In Pursuit of Porcine Pluripotent Stem Cells for Autologous Cell Therapy

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Abstract

Treatments to repair the human heart following regenerative diseases remain a challenge for medical science. Unlike lower vertebrate species the human heart lacks a regenerative pathway meaning that research has to be focused on cell transplantation. Porcines (Sus scrofa) are excellent models for cardiovascular disease and pluripotent stem cells (PSCs) generated from porcines will provide important clinical insights for cardiac cell therapy. This could open a new avenue of research into degenerative conditions as porcine is a more effective human proxy to work with. However, bona fide PSCs are currently available only in rodents (mouse, rat) and primates (monkey, human). Attempts to derive pluripotent stem cells (PSCs) from porcine have been going on for more than two decades with slow progress. Despite the fact that the porcine stem cells are under increasing glare of publicity due to milestone achievements in this area of research. Advances in stem cell technology, especially the genetic engineering, innovative cell culturing and induced pluripotency to generate stem cells has dramatically revolutionized the basic and applied investigations and applications of porcine stem cells. This review attempts to summarize the major signaling pathways involved in maintenance of pluripotency and the state of the art conceptual and technical progress for generating bona fide porcine PSCs.

Keywords

Porcine Model, Signaling Pathways, Stem Cells, Induced Pluripotent Stem Cells, Pluripotency, Regenerative Medicine, Cell Therapy, Epigenetics

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1. Introduction

Stem cells have huge potential for alleviating suffering for many diseases of humans for which effective therapy is currently not available. The PSCs are the unicellular equivalent to a whole animal derived from either pre-(embryonic stem cells or ESCs) or post- (epiblast stem cells or EpiSCs) implantation embryos [1]. Under precise culture conditions, the PSCs are theoretically capable of differentiating into all cell types of an adult animal.

In the late 1960s, Gordon’s pioneering work on nuclear transfer using amphibian cells demonstrated that differentiation was a reversible process [2]. Decades later in 2006, Yamanaka demonstrated that ectopic expression of transcriptional factors (Oct3/4, Sox2, Klf4 and c-Myc) could induce pluripotency in terminally differentiated somatic cells, first in mouse [3], then in humans [3]. Rigorous testing using molecular and functional studies have clearly demonstrated that iPSC are similar to ESCs including germ line competence [4].

While, mouse and human embryonic stem cells have been in the limelight for the last two decades, limited efforts have been on porcine embryonic stem cells despite reports of first isolation in early 1990s [5] and subsequent numerous attempts to develop porcine ESCs (pESCs). This is mainly because the pig is anatomically and physiologically more similar to humans than the rodents or other model animals. Although the information is available on porcine ESCs (pESCs) after pioneering work on porcine and murine ESCs (mESCs), the stem cell lines developed were poorly defined [6]. In addition, the stem cells reported were found to lose pluripotency over a relatively shorter number of passages [7]. It was suggested that slow epiblast development with prolonged pre-implantation stages in comparison to murine or human, might hinder the growth and pluripotency of porcine stem cells [7].

Recent advances in with reprogramming technology have provided ability to generate porcine iPSCs. Putative porcine iPS cells fulfilling several criteria of true pluripotency were reported after reprogramming fibroblasts with viral vectors [8]-[10]. Notably, in one study, chimeric offspring were also reported. It has been demonstrated that human factors could be used in porcine mesenchymal stem cells (pMSCs) to generate iPSCs capable of generating chimeras with germ line transmission. After breeding of chimeric porcine, 2 out of 43 F-1 offspring carried some of the reprogramming factors in their genome, suggesting germ line transmission [11]. However, both piglets with germ line transmitted reprogramming factors died around birth. Moreover, lack of true porcine ES cell lines prevents the unequivocal identification of stem cells after reprogramming towards iPSC cells in this species. Reports are scarce on bona fide piPSC, and their maintenance by endogenous factors with silenced exogenous reprogramming factors [12].

The paucity of information related to morphology, pluripotency markers, and differentiation capabilities hampers a thorough evaluation of the validity of putative stem cell lines. Data from the literature suggests that similar regulatory pathways are likely to exist among different species. Coupling of these pathways with their distinct expression patterns, the relative concentrations of pluripotency-related molecules, and timing of embryo development, along with supportive micro-environmental conditions, would appear to vary in a species-specific manner. It is envisaged that the understanding of signaling pathways and other subtle, but consequential diversities may endow with valuable information for isolating the genuine porcine PSCs.

2. Signaling Pathways Involved in Maintenance of Pluripotency and Self-Renewal in Murine, Humans and Porcines

The two most important and distinguishing traits that categorize PSCs, are pluripotency and self-renewal that allow the stem cells to divide continually in an undifferentiated manner. Both the traits orchestrate by a complex cascade of signaling pathways. However, literature indicates the involvement of some common signaling pathways among mice, primates and porcine stem cells, though the growth factors equilibrium may differ among different species [13]. Understanding of various regulatory pathways involved in pluripotency of porcine PS cells is still not clear. Till date bon fide porcine pluripotent stem cells have not been established. Therefore, the exposition and understanding of the mechanism underlying pluripotency of human and mice PS cells would be the stepping stone towards the successful establishment of the culture conditions for the bona fide PS cells.

