 hours post maturation (hpm) were vortexed in 80µl maturation media and 20µl hyaluronidase (0.1%, Sigma) for 3 minutes in Eppendorf tubes (Quantum Scientific). The oocytes were washed through handling media (Hepses buffered TCM199 with 5% FCS (199HF)) and those at the metaphase II stage (i.e. with the first polar body extruded) were selected for nuclear transfer (NT).

(c) Fibroblast cell collection and culture
Fibroblast cells were prepared from skin and muscle sections from approximately 50-60 day old bovine foetuses. Tissue sections were diced in PBS using sterile scalpels and tweezers prior to digestion in 0.25% trypsin at 37°C for 20-30 minutes. DMEM culture media containing 10% FCS was then added to the sample to inactivate the trypsin and, the sample centrifuged for 5
minutes to pellet the cells. Following the removal of the supernatant, the cells were resuspended in DMEM with 10% FCS and cultured for up to three passages. Prior to nuclear transfer, fibroblast cells at 70% confluency were cultured for a further 5-7 days in serum depleted media (DMEM plus 0.5% FCS).

(d) Granulosa cell collection and culture
Mural granulosa cells were collected from an elite superovulated calf using an ultrasound-guided transvaginal probe. Granulosa cells were present in the collection media (DMEM containing 20\(\mu\)g/ml Amphotericin B, 1mg/ml Kanomycin Sulphate, 40\(\mu\)g/ml Chloramphenicol, 100\(\mu\)g/ml Chlorotetacycline, 60\(\mu\)g/ml Penicillin and 100\(\mu\)g/ml Streptomycin, Sigma) as morphologically distinct cell sheets. Granulosa cell sheets were placed on a percoll gradient (Sigma) using a bi-layer of 50% and 25% percoll, and centrifuged at 600G for 20 minutes. Cells located at the interface were collected and washed twice in DMEM with 10% FCS. Granulosa cells were cultured in DMEM with 10% FCS for up to three passages. Prior to nuclear transfer, granulosa cells at 70% confluency were cultured for a further 5-7 days in serum depleted media (DMEM plus 0.5% FCS).

(d) Nuclear Transfer by Microinjection
After mechanical disruption of the donor cell membranes in 199HF using the injection pipette, mural granulosa cells were injected directly into the cytoplasts. The reconstructed embryos were transferred back into TCM199 + 10% FCS until activation.

(e) Nuclear Transfer by cell fusion (SUZI)
Bovine oocytes were enucleated at 18-22hpm in handling media containing cytochalasin B (7.5\(\mu\)g/ml, Sigma) by gentle aspiration of the polar body and metaphase plate in a small amount of cytoplasm using a glass pipette (inner diameter: 10-15\(\mu\)m). A donor cell is then injected into the oocytes perivitelline space, directly following enucleation. The oocyte-cell complexes are cultured in maturation medium for approximately half an hour to one hour prior to cell fusion. Oocyte-cell complexes are transferred to mannitol fusion media at room
temperature, aligned at 600KHz pre 6.0V AC and fused with two pulses of 80.0-90.0 V DC for 15-30µs, one second apart, using wire electrodes 0.5mm apart. The oocyte-cell complexes are then placed into the maturation medium to allow cytoplasmic fusion to occur (5-20 minutes).

Artificial activation was induced either 0.5 or 4 hours after fusion or injection by exposing the oocytes to 5µM calcium ionophore for 4 minutes, prior to culture in 2mM 6-DMAP for five hours.

Embryos were cultured in modified Synthetic Oviductal Fluid (SOF) culture media (Gardner et al, 1994) supplemented with amino acids (Sigma), 5% FCS, myo-inositol (0.05g/10ml, Sigma) and sodium tri citrate (1mg/1ml, Selby Scientific). Embryos were submerged in a Submarine-Incubation-System (SIS, Vajta et al, 1997). The 4-well plates were gassed in foil bags (Wests Packaging Services) with 5% O₂, 5% CO₂ and 90% N₂ and immersed in 39°C water for up to seven days.

