Pluripotent stem cells have been proposed as an unlimited source of pancreatic β cells for studying and treating diabetes. However, the long, multi-step differentiation protocols used to generate functional β cells inevitably exhibit considerable variability, particularly when applied to pluripotent cells from diverse genetic backgrounds. We have developed culture conditions that support long-term self-renewal of human multipotent pancreatic progenitors, which are developmentally more proximal to the specialized cells of the adult pancreas. These cultured pancreatic progenitor (cPP) cells express key pancreatic transcription factors, including PDX1 and SOX9, and exhibit transcriptomes closely related to their in vivo counterparts. Upon exposure to differentiation cues, cPP cells give rise to pancreatic endocrine, acinar, and ductal lineages, indicating multilineage potency. Furthermore, cPP cells generate insulin+ β-like cells in vitro and in vivo, suggesting that they offer a convenient alternative to pluripotent cells as a source of adult cell types for modeling pancreatic development and diabetes.
expensive to conduct, often resulting in low differentiation efficiencies, partly due to the variability inherent in long, multi-step differentiation protocols that seek to recapitulate the entire developmental history of a β cell. These issues are exacerbated when such protocols are applied to genetically diverse human embryonic stem cells (hESCs) and induced pluripotent stem cells (hiPSCs). A potential solution is to initiate differentiation from an alternative cell type that is developmentally more proximal to the β cell. The obvious candidate is the PDX1+SOX9+ progenitor cell population of the emerging pancreatic bud that is capable of extensive proliferation and gives rise to all of the mature functional cells of the pancreas.

We describe a cell culture platform that enables hESC- and hiPSC-derived pancreatic progenitors to be captured and expanded in vitro. These cultured pancreatic progenitor (cPP) cells express key pancreatic transcription factors, including PDX1 and SOX9, and are stable for >25 passages. Crucially, cPP cells are closely related to their in vivo counterparts at the transcriptome level and can be differentiated into cells of the endocrine, acinar, and ductal lineages. Therefore, cPP cultures represent a convenient alternative system for studying pancreatic development that circumvents the need to repeatedly generate progenitors from pluripotent cells by directed differentiation. Finally, replacing heterochronic differentiation cultures with comparatively stable and homogeneous cPP cultures will facilitate the development of more robust protocols for the generation of pancreatic cell types from genetically diverse patient-specific hiPSC lines.

RESULTS

Maintenance and Expansion of cPP Cells Derived from hESCs and hiPSCs

Directed differentiation guided by growth factors and small molecules facilitates the generation of diverse cell types from pluripotent stem cells. We chose to produce pancreatic progenitors from hESCs and hiPSCs (Figure S1) using reagents based on the early stages of a protocol designed to generate mature β cells (Figure S2A; Rezania et al., 2014). This differentiation strategy induced the sequential expression of PDX1 followed by NKX6-1 and yielded a median of 80% PDX1+NKX6-1+ cells after 15 days (PPd15 cells; Figures 1B and S2C). However, as is often observed during directed differentiation from pluripotent cells, the kinetics of PDX1 and NKX6-1 expression varied between cell lines (Figure S2B) (Cahan and Daley, 2013). Therefore, we sought to capture, synchronize, and expand PPD15 cells in culture.

The 3T3-J2 mouse embryonic fibroblast cell line has been used to culture progenitor cells derived from a variety of human tissues, including endoderm-derived intestinal stem cells (Rheinwald and Green, 1975a, 1975b; Wang et al., 2015). We therefore determined whether pancreatic progenitor cells could be similarly expanded, if provided with appropriate stimuli. We tested a series of signaling agonists and inhibitors previously shown to regulate pancreatic development, including EGFL7, BMP4, nicotinamide, LIF, WNT3A, R-Spondin-1, Forskolin (cAMP agonist), GSK3β inhibition (CHIR99021), and inhibitors of BMP (LDN-193189) and SHH (KAAD-cyclopamine) signaling. Ultimately, a combination of EGF, retinoic acid, and inhibitors of transforming growth factor β (TGF-β, SB431542) and Notch signaling (DAPT) was found to support long-term self-renewal of pancreatic progenitors (Figure 1A).