2.1. Murine and Human

It has already been established that mouse serum and the presence of the leukemia inhibitory factor (LIF: Interleukin 6 class cytokine) is required for maintenance of pluripotency in murine stem cells in vitro [14]. LIF binds
to specific LIF receptor (LIFR-α) at cellular surface, which forms a heterodimer with GP130 signal transducing unit (Figure 1). This leads to the activation of the JAK/STAT (Janus kinase/signal transducer and activator of transcription) and mitogen-activated protein kinase (MAPK) cascade. Once activated, the STAT3 is phosphorylated and subjected to dimerization, before being translocated to the nucleus where it acts as a transcription factor [15]. Withdrawal of LIF in culture medium pushes stem cells en route for differentiation, but they somehow maintain pluripotency. Hence, it is necessary to supplement the culture media with LIF to maintain murine stem cells in undifferentiated state. Genetically manipulated stem cells may remain undifferentiated in absence of LIF. For mouse ES cell pluripotency and self-renewal, STAT3 is indispensable [16] [17], and the downstream target of STAT3 is c-Myc [18]. LIF can sustain self-renewal only in the presence of serum (cheap proxy for BMP4). It means that key factor controlling this process is BMP (Bone Morphogenic Protein, a polypeptide related to the TGF-β superfamily of proteins). BMP4 binds to two distinct types of serine-threonine kinase receptors known as BMPR1 and BMPR2. Signal transduction via these receptors takes place via Smad and MAP kinase pathways to upshot transcription of its target genes. The downstream effectors of BMP4 are well-characterized SMAD1, 5 and 8 proteins. The role of BMP4 has been usually expressed more as controlling cell differentiation rather than promoting self-renewal [19] [20].

Modified form of murine stem cell culture conditions were effectively used to derive first monkey [21] and then human ESCs [22]. But so far, only a limited success has been achieved in other mammalian species. To understand the molecular mechanisms that contribute to pluripotency in vitro, mouse and primate PSCs are used as the main reference.

The LIF is not required for culturing and maintaining human ESCs [23]. However, LIF phosphorylates STAT3 in primate ES cells as in mouse ESCs [24]. In human PSCs, the TGF-β signaling pathway plays a key role in maintaining pluripotency and self-renewal (Figure 1). They signal via SMAD1/5 and activin/nodal branch which signal through SMAD 2/3/4. SMAD 1/5 branch transduces via type I receptors [Anaplastic lymphoma kinase (ALK) also known as ALK tyrosine kinase receptor], ALK1, ALK2, ALK3 and ALK6 (ACVERL1, ACVER1, BMPRIA and BMPR1B). TGF-β/activin/nodal involves activation of SMAD2/3 via ALK4, ALK5 and ALK7 (TGFBR1 and ACVR1C). On activation, via phosphorylation and alliance with common SMAD4, the receptor-activated SMADs enters into the nucleus where they regulate transcription of downstream genes [25].

In disparity to TGF-Beta/Activin/Nodal signaling, BMP signaling is incapable to sustain self-renewal and is linked with differentiation. In human PSCs, the BMP2 promotes extra-embryonic endoderm differentiation,
whereas the BMP4 induces differentiation into mesoderm and ectoderm. FGF2 (fibroblast growth factor-2) is proficient in maintaining human PSCs without serum and feeder cells via repression of BMP. FGF2 promotes self-renewal of Human PSCs by activating the PI3K pathway [26] [27].

Wnts (Wingless-Type MMTV Integration Site Family Members) proteins are also supposed to play significant role in maintaining pluripotency. In Canonical Wnt signaling, Wnt ligands bind to Frizzled receptor, which in turn activates Dsh (Dishevelled) that leads to inactivation of GSK-3β (glycogen synthase kinase-3β). GSK-3β negatively regulates the degradation of β-catenin which accumulates in cell nucleus and forms a transcriptionally active complex with T-cell specific factors (TCF) that activates target genes [28].

However, human PSCs cannot be maintained during long-term culture by activation of the Wnt pathway solely, and requires other extrinsic signals [29].

Apart from that, Nanog (NK2-family homeobox transcription factor) is reported to be vital for self-renewal in human PSCs. In comparison to Oct3/4 and Sox2, Nanog expression is elevated in human stem cells and is down regulated as cells undergo differentiation. Oct3/4 and Sox2 bind to the Nanog promoter to regulate Nanog transcription. Human PSCs display various signaling pathways implicated in self-renewal and pluripotency that are mutually dependent and exhibit array of cross-talk mechanisms [30].

Stem cells pluripotency and self-renewal capacity in both mouse and human are regulated by conserved transcription factors. However, downstream regulators are apparently not well conserved as compared to conventional pluripotency factors Oct4/Sox2 in both mouse and humans [31]. Substantial dissimilarities occur in the transcriptional networks and signaling pathways that regulates mouse and human PSC self-renewal and lineage development.

2.2. Porcine

Particular cell signaling pathways have been revealed to direct pluripotency in mouse and primates. The existing observations in porcine induced pluripotency and maintenance in murine and human stem cells advocate that, similar regulatory pathways might be conserved among them. However, species-related differences in the mechanisms controlling pluripotency are apparent. The cells isolated from primate pre-implantation of blastocyst, presumably originating from naive epiblast, as it occurs in mouse, spontaneously progress to the primed epiblast in vitro before giving rise to stable cell lines that despite the fact that have been named ESCs, are actually EpiSCs. EpiSC are different form typical ES cells in various aspects as listed in Table 1.

Pig epiblast has been shown to be dependent on activin/nodal signaling for self-renewal, as previously shown for human ESCs, indicating that maintenance of pluripotency by this signaling mechanism is conserved in mammals [32]. These authors demonstrated that pig EpiSCs express core pluripotency markers and maintain pluripotency via activin/nodal signaling pathway. Moreover, these cells could be induced to differentiate toward trophoderm and to germ cell precursors in response to BMP4 [32]. Blomberg et al., 2008, demonstrated inconsistent expression of LIF receptors in porcine epiblast cells cultured for 24 hr after separating them from inner cell mass (ICM) of the embryo [33]. Hall et al., 2009, confirmed that LIFR and bFGF are not expressed in epiblast, but within the trophoderm [34]. These findings revealed that cell signaling associated with maintaining pluripotency in human ESCs indicates the dispensability of LIF in supporting pluripotency in porcine species. However, Brevini et al., have shown that LIF is a key factor and it supports both attachment and self-renewal of stem cells [35]. They further hypothesized that LIF is unlikely to act through the GP130/LIFR/STAT3 signaling pathway, but rather via an alternative cascade involving phosphoinositide-3 kinase (PI 3K), serine/threonine protein kinase (AKT) (a key effector in the PI3K pathway) and phosphatase and tensin homolog deleted on chromosome 10 (PTEN) (a negative regulator of the same pathway), known to be responsive to LIF and has been previously shown to trigger the expression of NANOG and to facilitate efficient proliferation and survival of murine ESCs [36]. bFGF has been reported to be indispensable for proliferation, self-renewal and pluripotency of pig epiblast cells [37].