(f) Reverse Transcriptase Polymerase Chain Reaction (RT-PCR) embryo analysis

The protocols used for sample preparation, reverse transcription (RT) and polymerase chain reaction (PCR) amplification have been described previously (Daniels et al, 1997). Briefly, single oocytes or embryos were added to 5µl lysis buffer (0.8% Igepal, 5mM DTT, 1U/µl RNAsin), snap frozen in liquid nitrogen and stored at −80°C prior to use. When required, samples were heated to 80°C for 5 minutes, transferred straight to ice and the RT premix added. Reverse transcription was carried out in a final volume of 10µl comprising of the cell lysate, 1 × RT Buffer, 100U SuperScript H⁻ reverse transcriptase (GIBCO, BRL), 1.5 µg random primers (GIBCO, BRL), 5mM DTT and 1U/µl RNAsin. Reactions were held at 37°C for one hour. For granulosa cell samples, cells were scraped from the culture flask and pelleted in an Eppendorf tube in STE buffer (0.1M NaCl, 20mM Tris pH 7.4, 10mM EDTA pH 8.0). The supernatant was then removed, the cells resuspended in 40µl lysis buffer, snap frozen in liquid nitrogen and stored at −80°C. On use, cell lysates were thawed and centrifuged
at 12000g for 10 minutes to pellet cell debris. The supernatant was then transferred to a fresh Eppendorf tube and mRNA was extracted using a Dynal Beads mRNA purification kit (Dynal Pty. Ltd., Australia), as directed. Reverse transcription was carried out in a 20μl reaction mix with reagent concentrations as described for embryo analysis. Negative controls, omitting reverse transcriptase or added sample were always included.

PCR amplification was carried out on 2.5μl of the RT product from embryos or 1μl (approximately 20ng RNA or 2000 cells equivalent) from granulosa cell cDNA products. PCR cycles were as follows: 94°C x 5' followed by 50 cycles for embryos or 30 for cell samples of 94°C x 1'; 52°C x 1'; 72°C x 2'. Ten microlitres of the PCR products were visualised under ultra violet light on 2% agarose gels containing 1μg/ml ethidium Bromide. The PCR primer sequences for FGF4 were (5' to 3') TTCTCGTGCCATGAGCAG and AGGAAGTCGGTGACCTTCAT.

Results
In an initial experiment on nuclear transfer embryos derived from the microinjection of granulosa cell nuclei followed by activation of the resulting embryos 0.5 hrs after nuclear transfer, FGF4 transcripts were detected in only two of the nine embryos analysed at the morula and blastocyst stages. This is a significantly lower number (p<0.01) when compared to the detection of FGF4 transcripts in all ten IVF embryos analysed.

In a second experiment, a number of nuclear transfer embryos reconstructed with fibroblast nuclei were analysed for the presence of FGF4 transcripts. Embryos were produced by either microinjection and artificial activation either 0.5 (Group A) or 4 (Group B) hours after injection or cell fusion (SUZl) and activation 4 hours after fusion (Group C). In IVF embryos analysed at the blastocyst stage, FGF4 transcripts were detected in 37/43 embryos analysed (86%). However, FGF4 transcripts were detected in significantly fewer embryos at the blastocyst stage in group A (8/21, 38%, p<0.0005) and, in fewer embryos but with no significant difference in groups B (12/20, 60%) and C (13/21, 62%).
The results indicate that FGF4 is aberrantly expressed in a large proportion of nuclear transfer embryos produced with different donor cell nuclei and with different nuclear transfer techniques. Aberrant expression of FGF4 could indicate the abnormal development of the trophectoderm lineage.

Example 2: Trophectoderm enhancement of Nuclear Transfer Embryos
In order to assess the potential value of supporting the development of the trophectoderm lineage, embryos produced using SUZI nuclear transfer procedures, artificial activation 4 hours after fusion and fibroblast cells as the source of the donor nuclei, as described above, were separated into four groups. A control group of embryos were cultured to the day 7 blastocyst stage as described above and, three experimental groups were treated as described below.

1) Culture of embryos on a monolayer of trophectoderm cells.
At day 5 of culture, embryos at the morula stage of development were transferred in SOF media onto a monolayer of trophectoderm cells. Trophectoderm lineages were isolated from in vitro fertilised bovine embryos at the blastocyst stage as previously described (Tanaka et al., 1998; Flechon et al., 1995). The embryos were cultured for a further 48 hours prior to transfer to recipient cows.

2) Injection of trophectoderm cells into blastocyst cavity.
Day 6 embryos at the early blastocyst stage had approximately 10 trophectoderm cells injected into the blastocyst cavity. The trophectoderm cells were isolated from a trophectoderm cell lineage as described above. The embryos were cultured for a further 24 hours before being transferred into recipient cows.

3) Addition of recombinant FGF4 to embryo culture media.
At day 5 of culture, human recombinant FGF4 protein was added to the culture medium of embryos at the morula stage of development to a final concentration of 20ng/ml. The embryos were cultured for a further 48 hours prior to transfer to recipient cattle.
For each of the three experimental groups of embryos, ten recipient cows received two blastocyst stage embryos each. The cows were pregnancy tested at day 30 and day 60 using ultrasound techniques.

