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**Search for naive human pluripotent stem cells**

Fonseca SAS *et al*. Human naïve pluripotent stem cells

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**Abstract**

Normal mouse pluripotent stem cells were originally derived from the inner cell mass (ICM) of blastocysts and shown to be the *in vitro* equivalent of those pre-implantation embryonic cells, and thus were called embryonic stem cells (ESCs). More than a decade later, pluripotent cells were isolated from the ICM of human blastocysts. Despite being called human ESCs, these cells differ significantly from mouse ESCs, including different morphology and mechanisms of control of pluripotency, suggesting distinct embryonic origins of ESCs from the two species. Subsequently, mouse pluripotent stem cells were established from the ICM-derived epiblast of post-implantation embryos. These mouse epiblast stem cells (EpiSCs) are morphological and epigenetically more similar to human ESCs. This raised the question of whether cells from the human ICM are in a more advanced differentiation stage than their murine counterpart, or whether the available culture conditions were not adequate to maintain those human cells in their *in vivo* state, leading to a transition into EpiSC-like cells *in vitro*. More recently, novel culture conditions allowed the conversion of human ESCs into mouse ESC-like cells called naïve (or ground state) human ESCs, and the derivation of naïve human ESCs from blastocysts. Here we will review the characteristics of each type of pluripotent stem cells, how (and whether) these relate to different stages of embryonic development, and discuss the potential implications of naïve human ESCs in research and therapy.

**Key words:** Naïve pluripotent stem cells; Epiblast stem cells; Human embryonic stem cells; X chromosome inactivation

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**Core tip:** Mouse embryonic stem cells (ESCs) are the *in vitro* equivalent of the epiblast of pre-implantation embryos. Human ESCs on the other hand, although also pluripotent, appear to be in a more advanced developmental stage than their murine counterpart. Recently, several reports described culture conditions able to support mouse ESC-like human ESCs, called naïve human ESCs. Here we will review the search for human naïve ESCs, discussing their biological and practical relevance.

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**FROM THE MOUSE EMBRYO TO EMBRYONIC STEM CELLS**

The mammalian pre-implantation development is a process of cell multiplication and differentiation from the zygote to the totipotent blastomeres at the cleavage stage, and then to the pluripotent cells at the blastocyst stage. During preimplantation development, cell lineage specification and spatial segregation result in two cell fate decisions, which are believed to be sequential. The first segregation is into the trophectoderm (TE), involved in the formation of the fetal portion of the placenta, and the ICM at the early blastocyst stage. The second segregation divides the ICM into the primitive endoderm (PrE), that origins parietal and visceral endoderm of the yolk sac[1], and the epiblast, which will give rise to the embryo proper[2] (Figure 1).

Mouse ESCs are derived from the ICM of blastocysts and represent the epiblast lineage *in vitro*[3,4]. They are able to maintain an undifferentiated state (self-renewal) and to differentiate into tissues from all three germ layers *in vitro* and *in vivo*. In addition, when reintroduced in preimplantation embryos, mouse ESCs are able to get incorporated and give rise to all tissues of the resulting chimera.

In the pluripotent state, mouse ESCs are characterized by the presence of transcription factors such as OCT3/4[5] and Nanog[6], expression of alkaline phosphatase and specific cell surface glycoproteins such as SSEA-1 (stage-specific embryonic antigen 1)[7], and high telomerase activity[8]. In addition, like the epiblast lineage *in vivo*, female mouse ESCs have not undergone random X chromosome inactivation (XCI), and will do so upon differentiation *in vitro*[9] .

Maintenance of the undifferentiated state of ESC *in vitro* is greatly dependent on exogenous factors. The first mESC derived were maintained in medium conditioned by teratocarcinoma cells[4], or co-cultured with irradiated mouse embryonic fibroblasts (MEF) in an enriched culture medium which guaranteed self-renewal[3,10]. Later, maintenance of pluripotency was shown to be dependent on myeloid leukemia inhibitory factor (LIF), a molecule which induces differentiation of M1 myeloid leukemia cells[11]. LIF belongs to the interleukin-6 cytokine family and binds to a heterodimeric receptor consisting of the LIF-receptor and gp130[12]. LIF acts on mESC self-renewal mainly by activating Stat3 (signal transducer and activator of transcription 3) through the LIF/Stat3 pathway[13]. In addition, different matrices and scaffolds are also important for maintenance and/or determination of cell fate[14].