The presence of FGF receptor indicates the ability of porcine cells to bind bFGF and advocates a possible role of the FGF signaling pathway in self-renewal of porcine stem cells. Besides, it has been stated that bFGF can also bind and stimulate the PI3K/AKT cascade [38] and there is a possibility that bFGF and LIF acts in tandem to exert its pluripotency-related effect through this pathway as well as through the canonical one. Particularly, the PI3K/AKT cascade is known to modulate several regulatory pathways and to exert a potent triggering effect on Nanog expression which, in contrast to OCT4, is persistently transcribed and steadily detectable in pig pluri-
### Table 1. Comparative behaviour of naive and primed pluripotent state [1].

<table>
<thead>
<tr>
<th>Culture cell</th>
<th>Naive</th>
<th>Primed</th>
</tr>
</thead>
<tbody>
<tr>
<td>Embryonic tissue of origin</td>
<td>Mouse pre-implantation ICM</td>
<td>Mouse post implantation epiblast, primate pre-implantation ICM</td>
</tr>
<tr>
<td>Cell line</td>
<td>Rodent ESC</td>
<td>hESC, rodent EpiSC</td>
</tr>
<tr>
<td>Pluripotency in vitro</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Teratomas</td>
<td>Yes</td>
<td>Yes</td>
</tr>
<tr>
<td>Colony morphology</td>
<td>Domed</td>
<td>Flattened</td>
</tr>
<tr>
<td>Rapid self-renewal</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Slow self-renewal</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Response to LIF/BMP4</td>
<td>Self-renewal</td>
<td>None</td>
</tr>
<tr>
<td>Response to FGF2/ERK</td>
<td>Differentiation</td>
<td>None</td>
</tr>
<tr>
<td>Chimera formation</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Pluripotency markers</td>
<td>Oct4, Nanog, Sox2, Klf2, Klf4</td>
<td>Oct4, Sox2, Nanog</td>
</tr>
<tr>
<td>Single cell dissociation</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Clonigenecity</td>
<td>High</td>
<td>Low</td>
</tr>
<tr>
<td>X-inactivation</td>
<td>No</td>
<td>Yes</td>
</tr>
<tr>
<td>Response to 2i</td>
<td>Self-renewal</td>
<td>Cell death</td>
</tr>
</tbody>
</table>

potent cells. Altogether these revelations suggested that Nanog might be able to maintain pig pluripotent cells in an undifferentiated state, and may thus serve a vital molecule for self-renewal in porcine [32].

Evaluation of crucial signaling pathways like WNT, NOTCH, TGFβ1, VEGF and JAK-STAT in porcine, murine and human PSCs have indicated that the principal transcriptional network to uphold pluripotency and self-renewal in porcine had noteworthy resemblances to human but were diverse from that in murine. Moreover, porcine PSCs were found to be positive for prime state markers of Otx2 and Fabp7, which is the characteristic feature of human ESC and murine EpiSCs. However, porcine PSCs were lacking expression of specific naïve state markers like KLF2/4/5 [39]. These finding indicates that porcine PS cells are more like prime state cells.

Apart from that the bunch of imprinted genes were muzzled in porcine PSCs as earlier witnessed in murine PSCs that have inadequate potential to contribute to chimaeras [39]. These significant dissimilarities in imprinting and naïve state gene expression advocates that so far recognized poricne PS cell lines could be more analogous to primed state cells. Hence, the porcine PS cells seem to have the less potential to give rise to chimeras and cloned offspring’s [39].

### 3. Generation of Pluripotent Stem Cells (PSCs) from Porcine

Isolation of putative PSCs have been demonstrated from porcine blastocyst [33]. However, there is accord that these cell lines do not complete the needs of standard ESCs [40]. Alongside porcine ES cells, depiction of flattened cuboidal shape epithelial-like cells were also found [41]. The first few endeavors to establish primary cultures of epiblastic cells from 7 to 11 days post-conception blastocyst were carried out [42]. The lack of vimentin expression [42], high nuclear to cytoplasmic ratio and tendency to form clumps [5] [43] were used as the first gauge to assess presumptive ESCs phenotype. Alkaline phosphatase (AP), an established and reliable marker is used for the characterization of undifferentiated ESCs in various livestock species [44]-[46]. However, Wheeler et al., 1994 demonstrated the most stringent criterion of confirming pluripotency in the presumptive PSCs was their ability to produce chimeras [47]. This criteria to evaluate contribution of putative stem cell lines in chimera formation was not full proof, as they used coat color contribution and average daily gain in grams [47]. Interestingly, porcine ES cells isolated from day 24 - 25 embryos when cultured on inactivated MEFs (Murine embryonic fibroblasts) showed morphological resemblance to murine ESCs and differentiated into a wide range of cell types in vitro. In suspension culture, porcine ES cells formed embryoid bodies (EBs). The ES cells were able to differentiate and contribute to tissues of a chimeric piglet [48].