To establish stable cPP cell lines, PPD15 cells were replated on a layer of 3T3-J2 feeder cells in the presence of these factors. Thereafter, cPP cells were routinely passed once weekly as aggregates at an average split ratio of 1:6, although they were also capable of forming colonies at clonal density (Figure 1C). This suggests a doubling time of ~65 hr in culture, similar to the 61 hr we routinely observe for hESCs when cultured on a layer of mouse embryonic fibroblasts.

We were able to generate self-renewing cPP cell lines from four different genetic backgrounds using two hESC (H9 and HES3) and three hiPSC cell lines (AK5-11, AK6-8, and AK6-13 derived in house); these diverse cPP cells expressed comparable levels of genes encoding key pancreatic transcription factors, including PDX1 and SOX9 (Figure S2D). Two cPP cell lines selected for further analysis (H9#1 and
AK6-13) have been maintained in culture for >20 passages to date enabling >10^{18}-fold expansion over 20 weeks. Crucially, cPP cells can be frozen and thawed with no apparent loss of proliferation or viability, suggesting cPP cells could replace pluripotent cells as a starting point for further differentiation to mature pancreatic cell types such as insulin-secreting β cells.

To determine whether cPP cultures consist of a stable and homogeneous population of cells, we measured the expression of key pancreatic transcription factors at the mRNA and protein levels. Gene expression of numerous markers of pancreatic bud cells, including PDX1 and SOX9, remained constant over extended periods in culture, indicating that our culture conditions maintain a stable population of pancreatic progenitors (Figure 1D). To determine whether cPP cultures represent a homogeneous population, we carried out immunostaining for a selection of pancreatic markers and found these to be expressed near ubiquitously at the protein level (Figure 1E). Furthermore, flow cytometric analysis showed that approximately 85% of cPP cells were PDX1+ (Figure 1F).

However, NKX6-1 expression was rapidly downregulated in culture, and NKX6-1 protein was not detected by immunostaining. Furthermore, we were able to establish cPP cell lines from day 7, 10, and 15 differentiation cultures (data not shown), the earliest time point being prior to expression of NKX6-1 and suggesting that cPP culture conditions stabilize pancreatic progenitors in a developmental state that precedes NKX6-1 activation. Very few cells were NGN3+, which marks early endocrine progenitors, indicating that differentiation was blocked at the progenitor stage under our culture conditions. Finally, chromosome counting showed that five out of six cPP cell lines carried 46 chromosomes without signs of structural changes, such as presence of fragments or dicentric chromosomes (Figure 2A). Multiplex fluorescence in situ hybridization (M-FISH) analysis on the AK6-13 line at passage 20 confirmed the absence of chromosomal abnormalities at significantly higher resolution than chromosome counting alone. M-FISH of passage 20 AK6-13 cPP cells failed to detect aneuploidy, translocations or deletions in 19/20 spreads analyzed. A representative image of a single chromosome spread is shown.

Transcriptome Analysis Demonstrates cPP Cells Are Closely Related to Their In Vivo Counterparts

We next determined the transcriptome-wide gene counts by RNA-seq for cPP lines from three different genetic backgrounds and the PPd15 differentiation cultures from which they were established. Samples for RNA-seq were also taken from cPP cells at early, mid, and late passages. Gene
expression levels correlated strongly between different cPP samples, indicating that neither genetic background nor time in culture significantly affect the cPP transcriptome (Figure S3A). However, to completely eliminate donor-specific effects on gene expression, the following analysis used mean gene counts for cPP (early passage) and PPD15 cells derived from H9 and HES3 hESCs and AK6-13 hiPSCs.

To determine how similar cPP cells are to their in vitro and in vivo counterparts, we compared the cPP transcriptome with the published transcriptomes of pancreatic progenitors differentiated in vitro (Cebola PP) and from CS16-18 human embryos (CS16-18 PP), as previously described (Cebola et al., 2015), and a diverse collection of adult and embryonic tissues (Bernstein et al., 2010; Petryszak et al., 2013). Relative to non-pancreatic tissues, cPP cells exhibited similar patterns of gene expression to both PPD15 and Cebola PP cells (Figure 3A). Furthermore, cPP, PPD15, and Cebola PP cells closely resembled in vivo pancreatic progenitors at CS16-18, and all four cell populations expressed similar levels of genes associated with endodermal and pancreatic development (Figure 3B). However, as expected, cPP cells do not express the late-stage pancreatic progenitor markers NKX6-1, PTF1A, and CPA1. When taken together, these data demonstrate that the culture conditions described here maintain cPP cells in a developmental state closely related to both the embryonic human pancreas and pancreatic progenitors generated by directed differentiation.