Example 3: Blastocyst Development and Differential Staining of Bovine Embryos Treated with rhFGF4

a) Isolation and culturing of trophectoderm (TE) cells.

Blastocysts (9 day IVP-produced) were primarily seeded on bovine fibroblast feeder cells and subsequent cultures were grown on 1% gelatin layers, cultured in Dulbecco's Modified Eagles medium (Trace Biosciences) supplemented with L-glutamine (Gibco BRL, Invitrogen), sodium bicarbonate (BDH), 100IU/ml penicillin (Gibco BRL, Invitrogen), 100μg/ml streptomycin, 1% non essential amino acids (Sigma Chemical Co.), 15% FCS (Gibco, BRL Invitrogen), 1mg/ml heparin (Sigma Chemical Co.) and 15% conditioned medium, which was collected from the supernatant of embryonic carcinoma cells (ECC), known to secrete fibroblast growth factor-4 (FGF-4). ECC supernatant was found to be a useful source of FGF4. The use of 1% gelatin is a simple and effective feeder layer, forming an appropriate membrane for attachment and proliferation of the cells, which did not require the use of cell feeder layers.

Two weeks after initial attachment, cells were passaged by standard trypsinisation or mechanical lifting. Following one month in culture, TE cells were passaged weekly by mechanically lifting of the monolayers and vesicles. Morphological analysis of cultures suggested that following attachment of blastocysts to bovine feeder layers, ICM cells degenerated and trophectoderm cells grew as monolayers of epithelium with dome-like formations from the centre of most colonies (see Figure 1). Trophectoderm vesicles were abundant during culture, varying in size and appearing morphologically like enlarged embryos. Differentiation of trophectoderm cells into giant cells was noticeable at the periphery of some colonies, however, initial culture experiments showed
that differentiation appeared to be less prominent with the addition of conditioned medium containing FGF4. Pure cultures of TE cells were isolated and have been grown continuously for 6 months up to passage 13.

5 b) Cryopreservation
Trophectoderm cell lines were successfully frozen and thawed when vesicles were vitrified using standard Open-Pulled-Straw procedures. Viability of trophoderm cell lines using standard cell freezing was extremely low.

10 c) Characterisation
TE cells were identified by expression of interferon-tau (IFN-τ) gene transcripts. IFN-τ was expressed in trophoderm cells, as it is responsible for maternal recognition of pregnancy in the bovine, with expression highest at day 12-15 of development. Results were compared against actin expression as shown in Figure 2.

d) Addition of FGF4 to IVP Embryos
The following experimental treatments were added on day 5 after fertilisation of embryos to coincide with expression of FGF4.

20 I SOF + CS
II SOF + BSA
III SOF+CS plus rhFGF4 (20ng/ml, rhFGF4, and 1mg/ml heparin, Sigma Chemical Co.)
IV SOF+BSA plus rhFGF4
25 V SOF+BSA plus 5%CMed (conditioned medium of ECC cultures) and 5%FCS
VI SOF+BSA plus rhFGF4 and 5%c/tFCS (charcoal treated foetal calf serum)
VII SOF+BSA plus 5%c/tFCS
30 VIII SOF+BSA plus 5%CMed and 5%c/tFCS

Embryos (IVP- In vitro produced bovine embryos) were cultured for 7 days in SOFM before being differentially stained for TE:ICM Ratios.

35 Table 1 shows that results of blastocyst development and differential straining of bovine embryos treated with rhFGF4.
TABLE 1: Blastocyst development and differential staining of bovine embryos treated with rhFGF4 and CMed.