Apart from porcine ES cells, Chen and co-workers demonstrated the generation of pESCs from intact, early
hatched blastocysts and as well as isolated ICM (inner cell mass) of intermediate and late-hatched blastocysts. Furthermore, in vivo pluripotency of these cells was verified by birth of a chimeric piglet, as well as by pigmentation and DNA markers, and the ability to direct the development of nuclear-transfer embryos to the blastocyst stage. However, only one live piglet was born from 131 embryos transferred, which was chimeric in different tissues assessed by microsatellite markers. Unfortunately germ line contribution was not observed [49]. Primary explants were isolated from in vitro produced (IVF) blastocysts, and the cells were injected into the blastocysts. The injected cells could contribute to embryos at the blastocyst stage [50].

Furthermore, pESCs have also been generated from epiblast of elongating embryos [32] as well as from ICM of early blastocysts using a cocktail of growth factors [51]. However, the status of these cells (naive or primed) and their germline transmission has not been demonstrated. ICM cells derived porcine ESC lines have been established using inhibitors of glycogen synthase kinase 3β and MAPK1 [52]. These lines were derived in standard hESC culture medium but later transferred to ESC medium supplemented with glycogen synthase kinase 3β and MAPK1. Morphology of these colonies was reported to be analogous to mouse ESCs.

While till date there is no established bona fide pESC reported, multiple reports have shown derivation of piPSCs from a variety of somatic tissues as listed in Table 2. Interestingly, porcine induced pluripotent stem cells (piPSC) have been generated using mouse [8] or human [9]-[11] transcription factors as mentioned in Table 2.

Table 2. Summary of porcine induced pluripotent stem cells (iPSCs) generated from various somatic cell. The abbreviations used have already been explained in the text.

<table>
<thead>
<tr>
<th>Cell type</th>
<th>RF</th>
<th>Mode of delivery</th>
<th>Medium supplement</th>
<th>Transgene activation</th>
<th>Teratoma</th>
<th>Chimeras</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Foetal fibroblasts</td>
<td>OSKM</td>
<td>Lentiviral vector</td>
<td>bFGF</td>
<td>Not tested</td>
<td>Yes</td>
<td>No</td>
<td>[10]</td>
</tr>
<tr>
<td>Embryonic fibroblasts</td>
<td>OSKM</td>
<td>Retroviral vectors</td>
<td>bFGF</td>
<td>Not tested</td>
<td>Yes</td>
<td>No</td>
<td>[8]</td>
</tr>
<tr>
<td>Mesenchymal stem cells</td>
<td>OSKMLN</td>
<td>Lentiviral vectors</td>
<td>bFGF</td>
<td>Not tested</td>
<td>No</td>
<td>Yes, 29 offsprings</td>
<td>[11]</td>
</tr>
<tr>
<td>Adult fibroblast Bone marrow cells</td>
<td>OSKMLN</td>
<td>Lentiviral vectors</td>
<td>Doxycillin inducible Transgene expression</td>
<td>Not tested</td>
<td>Yes</td>
<td>No</td>
<td>[9]</td>
</tr>
<tr>
<td>Adult fibroblasts</td>
<td>SKM</td>
<td>Retroviral vectors</td>
<td>bFGF, LIF</td>
<td>Not tested</td>
<td>Yes</td>
<td>No</td>
<td>[71]</td>
</tr>
<tr>
<td>Foetal fibroblasts</td>
<td>OSKM</td>
<td>Transposon mediated</td>
<td>bFGF</td>
<td>Not tested</td>
<td>Yes</td>
<td>No</td>
<td>[73]</td>
</tr>
<tr>
<td>Embryonic fibroblasts</td>
<td>OSKM</td>
<td>Non-viral</td>
<td>bFGF</td>
<td>Not tested</td>
<td>No</td>
<td>No</td>
<td>[70]</td>
</tr>
<tr>
<td>Embryonic fibroblasts</td>
<td>OSKM</td>
<td>Retroviral</td>
<td>Porcine LIF</td>
<td>Not tested</td>
<td>Yes</td>
<td>Yes</td>
<td>[58]</td>
</tr>
</tbody>
</table>

Abbreviations: RF: Reprograming factors; O: Oct4; S: Sox2; K: Klf4; M: c-Myc; L: Lin28; N: Nanog.
Numerous methods of generating induced pluripotent stem cells. Only few representative methods are shown here.

larly, in pig the Oct4 is present in the ICM, trophectoderm, and during hatching it is expressed in the epiblast component of the embryonic disc [59] [62]. Apart from this there is a gradual down-regulation of Oct4 expression noted in porcine ESCs [63].

Like Oct-4, Nanog too shows marked differences in porcine species as compared to human or rodent systems. In porcine, Nanog expression is not only limited to the ICM and epiblast [64] but its expression could be detected in day E7.5 to day E10.5 embryos, which is five days later blastocyst formation [34]. Unlike the mouse, expression of Nanog is restricted to only a few cells of the ICM around implantation [65]. These observations clearly suggest that different species have different timing of Nanog expression. Nevertheless, chimeric animals can be generated by injecting porcine ICM clumps into blastocyst which suggests that few cells in porcine ICM are pluripotent or attain pluripotency [66]. Somehow, putative porcine pluripotent stem cells lose their pluripotency over a relatively short number of passages as compared to their mouse or human counterparts. These observations clearly suggest that other factors may be responsible for the maintenance of self-renewal of these cells in an undifferentiated status.