To further characterize the transcriptional identity of cPP cells, we sought to identify genes that distinguish them from other lineages. Specifically expressed genes were defined as those that are variably expressed across the aforementioned panel of 25 tissues (coefficient of variance >1) and whose expression is upregulated in cPP cells (Z score >1), as previously described (Cebola et al., 2015). In total, 1,366 genes were identified, including numerous well-characterized markers of pancreatic progenitor cells, such as PDX1, SOX9, MNX1, and RFX6 (Figure 3C). To confirm the validity of this method, we demonstrated that these genes are not expressed by other endodermal derivatives, including liver, colon, and lung (Figure S3B). Encouragingly, around 80% of genes specifically expressed by cPP cells were shared with CS16-18 pancreatic progenitors and/or PPD15 cells. Furthermore, gene Z scores were highly correlated between these three pancreatic cell types but not with liver (Figure S3C), further demonstrating the transcriptional similarities between cPP cells and other pancreatic progenitors.

To determine the functional roles of cPP-specific genes, we analyzed associated Gene Ontology (GO) terms. The most enriched terms were those associated with endocrine pancreas development (Figure 3D, above). In order to determine how our culture conditions affect the behavior of cPP cells, we analyzed GO terms associated with genes expressed by cPP cells but not PPD15 or CS16-18 pancreatic progenitor cells (Figure 3D, below). Interestingly, the most enriched terms were those associated with aspects of cell division and telomere maintenance. Indeed, genes associated with these enriched terms, such as those encoding telomerase reverse transcriptase (TERT) and proliferating cell nuclear antigen (PCNA), were consistently upregulated in cPP cells from different genetic backgrounds, compared with the PPD15 populations from which they were derived (Figures 3E and 3F). We conclude that our feeder-based culture system maintains pancreatic progenitors as a stable population while upregulating genes required for long-term self-renewal.

A Feeder Layer of 3T3-J2 Cells Prevents cPP Differentiation while Exogenous Signals Promote Proliferation

We next investigated the roles played by the individual components of our culture system, specifically the layer of irradiated 3T3-J2 feeder cells, stimulation with EGF, FGF10, and retinoic acid (RA), and inhibition of the TGFβ and Notch signaling pathways. To assess the importance of the feeder layer, cPP cells were subcultured onto a layer of 3T3-J2 cells plated at decreasing densities and maintained in complete cPP culture media for 7 days. At reduced feeder densities, cPP cells continued to proliferate rapidly but quickly altered their morphology and could not be serially passaged (Figure 4A). The levels of PDX1 and SOX9 remained stable, indicating cPP cells are committed to the pancreatic lineage, while markers of duct (KRT19 and CA2) and acinar (CPA1 and AMY2B) differentiation were upregulated (Figure 4B). However, we did not observe upregulation of endocrine markers (NGN3 and NKX2-2), suggesting that 3T3-J2 feeder cells are required to block further differentiation toward the ductal and acinar lineages.