**HUMAN ESCS**

The first lines of human ESCs were established in 1998 from the ICM of human blastocysts, in culture conditions equivalent to those of mouse ESCs[15,16]. Subsequently, it was shown that instead of LIF, bFGF was required for the prolonged culture of human ESCs in the pluripotent state[17].

Despite being derived from pre-implantation embryos in apparently the same stage of development, human ESCs differ from their murine counterparts in many ways, including morphology (flat vs. domed shaped colonies), passage in culture (clusters *vs* single cell passage), growth factors dependence (bFGF and TGFB/ACTIVIN/NODAL *vs* LIF/STAT3 signaling)[16], genomic stability[18], and the epigenetic state of the X chromosome[19,20]. Furthermore, control of induced specific differentiation of mouse ESCs was shown to differ from human ESCs in many cases[21] (, where protocols established with mouse ESCs do not necessarily yield the same differentiated cell population from their human counterpart. This suggests distinct epigenetic states and/or developmental pathways between ESCs from the two species.

Regarding the X chromosome, although female human ESCs in a pre-XCI status equivalent to that of mouse ESCs have been described[22], this appears to be an unstable epigenetic state, which is lost with time in culture leading to pluripotent cells with one active and one inactive X chromosome[19,23,24]. This suggested that human ESCs may be in a different developmental stage than mouse ESCs. However, the kinetics of XCI during human pre-implantation development is still not clear, and thus it is not known whether human ESCs are indeed the *in vitro* equivalent of the epiblast lineage, or whether they are altered during establishment/culture by the current *in vitro* conditions[25,26].

**INDUCED PLURIPOTENT STEM CELLS**

A novel source of pluripotent stem cells was established in 2006: those derived through nuclear reprograming of somatic cells by exogenous expression of Oct4, Klf4, c-Myc and Sox2[27]. These induced pluripotent stem cells (iPSCs) were first established from mouse fibroblasts, and the same strategy was able to convert human fibroblasts also into a pluripotent state[28].

Human and mouse iPSCs are equivalent to their embryo-derived counterparts human and mouse ESCs, respectively[27,28], regarding morphology, growth factor dependence and XCI state. Thus, human and mouse pluripotent stem cells derived from developmentally equivalent cells (adult fibroblasts) are still distinct in several important aspects. This may suggest that the differences between pluripotent stem cells from the two species are not due to distinct origins of the starting embryonic cell, but rather to distinct pathways defining and controlling pluripotency.

**MOUSE ESCS**

Indirect evidence that human ESCs are not the developmental equivalent of mouse ESCs came from the establishment of pluripotent SCs derived from the epiblast of post-implantation mouse embryo - EpiSCs[29,30]. Although these cells form teratomas when injected in SCID mice, when injected into preimplantation embryos EpiSCs do not integrate to form chimeras like mouse ESCs do. EpiSCs keep their pluripotency *in vitro* when cultured under human ESCs´ conditions, present the same flat colony-morphology of the human cells, and are in a post-XCI state. In addition, EpiSC and human ESCs present expression profiles characteristic of the epiblast, suggesting that indeed the human pluripotent cells have embryonic origin distinct of mouse ESCs, and similar to mouse EpiSC.

 Therefore, two distinct classes of pluripotent SCs were defined: ground state, or naïve-pluripotent SCs, like mouse ESCs/iPSCs; and primed-pluripotent SC, like human ESCs/iPSCs and mouse EpiSCs, which are in a latter developmental stage[31]. And thus, the question was rephrased: can human naïve ESCs be established?