Porcine iPS cells derived from embryonic fibroblasts in serum containing medium morphologically resemble human ES cells with flatten, compact colonies that stain positively for AP, SSEA-4, Nanog and Rex1 [8]. Porcine induced pluripotent stem cells (piPSCs) derived from primary ear fibroblasts or bone marrow cells and ESC-specific culture medium without bFGF, also express human ES cell surface markers [9]. However, iPS cells generated from porcine fetal fibroblasts cultured in hESC-specific medium supplemented with bFGF stain positive for AP and SSEA-1, and negative for SSEA-3, SSEA-4, TRA1-60 and TRA1-81 [10], which is very unusual compared to other stem cells cultured in hESC medium. The patterns of methylation in iPSCs generated also resemble the patterns observed in ESCs, especially for genes related to pluripotency, such as Oct4. Nevertheless, all these iPSCs differentiate into cells representing the three germ layers via EB and teratoma formation. The use of episomal factors in combination with 2i has also proven successful for piPSCs generation [67]. The cells display similarities to their LIF-dependent ESCs counterparts [68]. In spite of that, the cell lines showed either vector integration or persistence of episomal plasmids even after several passages [67].

The most challenging criterion of pluripotency was accomplished by West, and his colleagues, where 30 of 36 pigs produced were chimeric, displaying varying levels of chimerism. In this study, the forced expression of transcription factors enabled the derivation and establishment of iPSCs in pig that have displayed pluripotency [11] [69]. LIF-dependent naive pluripotent cells were generated from cultured porcine ICM, using combinations of DOX-driven Oct4, Klf4 human transgenes and stringent culture conditions but there was no evidence of germ-line transmission.

The PSCs derived from embryos and iPSCs generated from embryonic fibroblasts in porcine model has been found to possess a primed pluripotent state analogous to that of mEpiSCs or hESCs, rather than to that of mESCs [70].Montserrat et al., exhibited that over expression of exogenous Oct4 directly or indirectly are not critical to attain and uphold pluripotency in the generation of porcine PSCs [71]. These authors were able to establish PSCs without Oct4 over-expression and by ectopic expression of Sox2, Klf4 and c-Myc. However, in vitro produced porcine blastocysts express Oct 4 both in ICM and trophectodermal cells [59] [62].
Studies have demonstrated that piPSCs generated from fibroblast using OSKM, displayed mES like morphology when they were maintained in LIF/STAT3-dependent manner and they exhibited potential of generating chimeric and reconstructed NT (Nuclear Transfer) embryos [72]. However, in this study, the transgenes were clearly down regulated, but were not silent completely, and it was in agreement with previous studies [70] [71]. Kues et al. displayed the utility of porcine Oct4-EGFP cells and transposon mediated reprogramming for the derivation of porcine iPS cells. However, this study also indicated the limitation of culture conditions and culture media to support pluri potency [73]. Petkov et al., demonstrated that the outcome of porcine somatic cell reprogramming experiments depends on the selection of expression vector promoter because it controls the expression levels of the reprogramming transcription factors. In this study EF1a and CAG promoters were found to be more efficient than doxycycline-induced tet operator (TetO) promoters [74]. These finding suggest that the endogenous pluripotent genes and their networks could not sustain pluripotency of putative porcine PSCs. Thus, establishment and optimization of culture conditions specific for porcine pluripotent stem cells is prerequisite for any further advancement in porcine stem cell research.

3.1. Culture Conditions

It is evident from earlier reports that culture conditions required to sustain pPSCs have not been well established. The stumbling block faced by laboratories worldwide has been that putative PSCs lose their pluripotency over a relatively short number of passages. Bona fide Porcine PSCs in which pluripotency maintained by the endogenous pluripotency factors have not yet been reported [12]. In other words, the incessant expression of the exogenous transgenic transcription factors is still obligatory to sustain the pluripotent phenotype of porcine iPSC-like cells [9]-[11].

Over the past few decades, all endeavours to generate livestock PSCs have failed. Researchers have used mouse or human protocols as reference for culturing of ungulates PS cells, which include feeders, serum and supplementation with LIF [45] [75] [76] or FGF2 [75], epidermal growth factor and stem cell factor (SCF) [76].

Usually for the pPSC culturing, standard human stem cell medium [69] or mouse and human standard culturing medium are mixed in 1:1 ratio [8] [77] because of the paucity of the information related to presence of receptors on pPSCs [35] demonstrated that porcine pluripotent cell lines do not express LIF receptors [13]. However, they do express FGFR-2 [13]. Apart from that their finding also suggests that LIF is not essential for the maintenance of pluripotency, but its presence inhibits the differentiation process. They also indicated that LIF do not act through LIF receptor signaling pathway.

Researchers also tried to culture ICM and epiblast in feeder layer and serum-free culture conditions. A study was based on modified TeSR medium and matrigel. Almost half of the primary cultures got differentiated within 3 - 4 passages. They used mechanical passaging for sub-culturing [64]. However, mTeSR1 based on Ludwig et al., is most widely used feeder and serum free culture medium for hESCs and iPSCs [78].

Vassiliev et al., cultured ICM cells in MEM medium supplemented with EGF (epidermal growth factor) and activin [51]. They were able to generate chimeras by using putative porcine ES cells. But it was not vivid from their study whether they were able to keep the putative pESCs in undifferentiated self-renewal state beyond 14 passages [51]. Various type of culture conditions were used to culture ICM/epiblast primary outgrowths [26]. But, none of the culture conditions supported to keep the putative porcine ES cells in undifferentiated state after certain passages.

A very different culture regime i.e., 2i/LIF medium led to the first isolation and establishment of bona fide ESCs from rat embryos [53] [54], and noncompliant mouse strains [79]. Human embryo-derived stem cells cannot be maintained in 2i/LIF alone [80]. However, it facilitated to establish mouse ESC-like cells in presence of forskolin [55]. Consequently, human cells with an epigenetic and transcriptional profile analogous to naïve ESC lines could be established.

Haraguchi et al., and colleagues were able to generate presumptive porcine ES cell lines from the ICM of porcine embryos by using inhibitors of glycogen synthase kinase 3β and MAPK1. They were able to maintain the cell lines over 100 passages only [52].