To establish the roles played by the growth factors and small molecules in our culture media, we removed each individually and assessed the effect on differentiation and proliferation. Exclusion of EGF or RA prevented cPP expansion, while removal of the TGF-β inhibitor SB431542 caused colonies to detach from the feeder layer (Figure 4C). Removal of either FGF10 or the γ-secretase inhibitor DAPT did not significantly affect colony size or morphology in the short term but, when removed from the culture media over multiple passages, led to a noticeable loss of viability. Interestingly, none of the growth factors or signaling inhibitors was individually required for maintenance of PDX1 or SOX9 expression (Figure 4D). Indeed, removal of RA actually increased PDX1 expression. These results suggest that the growth factors and inhibitors present in our culture media are primarily required to drive proliferation of cPP cells rather than maintain their developmental state.
To quantify the effect of exogenous signaling molecules on the maintenance and expansion of cPP cells, we used a microbioreactor array (MBA) screening platform to measure differentiation and proliferation (Titmarsh et al., 2012). Single cPP cells were seeded in Matrigel-coated culture chambers in the absence of feeders and exposed for 3 days to complete cPP culture media in which the levels of EGF, RA, and DAPT were varied (Figure S4). We then used an image-segmentation algorithm to identify individual nuclei and quantify immunofluorescence staining for PDX1 and SOX9, thereby enabling us to determine the percentage of double-positive cells following exposure to different growth factor regimes. Reducing the levels of any of the three factors led to a reduction in both the total number of cells and the number of PDX1+SOX9+ cells (Figure 4E). However, neither the mean levels of PDX1/ SOX9 nor the percentage of PDX1+SOX9+ cells were dependent on the levels of these factors, suggesting they act primarily as mitogens. Interestingly, we noticed an increase in the number and percentage of PDX1+SOX9+ cells, but no change in the overall proliferation rate, when cells were exposed to higher concentrations of autocrine signals, particularly when provided with maximal levels of EGF, RA, and DAPT (Figure S4D). Exposure to endogenous soluble signaling molecules is therefore required to maintain PDX1 and SOX9 independently of proliferation.

When taken together, these observations demonstrate that self-renewal of cPP cells is dependent on activation of the EGF, FGF10, and RA pathways and inhibition of Notch signaling. Indeed, cPP cells and their in vitro (PPd15) and in vivo (CS16-18 pancreatic progenitor) equivalents expressed high levels of multiple receptors of EGF, FGF, RA, and Notch signaling, as well as the TGFβ receptors ALK4 and ALK5 (encoded by ACVR1B and TGFBR1, respectively) that are inhibited by SB431542 (Figure 4F). Consistent with our observations, production of FGF10 and RA by the surrounding mesenchyme is essential for expansion of the murine pancreatic bud (Bhushan et al., 2001; Martin et al., 2005; Ye et al., 2005), while EGFR is expressed throughout the pancreas and regulates islet development (Miettinen et al., 2000). Intracellular Notch signaling promotes expansion of pancreatic progenitors and prevents their further differentiation into endocrine cells (Hald et al., 2003; Murtaugh et al., 2003). Therefore, our observation that the γ-secretase inhibitor DAPT promotes proliferation of cPP cells is somewhat surprising. However, FGF10 has been shown to promote Notch activity in the developing pancreatic epithelium (Hart et al., 2003), and cPP cells express intermediate levels of the Notch effector HES1 relative to the 23 tissues described in Figure 3A (data not shown). Therefore, the relatively low concentration of DAPT added to cPP cultures most likely serves to temper Notch activity, and exceptionally high levels of Notch activity might actually suppress proliferation.

**Differentiation of cPP Cells into Pancreatic Cell Types In Vitro and In Vivo**

The canonical property of pancreatic progenitors is their ability to differentiate into each of the three lineages that constitute the pancreas as well as their functional derivatives. Initially, we sought to determine whether cPP cells are capable of commitment to the endocrine, duct, and acinar lineages in vitro. Since robust protocols for the directed differentiation of pancreatic duct and acinar cells have yet to be developed, cPP cells were replated in the absence of feeders and exposed to a minimal signaling regime that promotes multilineage differentiation (Figure 5A). Over the course of 12 days, we observed

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**Figure 3. Transcriptome Analysis of cPP Cells by RNA-Seq**

(A) Hierarchical clustering of Euclidean distances between transcriptomes of diverse adult and embryonic tissues shows that in vitro and in vivo pancreatic progenitors exhibit similar patterns of gene expression. Log2-transformed gene count values were used to calculate Euclidian distances. For detailed information on the sources of data used here, see Table S1.

(B) Heatmaps showing log2-transformed gene expression levels of key endodermal and pancreatic markers by in vitro and in vivo pancreatic progenitors. Levels in brain are shown for comparison.

(C) Genes specifically expressed by cPP, PPd15, and CS16-18 pancreatic progenitors. The coefficient of variance (CV) for each protein-coding gene across the 25 tissues shown in (A) was plotted against the corresponding Z score (see Supplemental Experimental Procedures). Specifically expressed genes are located in the upper right-hand quadrant (CV >1 and Z score >1) and include genes with well-characterized roles in early pancreatic development (labeled). The color scale denotes the number of genes. The Venn diagram shows overlap between genes specifically expressed by cPP, PPd15, and CS16-18 pancreatic progenitors.