<table>
<thead>
<tr>
<th>Treatment Groups</th>
<th>Blastocysts (%)</th>
<th>No.</th>
<th>Differential Staining</th>
<th>Total Cell No.</th>
<th>Ratio ICM:TE</th>
</tr>
</thead>
<tbody>
<tr>
<td>I). IVP CS</td>
<td>254/927 (27)</td>
<td>23</td>
<td>34.3 ± 2.20&lt;sup&gt;a&lt;/sup&gt;</td>
<td>117.0 ± 7.86&lt;sup&gt;b&lt;/sup&gt;</td>
<td>151.3 ± 8.26&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>II). IVP BSA</td>
<td>201/814 (25)</td>
<td>23</td>
<td>33.70 ± 2.87&lt;sup&gt;b&lt;/sup&gt;</td>
<td>110.8 ± 5.12&lt;sup&gt;b&lt;/sup&gt;</td>
<td>144.5 ± 6.54&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>III). IVP CS:rhFGF4</td>
<td>215/887 (24)&lt;sup&gt;a&lt;/sup&gt;</td>
<td>19</td>
<td>28.16 ± 2.23&lt;sup&gt;b&lt;/sup&gt;</td>
<td>97.21 ± 5.19</td>
<td>125.4 ± 6.48&lt;sup&gt;a&lt;/sup&gt;</td>
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<tr>
<td>IV). IVP BSA:rhFGF4</td>
<td>251/865 (29)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>31</td>
<td>32.96 ± 2.09&lt;sup&gt;b&lt;/sup&gt;</td>
<td>98.48 ± 4.63</td>
<td>127.5 ± 5.04&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>V). IVP BSA:CMed+FCS</td>
<td>149/573 (26)</td>
<td>20</td>
<td>35.85 ± 2.14</td>
<td>95.85 ± 5.28&lt;sup&gt;a&lt;/sup&gt;</td>
<td>132 ± 5.38</td>
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<tr>
<td>VI). IVP BSA:rhFGF4+c/FCS</td>
<td>127/399 (32)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>35</td>
<td>37.97 ± 1.49</td>
<td>94.77 ± 2.54&lt;sup&gt;a&lt;/sup&gt;</td>
<td>132.8 ± 3.32</td>
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<tr>
<td>VII). IVP BSA:c/FCS</td>
<td>225/713 (32)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>21</td>
<td>36.81 ± 2.02</td>
<td>108.3 ± 5.45</td>
<td>145.1 ± 6.87&lt;sup&gt;b&lt;/sup&gt;</td>
</tr>
<tr>
<td>VIII). IVP BSA:CMed+c/FCS</td>
<td>288/1002 (29)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>22</td>
<td>44.50 ± 3.14&lt;sup&gt;a&lt;/sup&gt;</td>
<td>102.0 ± 3.47</td>
<td>146.5 ± 5.15&lt;sup&gt;b&lt;/sup&gt;</td>
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</tbody>
</table>

Comparison between a vs b within columns are significantly different (p<0.05)
Results show the following:

- The ratio of ICM:TE in all treatment groups is approximately 1:3.0. Addition of (VIII) CMed and c/tFCS on day 5 resulted in significantly higher numbers of ICM cell (p<0.05) when compared to control and rhFGF4 treatment groups (I, II, III and IV).

- Addition of FGF4 (either with rhFGF4 or CMed) appeared not to increase TE proliferation, and in groups (V and VI) was significantly lower (P<0.05) when compared to controls (I and II). However, addition of FGF4 or CMed may provide conditions that establish a tighter control over the ratio of ICM:TE (Reports show that the developmental competence of a transferred blastocyst is related to the establishment of a ICM:TE ratio of 1:3).

- Initial testing of in vivo development following embryo transfer indicated embryos from groups I, II, III, IV and VIII transferred to recipients (2 embryos per recipient) and assessed for pregnancy by ultrasound between day 30 and 60 resulted in 24/53(45%), 12/20(60%), 4/10(40%), 6/9(67%) and 5/7(71%) pregnancy rate, respectively. It appears that addition of rhFGF4 or CMed does not have a detrimental effect on embryo implantation.

Finally, it is to be understood that various other modifications and/or alterations may be made without departing from the spirit of the present invention as outlined herein.
REFERENCES


Flechon et al., 1995. Placenta 16:643-658


CLAIMS:

1. A method of culturing an embryo to improve development potential, said method comprising:
   obtaining an embryo; and
   culturing the embryo to enhance trophectoderm development of the embryo.

2. A method according to claim 1 wherein the embryo is selected from the group including naturally conceived embryos, artificially fertilised embryos, nuclear transfer embryos, cloned nuclear transfer embryos or genetically modified embryos.

3. A method according to claim 1 or 2 wherein the embryo is derived from a source selected from the group including bovine, ovine, porcine, caprine, murine or human.

4. A method according to any one of claims 1 to 3 wherein the embryo is a nuclear transfer embryo.

5. A method according to any one of claims 1 to 4 wherein the trophectoderm development is enhanced by exposure of the embryo directly or indirectly to trophectoderm cells, exposure of the embryo to a trophectoderm stimulating agent, or exposure to a supernatant of a trophectoderm cell culture.

6. A method according to any one of claims 1 to 5 further including the steps of:
   obtaining a source of trophectoderm cells; and
   culturing the embryo in the presence of the trophectoderm cells.

7. A method according to claim 6 wherein the trophectoderm cells are selected from the group including mature trophectoderm cells, trophectoderm stem cells, trophectoderm vesicles or trophectoderm like cells identifiable by the
expression of growth factors selected from the group including TP, FGFr-2, LIF, EGF, HB-EGF or EGFR.