We used the 2i (GSK 3β inhibitor, and MAPK inhibitor)/LIF culture regime (chemically defined feeder-free culture without supplementing exogenous growth factors) for deriving bovine PSCs [46]. Under these minimal conditions, Verma et al., investigated the subsistence of a pluripotent ground state, stabilized through shielding from FGF/MAPK signaling and Wnt activation, in bovine ICM-derived cell cultures. The GSK 3β facilitate Wnt3a signaling pathway, and the deficiency of the Wnt signal roots GSK3β to inactivate targets, such as
β-catenin and c-Myc. CHIR99021, a GSK3β inhibitor, effects Wnt signaling activation [81] [82]. These findings were in conjugation with several previous reports, where they have shown that CHIR99021 augments the reprogramming efficiency [82].

Rodriguez et al., demonstrated that blocking MEK signaling increases the proportion of Nanog expressing cells in the ICM of porcine embryos [83]. From their findings, it is apparent that FGF signaling participates in the formation of the founders of the ICM. PDO325901 (MEK pathway inhibitor), CHIR (GSK3β inhibitor) and LIF supplementation were used to modulate pluripotency in iPSCs cells to acquire features of naive/ground pluripotency. It was demonstrated that small molecule inhibitors can be used to increase the homogeneity of iPSC cultures [83]. Oxygen tension is also an imperative trait for PSC’s maintenance and differentiation. The iPSCs induction executed in hypoxic conditions (5% O2) demonstrated augmented reprogramming efficiency [84].

3.2. Epigenetic Modifiers

Apart from mRNA and episomal vector approach to safely reprogram somatic cells, another approach is usage of concoction of small molecules that are allied with epigenetic modifiers and key signaling pathways listed in Table 3. Small molecules that alter chromatin and gene expression enhance the reprogramming efficiency in iPSC generation. Inhibitors of histone deacetylases (HDACs), histone demethylases (HDMs), and histone methyltransferases (HMTs), which regulate chromatin remodeling, have been recognized as small molecules for terminally differentiated cells reprogramming. For instance, Huangfu et al., used chromatin modification inhibitors, the 5’-azacytidine (5-aza) (DNA methyltransferase inhibitor), dexamethazone (synthetic glucocorticoid) and valproic acid (VPA, a histone deacetylase inhibitor) to investigate their effects on reprogramming efficiency [85]. It was inferred that 5-aza, dexamethazone and VPA enhanced the efficiency of cellular reprogramming by 10-fold, 2.6-fold and >100-fold respectively. The effect of VPA was found to be much stronger than that of 5’-azacytidine and other HDAC inhibitors tested [85]. Other noticeable epigenetic modifiers which are used during the reprogramming are, parnate, a histone demethylase inhibitor [86], 5-aza and RG108, a DNA methyltransferase inhibitors [87]. Butyrate (a HDAC inhibitor) also improves iPSC generation by augmenting acetylation of histone H3 and demethylation of genes promoter, related to of pluripotency [88] [89].

Lee et al., demonstrated that Sirtuin 1 (SIRT1) (a member of the sirtuin family of NAD+-dependent protein deacetylases) a class III HDAC assists in generation of iPSCs from MEFs via miR-34a and p53 Pathways [90]. It was hypothesized that SIRT1 works as a positive epigenetic regulator in reprogramming somatic cells.

Korean scientists discovered a new molecular compound, RSC133 (inhibits histone deacetylase and DNA methyltransferase) that could enhance the reprogramming efficiency of human adult cell to iPSCs. It was inferred that compound derivative acts as the “booster of pluripotency”, and it potently improves the reprogramming of human somatic cells into a pluripotent state and aids the growth and maintenance of human pluripotent stem cells [91].

Even in livestock, there are several reports that suggest that the reprogramming process is mired by faulty epigenetic modifications during the reprogramming process, resulting into low survival rates among clones [92] and inadequacy of iPS reprogramming. Histone deacetylation is considered to be an unwanted event in nuclear reprogramming assays. Trichostatin A (TSA) has been used to improve cloning efficiency by as much as 75% in various species [92].

Kim et al., have shown that the usage of decitabine (5-aza-2’-deoxycytidine) in ESC culture improves the number of colonies obtained [93]. Treatment of early pre-implantation murine and bovine embryos with 5-AzaC exhibited a simple and efficient method for isolation of putative bovine PS cells [94]. However, there are still safety concerns regarding the usage of epigenetic modifiers or cocktail of small molecules as they could activate cell growth pathway or inhibit tumor suppressor pathways.