 In 2009, Bao *et al*[32] demonstrated that mouse EpiSC were able to revert to the more primitive epigenetic state of ESCs just by culture in LIF/MEFs. After few passages, ESC-like cells started appearing among EpiSCs, and eventually a sub-line of reprogrammed epiblast ESC-like cells was derived. As part of the reprogramming, the cells reactivated the inactive X chromosome, changing into a pre-XCI state like mouse ESCs and iPSCs. Thus, if mouse EpiSCs could be reverted into ESC-like cells, could human ESCs also be reverted into a naïve pluripotent state by simply changing media conditions?

**CONTROL OF PLURIPOTENCY BY SMALL MOLECULES**

One important improvement in the dissection of the pluripotent state was the demonstration that it was not dependent on external *stimuli* like serum and MEFs, but instead that pluripotency could be maintained by inhibition of endogenous signaling by mitogen-activated protein kinase (MEK), fibroblast growth factor (FGF) receptor tyrosine kinase and glycogen synthase kinase 3 (GSK3). This lead to the development of the 3i culture condition, which involves small molecules-inhibition of GSK3, MEK and FGF receptor tyrosine kinase signaling by small molecules CHIR99021, PD18352 and SU5402, respectively[33]. While inhibition of GSK3 promotes cell growth and viability, inhibition of FGF receptor tyrosine kinase blocks phosphorilation of MEK, thus hampering its action in promoting cell commitment (Figure 2). In comparison, in the original culture conditions of mouse ESCs, LIF acts only in blocking cell commitment, which would be one step later that ERK phosphorilation, and other factors present in serum and secreted from MEFs promote cell growth and viability[13,34].

Subsequently it was found that inhibition of MEK by PD0325901 was more potent and could substitute both PD18352 and SU5402, in what was called 2i media (CHIR99021+PD0325901)[33,35].The 3i/2i culture conditions allowed the derivation of germ line competent pluripotent stem cells from other species like rat[36,37] and pigs[38-43].

Therefore, could this media support derivation of human naïve ESCs? Although one report described the generation of apparently naïve hiPSCs in 3i media[44], those results were not replicated by any other group, nor were the human cell lines derived ever used in other reports. Thus, by the lack of publications at that time on the derivation of human naïve ESCs, it became obvious that those relatively simple conditions that worked for many mammalian species were not enough for human cells.

 Nevertheless, small molecules took the spot light in pluripotent stem cell research, including epi-drugs like inhibitors of DNA methyltransferases and histone deacetylases, shown to have important roles on cell reprograming and maintenance of pluripotency[45].

**HUMAN NAÏVE ESCS**

Indeed, it was subsequently shown that naïve human pluripotent SCs could only be obtained in 2i when expression of exogenous reprogramming factors was maintained in the cells[46,47]. Generation of hiPSCs with constant ectopic expression of Oct4, Sox2, Klf4, c-Myc, and Nanog in the presence of LIF lead to human pluripotent SCs with behavior in culture and ease to be genetically manipulated equivalent to those of mouse ESCs[46]. Similarly, ectopic induction of Oct4, Klf4 and Klf2 in human ESCs in 2i/LIF media reprogrammed the cells into a naïve state transcriptionally similar to that of mouse ESCs, including the reactivation of the inactive X[47]. Nevertheless, both strategies required continuous expression of the exogenous factors for maintenance of the naïve state.

 The next year, Wang *et al*[48] reported the efficient generation of exogenous factor-independent naïve hiPSC by transient expression of *Oct4, c-Myc, Klf4, Sox2, Rarg* and *Lrh-1* in adult fibroblasts, demonstrating a pivotal role of retinoic acid receptor signaling in reprogramming. However, to our knowledge their results were not replicated by any other group.

 After a hiatus of almost two years on reports of naïve human PSCs, within 3 months four articles described the identification of culture conditions adequate for culture and/or derivation of human naïve ESCs and iPSCs without expression of external factors[49-52] (Table 1). The first report performed a functional screen of 16 small molecules able to support *OCT4* expression in hiPSCs in LIF/2i. This lead to the identification of a combination of 7 factors (including bFGF) able to maintain hiPSCs in a naïve state and to support derivation of naive human ESCs from blastocysts[49]. When introduced to mouse embryos, the naïve cells were able to contribute to the formation of E10.5 chimeras, a hallmark of mouse ESCs and not EpiSCs.