8. A method according to claim 6 or 7 wherein the trophectoderm cells are cultured as a monolayer or as a cell suspension with the embryo.

9. A method according to claim 6 or 7 wherein the trophectoderm cells are aggregated with the embryo.

10. A method according to claim 9 wherein the trophectoderm cells are placed on the embryo.

11. A method according to claim 9 wherein the trophectoderm cells are cultured under the zona pellucida of the embryo.

12. A method according to any one of claims 1 to 11 wherein the embryo is developed to a morula stage.

13. A method according to any one of claims 1 to 11 wherein the embryo is developed to a blastocyst stage.

14. A method according to claim 13 wherein the trophectoderm cells are introduced into the blastocyst by injecting into the blastocyst cavity.

15. A method according to claim 14 wherein at least one trophectoderm cell is injected into the blastocyst cavity.

16. A method according to any one of claims 1 to 15 wherein the embryo is cultured to a hatching blastocyst stage.

17. A method according to any one of claims 1 to 16 wherein the embryo is cultured in the presence of a trophectoderm stimulating agent.
18. A method according to claim 17 wherein the trophectoderm stimulating agent is fibroblast growth factor-4 protein (FGF-4).

19. A method according to claim 18 wherein the FGF-4 is natural or recombinantly produced.

20. A method according to any one of claims 17 to 19 wherein the trophectoderm stimulating agent is cultured with the embryo at a stage equivalent to the morula stage of development.

21. A method according to any one of claims 17 to 20 wherein the trophectoderm stimulating agent is present at a concentration of 15 to 25 ng/ml.

22. A method according to any one one of claims 17 to 21 wherein the embryo is a genetically modified embryo which is modified to express FGF-4.

23. A method according to any one of claims 1 to 22 wherein the embryo is cultured for a period of at least 24 hours prior to transferring to a receptive animal.

24. An embryo with improved development potential prepared by a method according to any one of claims 1 to 23.

25. A method of developing an animal, said method comprising:
   - obtaining an embryo with improved development potential according to claim 24;
   - obtaining a receptive animal capable of incubating an embryo to term;
   - implanting the embryo into the receptive animal; and
   - allowing the receptive animal to incubate the embryo to term.

26. A method according to claim 25 wherein the receptive animal is a female animal in a breeding cycle or is artificially induced.

27. An animal prepared by the method according to claim 25 or 26.
28. A method according to claim 1 substantially as hereinbefore described with reference to the examples.
Figure 2

M  F  Pl  TE  -  F  Pl  TE  -

Actin  IFN-tau
INTERNATIONAL SEARCH REPORT

International application No.
PCT/AU01/00937

A. CLASSIFICATION OF SUBJECT MATTER

Int. Cl. 7: A01K 67/02

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

AS BELOW.

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

AS BELOW.

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
WPIDS, Chemical Abstracts, Medline : Keywords ; embryo, blastocyst, fibroblast growth factor 4, fgf4,

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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Further documents are listed in the continuation of Box C

See patent family annex

* Special categories of cited documents:
  "A" document defining the general state of the art which is not considered to be of particular relevance
  "E" earlier application or patent but published on or after the international filing date
  "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
  "O" document referring to an oral disclosure, use, exhibition or other means
  "P" document published prior to the international filing date but later than the priority date claimed
  "T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
  "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
  "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
  "&" document member of the same patent family

Date of the actual completion of the international search

19 September 2001

Name and mailing address of the ISA/AU

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Date of mailing of the international search report

17 September 2001

Authorized officer

ALISTAIR BESTOW
Telephone No : (02) 6283 2450
<table>
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<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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**INTERNATIONAL SEARCH REPORT**

**Box I**  
Observations where certain claims were found unsearchable (Continuation of item 2 of first sheet)

This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. ☐ Claims Nos:
   
   because they relate to subject matter not required to be searched by this Authority, namely:

2. ☒ Claims Nos: 1 - 17

   because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
   
   Claims 1 - 17 are so broad and of indeterminate scope, that a complete search could not be undertaken for these claims. The scope of the means by which the 'culturing the embryo to enhance trophectoderm development' occurs, is quite unknown, except where such means is by use of fibroblast growth factor 4 (fgf4). Therefore the above claims are searched only in as far as the claimed methods use fgf4.

3. ☐ Claims Nos:

   because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a)

**Box II**  
Observations where unity of invention is lacking (Continuation of item 3 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

1. ☐ As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims

2. ☐ As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.

3. ☐ As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:

4. ☒ No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

**Remark on Protest**

☐ The additional search fees were accompanied by the applicant's protest.

☐ No protest accompanied the payment of additional search fees.