3.3. Post-Translational Modulation of Pluripotency Genes

Ever since the first report of the existence of miRNA [95], approximately thousand types of microRNAs (miRNAs) documented till date, small RNAs have been identified in humans. The typical feature of stem cells is self-renewal and abnormal pluripotency was reported in ES cells lacking DGCR8 or Dicer [96] [97] which implies their vital function in the biosynthetic pathway of miRNA. Moreover, it’s been showed that miRNA have reduced expression in differentiated ES cells. In addition reprogramming factors and chemicals as epigenetic
<table>
<thead>
<tr>
<th>Names</th>
<th>Mode of action</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Forskolin</td>
<td>Stimulates adenylate cyclase activity and increases cAMP</td>
<td>[58]</td>
</tr>
<tr>
<td>Valproic acid</td>
<td>Histone deacetylase inhibitor</td>
<td>[85]</td>
</tr>
<tr>
<td>BIO (6-Bromindirubin-3’-oxime)</td>
<td>Inhibitor for GSK-3 α/β in the Wnt signaling pathway.</td>
<td>[94]</td>
</tr>
<tr>
<td>Lithium</td>
<td>GSK3 Inhibition (Wnt+)</td>
<td>[26]</td>
</tr>
<tr>
<td>CHIR99021</td>
<td>GSK3 inhibitor</td>
<td>[46]</td>
</tr>
<tr>
<td>Trichostatin</td>
<td>A histone deacetylase inhibitor</td>
<td>[85]</td>
</tr>
<tr>
<td>Sodium Butyrate</td>
<td>Sodium butyrate is a known inhibitor of histone deacetylases and inhibits both the mRNA and protein content of cyclin D1</td>
<td>[85]</td>
</tr>
<tr>
<td>Parnate</td>
<td>Histone demethylase inhibitor</td>
<td>[86]</td>
</tr>
<tr>
<td>5-azacytidine</td>
<td>DNA methyltransferase inhibitor</td>
<td>[85]</td>
</tr>
<tr>
<td>SC1 (Pluripotin)</td>
<td>Can act as a LIF replacement molecule for mouse embryonic (mES) cell self-renewal</td>
<td>[113]</td>
</tr>
<tr>
<td>RG108</td>
<td>DNA methyltransferase inhibitor</td>
<td>[87]</td>
</tr>
<tr>
<td>PD0325901</td>
<td>Selectively binds and inhibits MEK</td>
<td>[46]</td>
</tr>
<tr>
<td>SIRT1</td>
<td>A class III HDAC assists in Generation of iPSCs</td>
<td>[90]</td>
</tr>
<tr>
<td>RSC133</td>
<td>Inhibits histone deacetylase and DNA methyltransferase</td>
<td>[91]</td>
</tr>
<tr>
<td>RepSox</td>
<td>Tgfbr1 kinase inhibitor</td>
<td>[114]</td>
</tr>
<tr>
<td>Etoposide</td>
<td>Notch1 upregulation</td>
<td>[109]</td>
</tr>
<tr>
<td>Cytochalasin D</td>
<td>Wnt5a upregulation</td>
<td>[109]</td>
</tr>
<tr>
<td>Vitamin C</td>
<td>Nutrient vital that lower reactiveoxygen species</td>
<td>[110]</td>
</tr>
<tr>
<td>Chaetocin</td>
<td>Histone Methyltransferase Inhibition</td>
<td>[109]</td>
</tr>
<tr>
<td>Pifithrin-α</td>
<td>Inhibitor of p53-dependent apoptosis. Moreover, reduced p53 activity augments reprogramming efficiency of human and mouse somatic cells</td>
<td>[111]</td>
</tr>
<tr>
<td>Pifithrin-μ</td>
<td>Inhibitor of p53 mitochondrial pathway by reducing its affinity for antiapoptotic proteins Bcl-2 and Bcl-XL but does not affect any additional transcriptional functions of p53</td>
<td>[112]</td>
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<tr>
<td>DNMT1</td>
<td>Inhibition promote fully reprogramming</td>
<td>[79]</td>
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<td>HDACs</td>
<td>Inhibition increase the efficiency of reprogramming</td>
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<tr>
<td>G9a</td>
<td>Increase the efficiency of reprogramming</td>
<td>[53]</td>
</tr>
</tbody>
</table>

modifiers, the miRNAs play a significant part in the reprogramming of terminally differentiated cells.

Several miRNAs have been shown to enhance iPSC reprogramming when expressed along with combinations of the OSKM factors. Introduction of miRNAs specific to ESCs improve the generation of iPSCs in mouse. The miRNAs miR-291-3p, miR-294 and miR-295 augment the reprogramming efficiency [98]. The miRNAs did not improve reprogramming efficiency in the presence of c-Myc transgene; it suggests that they could be downstream target of c-Myc. These miRNA seems to have the ability to control cell cycle, to effect the augmentation of iPSC reprogramming [98].

Oct-4 and Sox2 are the decisive factors requisite for iPSC reprogramming and the miR302/367 clusters has been revealed to be the direct target of these two key genes [99]. Morrisey’s group demonstrated that expression of the miR302/367 cluster reprograms mouse and human somatic cells to pluripotent state very efficiently, in the absence of exogenous transcription factors or Yamanaka factors [100]. This miRNA-based reprogramming regime was found to be more efficient than conventional OSKM-mediated methods. Moreover, mouse and human miR302/367 iPSCs demonstrated features including pluripotency marker expression, teratoma formation and for mouse cells, chimera contribution and germine contribution was found to be analogous to Oct4/Sox2/Klf4/c-Myc iPSCs [100]. It was also shown that miR367 expression is requisite for miR302/367-mediated reprogram-
V. Verma et al.

ming and stimulates expression of Oct4 gene, and that suppression/degradation of HDAC2 by using Valproic acid is also vital [100]. As far as usage of miRNA for reprogramming is concerned, there is no authentic report available.

3.4. Differentiation Potential of Presumptive Porcine PS Cells

Despite non-stabilized defined state of pluripotency of pPSCs [74], in vitro differentiation of presumptive porcine iPSC into rod photoreceptors and their integration into the retinas of recipient pigs has been reported [101]. The impending therapeutic benefits of stem cell-based treatment to recover the cardiac function have been customary in small animal models of myocardial infarction (MI) [102]-[106]. Human PSCs transplanted to treat immune-deficient porcine or murine models of acute myocardial infarction (AMI) [107], but intra-myocardial injection of livestock stem cells, particularly porcine PS cells in a pig model of AMI has not yet been gauged. Animal models that imitate human cardiac disease, such as MI, AMI and ischemia-reperfusion (IR) that induces heart failure are useful models to study cardiovascular disease (CVD). Recently, a group of researchers examined the possibility of using porcine PS cells to prevent LV dysfunction and post in infarction left ventricular remodelling in hearts suffering from AMI. They demonstrated that porcine PS cell could be an effective treatment for myocardial infarction (MI). In this study, engrafted porcine PS cells showed the differentiation potential by differentiating into vessel cells, which further augmented the formation of new vessels in infarcted heart [108]. However, no observe myocyte generation was observed. piPSCs may have the prospective to be used in ischemic cardiac diseases. This study had some serious limitations and further research is required to translate cell therapy form bench to bed side. However, a porcine acute myocardial infarction (AMI) model can fill the space between treatment for small animals and treatment for humans.