In another report, based on the finding that *NANOG* expression is higher in the epiblast of human pre-implantation epiblast than in human ESCs[53], Chan *et al*[50] tested 11 small molecules that would increase *NANOG* expression in hESCs. They identified a combination of 3 factors that rewired human ESCs into a naïve state responsive to LIF, which was required for self-renewal of the cells (Table 1). In addition, the authors showed that the naïve cells had an expression profile more similar to that of human blastocyst cells than conventional (or primed) human ESCs.

 A third report of conditions for naïve human PSCs derivation and culture described a simpler strategy, where human ESCs were exposed to histone deacetylase inhibitors for a few passages before being transitioned to 2i media with bFGF[51] (Table 1). The 2i+bFGF also supported the derivation of a new line of naïve human ESC that could then be transitioned in to 3i media with LIF. Nevertheless, the efficiency of derivation was very low, and cells in 3i+LIF showed significantly higher rates of differentiation than those kept in 2i+bFGF. Furthermore, although cloning efficiency of the naïve human ESC line established was twice of the primed cell line, it was still relatively low (20%) when compared to other reports, whereas there was no improvement of cloning efficiency of naïve H1 compared to the original primed H1. Thus, the strategy appeared to generate cells in an intermediate naïve state.

 Another alternative media for maintenance of transgene-expression free naïve hiPSCs was described by Valamehr *et al*[52] (Table 1). While developing culture conditions for high throughput generation of hiPSCs, the group developed a media containing inhibitors of ROCK, GSK3 and MEK pathways, LIF and bFGF that allowed long term culturing of naïve hiPSCs derived in a previously described cocktail of small molecules[54]. Although those naïve hiPSCs did not have mouse ESC-like morphology, they showed high survival rates in single-cell dissociation, and appeared to be in a pre-XCI state.

 More recently, Theunissen *et al*[55] performed an alternative functional screen of small molecules able to support growth of human naïve ESCs based on the finding that naïve pluripotent stem cells use the distal while primed cells used the proximal enhancer of *OCT4*. Using a GFP-reporter gene under the control of the distal enhancer of *OCT4* in human ESCs, the authors screened a kinase inhibitor library in the presence of 2i/LIF, identifying a combination of 5 compounds which, together with Activin A and FGF, supports conversion of primed into naïve human ESCs and the derivation of new naïve human ESCs and iPSCs (Table 1). It is important to note that, although these converted or newly established naïve cells had the expected dome-shaped morphology and expression profile more similar to mouse ESCs than to primed human ESCs, they presented a few undesired characteristics, namely, (1) high frequency of aneuploidies, and (2) XCI. The authors suggest that, rather than limitations of their “naïve” cells, these may represent differences between naïve human and murine ESCs.

**HUMAN NAÏVE ESCS: FACT OR ARTIFACT?**

Which brings us to a crucial point: what is a human naïve ESC? When searching for the human naïve pluripotent state, are we looking for the *in vitro* equivalent of the human epiblast, or for a human pluripotent stem cell as amenable to manipulation in culture as mouse ESCs? Depending on the goal, the characteristics of the resulting naïve human cells can be very different.

 The methods for culturing human primed ESCs have significantly improved in the last 10 years with the development of defined and feeder-free culture conditions and of solutions to substitute manual passage. Nevertheless, these cells are still harder to culture and to genetically manipulate than their mouse counterpart. Thus, the development of culture conditions that make human ESCs behave like mouse ESCs may facilitate their use in research and therapy, regardless of whether this naïve state has any relevance as a model for human development. In that case, we know that these human naïve ESCs should have enhanced proliferation, be passaged as single cells (high cloning efficiency), be genetically stable and amenable to efficient genetic manipulation *in vitro*. In addition, being in a pre-XCI state and undergoing random XCI upon differentiation would be desirable for the generation of naïve hiPSCs to model X-linked diseases[56].

 On the other hand, if the goal is to obtain cells that model the pluripotent epiblast from human blastocysts, then we must step back and look at what we know about these cells *in vivo*. Despite the difficulties in studying the human embryo, enough has been done to know that there are significant differences between human and mouse blastocysts so that one cannot completely extrapolate data from one species to the other[57]. For instance, at the blastocyst stage, while *Oct4* expression is restricted to epiblast cells in mouse[58], it is also found in the human throphectoderm[59]. Thus, the organization and/or the markers of pluripotent cells in the human blastocyst may differ from mouse.

 Recent advances in single-cell RNA sequencing (RNAseq) have allowed the study of expression profiles of individual cells during human pre-implantation development[60,53]. Analysis of RNAseq data from 30 single-cells derived from human blastocysts identified three distinct clusters corresponding to cells from the epiblast, primitive endoderm and throphectoderm. Furthermore, comparison of the expression profile of human epiblast cells with human ESCs at different stages of derivation showed that, although similar, there are significant changes in gene expression in the embryo-derived cells at the initial primary outgrow of hESCs[53]. That is not surprising given that the cells are adapting to the *in vitro* conditions, but how distinct can they be while still modeling the epiblast *in vitro*? It will be interesting to do the same comparison in mice in order to evaluate whether these changes between murine epiblast and ES cells are less accentuated. Nevertheless, it would be reasonable to use the single-cell RNASeq data from human blastocysts as a signature of “true” naïve human ESCs. Indeed, Chan *et al*[50] showed that the expression profile of their naïve human ESCs was more similar to those of single cells from human blastocysts. The same analysis should be performed with the other reported naïve human ESCs for comparison.

 Regarding XCI, the kinetics of the process is very well characterized during mouse pre-implantation development, starting at the two-cell stage with the imprinted inactivation of the paternal X (pX) in all cells of the embryo. Then, at the blastocyst stage, cells of the ICM reactivate the pX, and go through a second wave of XCI, where each cell will randomly choose the Xi. In contrast, much less is known about the XCI status of the human pre-implantation embryo, and results are conflicting[23]. It would be extremely valuable if “true” naïve human ESCs were in a pre-XCI state like their murine counterpart in order to study the mechanisms of this epigenetic phenomenon *in vitro*. However, we still don´t know whether this would represent the *in vivo* situation, and thus, whether the pre-XCI state is a prerequisite for being a “true” naïve human ESC.

 While this manuscript was in revision, Wang *et al*[61] showed that small populations of naïve human pluripotent SCs exist within cultures of primed human ESCs and iPSCs. These naïve cells can be purified and maintained in 2i/LIF media. In addition, the authors identified increased expression of the primate-specific endogenous retrovirus HERVH as a genetic signature of naïve human pluripotent SCs, which can be used for the development of culture conditions to support human naïve PSC culture. More importantly, the study of HERVH-associated transcription during human pre-implantation embryonic development may provide more insights into this still much-unknown process.

In conclusion, in the last year different culture conditions were developed that support the derivation and growth of human ESCs more similar to mouse ESCs. Whether the distinct methods generate similar or distinct epigenetic states of human pluripotent SCs, and more importantly, which of these potentially naïve human pluripotent SCs are more similar to the cells of the human epiblast remain to be determined. These studies will have great impact on the establishment of a good *in vitro* model for human embryonic development.

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**B**

**A**

**Figure 1** First two cell segregations during pré-implantation embryonic development. A: In blue, cells from the inner cell mass (ICM) in the early blastocyst, and white, cells from the trophectoderm; B: At the late blastocyst stage, cells from the ICM segregate into cells from the epiblast (green) and the primitive endoderm (orange).