4. Future Directions/Opportunities and Challenges

It is clear that pig and humans share far more anatomical, histological, biochemical and physiological properties than do the mice and humans. As a result, the pig has emerged as an excellent model for cardiovascular research and many other basic and applied biomedical applications.

The supply of human organs and tissues will be scarce to satisfy burgeoning demands of organ transplantation, making xenotransplantation a viable alternative. In this concern, pig is a promising effective translational model, and represents a candidate species for studying CVD, besides several other applications in regenerative medicine. Bona fide stem cells generated from porcine could be used in multiple applications such as creating models for human genetic diseases, engineering organs for transplantation therapies.

Understanding and unravelling the basic cellular and molecular mechanisms that regulate cell-mediated tissue regeneration is the key to harness this potential. Future research in the field could open novel prospects of research into degenerative conditions as pig is a more effective human proxy to work with. The research on porcine stem cells will lay a foundation for future studies of stem cell transplantation and regenerative medicine. In vitro culturing of porcine stem cell-derived cardiomyocytes could be transplanted into porcine MI model, which would give the researchers and clinicians valuable insights for cell therapy. There is an urgent need to develop validated reagents and cell culture methods to enhance use of porcine stem cells in regenerative medicine. To bestow important clinical insights for cell therapy or transplantation, necessary investigation is required in large animals.

The general ethical and logistical issues impede the use of humans or human-derived tissues for research and discovery pertaining to biomedical applications, which highlight the need for research model that closely mimic human anatomy, physiology, disease and injury processes. In view of significant differences between rodents and humans, the murine (rat and mice) confound or prohibit their use in translational studies.

The pig will continue to be ideal species to serve this purpose. The pigs conserve immunological and physiological attributes of human hearts apart from resemblance with the human heart. Hence, they can provide a potent research tool for pre-clinical study and particularly for transplantation medicine.

The Failure to identify or validate bona fide porcine/livestock PSCs is still a bewildering issue because of the several discrepancies in the presumed markers reported in various studies as listed in Table 4. On contrary, pig and humans share far more anatomical, histological, biochemical and physiological properties than mice and humans. The study of stem cells is one of the most exciting areas of contemporary biomedical research across
Table 4. Synopsis of the characterization of porcine iPSCs (induced pluripotent stem cells) from various laboratories.

<table>
<thead>
<tr>
<th>Mode of delivery and characterization of porcine iPSCs</th>
<th>[8]</th>
<th>[10]</th>
<th>[9]</th>
<th>[11]</th>
<th>[70]</th>
<th>[68]</th>
<th>[71]</th>
<th>[74]*</th>
<th>[74]#</th>
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<tbody>
<tr>
<td><strong>Delivery method</strong></td>
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<tr>
<td>Viral</td>
<td>Oct4</td>
<td>ND</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Viral</td>
<td>Sox2</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td><strong>Pluripotency markers</strong></td>
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<tr>
<td>Viral</td>
<td>Nanog</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Viral</td>
<td>Rex1</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>ND</td>
<td>+</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Viral</td>
<td>CDH1</td>
<td>ND</td>
<td>+</td>
<td>+</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>Viral</td>
<td>SSEA1</td>
<td>ND</td>
<td>+</td>
<td>−</td>
<td>+</td>
<td>−</td>
<td>+</td>
<td>ND</td>
<td>−</td>
</tr>
<tr>
<td>Viral</td>
<td>SSEA3</td>
<td>ND</td>
<td>−</td>
<td>+</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>ND</td>
<td>ND</td>
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<tr>
<td><strong>Surface markers</strong></td>
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<tr>
<td>Viral</td>
<td>SSEA4</td>
<td>+</td>
<td>−</td>
<td>+</td>
<td>−</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>−</td>
</tr>
<tr>
<td>Viral</td>
<td>Tra-1-60</td>
<td>ND</td>
<td>±</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>ND</td>
<td>+</td>
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<tr>
<td>Viral</td>
<td>Tra-1-81</td>
<td>ND</td>
<td>−</td>
<td>+</td>
<td>−</td>
<td>−</td>
<td>−</td>
<td>ND</td>
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<td><strong>Functional assays</strong></td>
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<tr>
<td>Viral</td>
<td>Teratoma</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>ND</td>
<td>+</td>
<td>+</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>Viral</td>
<td>EBs</td>
<td>ND</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Viral</td>
<td>Chimera</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
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*Tet O gp; Ef1a and CAG gp; ND: Not Determined.

The globe. So far, generation of bona fide porcine PSCs has proven elusive, despite overwhelming knowledge and availability of numerous types of epigenetic modifiers, small molecules, specialized reagents and culture details from mouse and human stem cells work. Presumably, this information cannot directly extrapolate to other species as such. Building on the available knowledge, we can innovate on techniques and ideas very specific to porcine species.

5. Conclusion

The establishment of methods for generating porcine stem cells for clinical applications is still a work in progress. Porcine pre and peri-implantation process as well as mechanism behind different cell fate specification is considerably dissimilar from mouse and human ones, that’s why presumptive porcine PS cells behave differently from existing human and mouse stem cell lines. Consequently, a comprehensive understanding of species specific developmental processes, signaling pathways and other related transcription factors would tender a roadmap for the generation of bona fide PSCs and their in vitro differentiation into cardiomyocytes. Until now none of the research group has documented successful differentiation of pPSCs into cardiomyocytes. To recapitulate, porcine PSCs will fill the vacuum between mouse and human pluripotent stem cells. However, noteworthy technical obstacles for the generation of bona fide porcine PS cells still persist that will only prevail over via ages of exhaustive research.

Disclosure of Potential Conflicts of Interest

The authors indicate no potential conflicts of interest.

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