**SELF RENEWAL**

**GSK3**

**LIF**

***PD18352***

**COMMITMENT**

***SU5402***

**MEK**

***CHIR99021***

**FGFr**

**(i)**

**(ii)**

**Figure 2** **Maintenance of pluripotency by LIF and small molecules. Green arrows indicate activation, red blunted lines indicate inhibition.** The pluripotent cell must self-renew without committing to any specific cell type. This may be achieved by promoting self-renewal and inhibiting commitment by LIF; or by small molecules (in italic) which inhibit (i) signaling through the FGF receptor tyrosine kinase (FGFr), thus inhibiting commitment; and (ii) GSK3, promoting self-renewal. Adapted from[33]. LIF: Leukemia inhibitory factor; FGFr: Fibroblast growth factor receptor-specific tyrosine kinase; GSK3: Fibroblast growth factor (FGF) receptor tyrosine kinase and glycogen synthase kinase 3); MEK: Mitogen-activated protein kinase.

**Table 1 Comparison between different protocols to convert/derive human pluripotent stem cell into naïve stem cell**

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
|  | **Compounds (inhibitor of)** | **Conversion** | **Derivation**  | **XCI status** |
| **Gafni *et al*[49]** | PD0325901(MEKi)1, CHIR99021(GSKi)1, SP600125 (JNKi), SB203580 (MAPKi) , LIF1, IGF1, FGF2, TGFB1 | hESC/hiPSC | hESC/hiPSC | *XIST* RNA (-)H3K27me3 (-) XaXa |
| **Chan *et al*[50]** | PD0325901(MEKi)1, BIO (GSKi)1, Dorsomorphin (BMPi), LIF1,2 | hESC/hiPSCs | hiPSCs | nt |
| **Ware *et al*[51]** | PD0325901(MEKi)1, CHIR99021 (GSKi)1, FGF2, LIF1, Activin A, NSC74859 (STAT3i), SU5402(FGFRi)  | hESC/hiPSC | hESC | *XIST* RNA (-)XaXa |
| **Valamehr *et al*[52]** | PD 0325901(MEKi)1, CHIR99021(GSKi)1, Thiazovivin (ROCKi), FGF2, LIF1 , SB431542 (ALKi/TGFB1R) | - | hiPSC | *XIST* RNA (downreg.)H3K27me3 (<10%)XaXa |
| **Theunissen *et al*[55]** | PD0325901 (MEKi)1, CHIR99021 (GSKi)1, SB590885 (BRAFi), Y27632 (Rocki), WH-4-023 (pan-RTKi), LIF1, FGF, Activin A2 | hESC  | hESC/hiPSC | *XIST* RNA (+)XaXi |

1Compounds used in most works; 2Addition of MEFs. NT: Not tested; ALKi: Inhibitor of transforming growth factor-beta superfamily type I activing; TGFB1: Receptor-like kinase (ALK) receptors; BMPi: Inhibitor of bone morphogenic protein; BRAF1: Serine/threonine-specific protein kinase participating of RAS-RAF-MEK-ERK signaling transduction cascade; FGF: Fibroblast growth factor; FGF2: Fibroblast growth factor 2; FGFRi: Fibroblast growth factor receptor (FGFR)-specific tyrosine kinase inhibitor; GSK3: Fibroblast growth factor; FGF: Receptor tyrosine kinase and glycogen synthase kinase 3; GSK3i: Fibroblast growth factor; FGF: Receptor tyrosine kinase and glycogen synthase kinase 3 inhibitor; H3K27me3: Tri-methylation of lysine 27 on histone H3; hESC: Human embryonic stem cells; hiPSC: Human induced pluripotent stem cells; IGF1: Insulin-like growth factor 1; JNKi: Inhibitor of c-Jun N-terminal kinase; LIF: Leukemia Inhibitory Factor; MAPKi: p38 mitogen-activated protein (MAP) kinase inhibitor; MEK: Mitogen-activated protein kinase; MEKi: Mitogen-activated protein kinase inhibitor; Pan-RTKi: Receptor tyrosine kinase (RTK) Inhibitor; ROCKi: Inhibitor of Rho-associated coiled-coil containing protein kinase; STAT3i: Signal transducer and activator of transcription 3 inhibitor; TGFB1: Transforming growth factor beta 1; Xa: Active X chromosome; Xi: Inactive X chromosome; *XIST* RNA: RNA FISH signals from X-inactive specific transcript - XIST.