(D) Biological process Gene Ontology (GO) terms associated with all genes specifically expressed by cPP cells (above) or genes specifically expressed by cPP cells but not PPd15 or CS16-18 pancreatic progenitors (below). Only GO terms associated with >5 genes and/or an adjusted p value <0.01 are shown.

(E) Heatmap of expression levels of genes associated with the enriched GO terms mitotic recombination, DNA strand elongation, telomere maintenance, and DNA packaging. Levels are shown for individual cPP and PPd15 populations derived from three different genetic backgrounds (H9, Ak6-13, and HES3) relative to the maximum detected value across the 25 different tissues shown in (A).

(F) Expression of selected telomerase pathway genes as measured by qRT-PCR in cPP and PPd15 cells. Error bars represent the SE of three technical replicates.
Depletion of 3T3-J2 Feeders Causes Morphological Changes and Differentiation

A

H9

AK6-13

3T3-Feeder x 10^6 cells/cm²

B

PDX1

SOX9

NGN3

NKX2-2

KRT19

CA2

CPA1

AMY2B

3T3-Feeder x 10^6 cells/cm²

Removal of Media Components Impedes Colony Formation

C

Complete Media

- EGF

- RA

- SB431542

- DAPT

- FGF10

D

Normalized expression relative to complete media

RDX1

SOX9

E

Exogenous Signaling Molecules Promote cPP Proliferation

F

Signaling Pathway Component RNA-seq Expression

(legend on next page)
upregulation of endocrine (NKK6-1, INS, and GCG), acinar (CPA1, AMY2B, and TRYP3), and duct (SOX9, KRT19, and CA2) markers, demonstrating that cPP cells retain multi-lineage potency in vitro (Figure 5B).

Of particular interest is the ability to generate β-like cells capable of secreting insulin in response to elevated glucose levels. Several groups recently published protocols that describe the differentiation of particular hESC and hiPSC cell lines into β-like cells. Activation of NKK6-1 prior to expression of NGN3 is thought to be essential for the formation of mature, functional β cells (Nostro et al., 2015; Russ et al., 2015). Therefore, we selected the four most promising protocols and assessed their ability to induce NKK6-1 expression while maintaining low levels of NGN3 (Pagliuca et al., 2014; Rezania et al., 2014; Russ et al., 2015; Zhang et al., 2009). Specifically, cPP cells were cultured as monolayers or aggregates, then exposed to the section of each differentiation protocol shown to induce NKK6-1 expression (Figure S5A). The protocol described by Russ et al. (2015) produced the highest levels of NKK6-1 expression and minimal activation of NGN3, with monolayer and suspension cultures yielding a very similar response (Figure S5B). Since the original protocol demonstrated the generation of insulin-secreting β-like cells when cells were differentiated as aggregates, we chose to use the 3D suspension platform for subsequent experiments.

Using the Russ et al. (2015) protocol, we found that around 40% of cPP cells reactivate NKK6-1. However, doubling the length of each of the first two treatments enabled the generation of nearly 70% double-positive cells, similar to the number originally reported (Figures 5E, SSC, and SSD). Interestingly, these PD1X1+NKK6-1+ cells generated convoluted structures reminiscent of the branching morphogenesis of the embryonic pancreas (Figures 5D and 5F). Further differentiation induced expression of the endocrine markers NKK2-2 and NGN3, the latter in a smaller subset of cells, reflecting its transient expression during endocrine commitment (Schwitzgebel et al., 2000; Figure 5G). Finally, after 16 days, 20% of cells contained C-peptide, a proxy for insulin production, similar to the 25% reported by Russ et al. (2015). Crucially, C-peptide+ cells did not co-express the β cell hormone glucagon, suggesting that these cells are unlike the polyhormonal cells produced by earlier generations of protocols, which are unable to secrete insulin in response to elevated glucose levels. However, NGN3 levels remained high at the end of the protocol and INS mRNA levels were significantly lower than in isolated human islets, suggesting that further optimization of the protocol is required (Figure 5K).

The most stringent test of developmental potency is whether a progenitor can differentiate into a particular lineage in vivo. To assess the potency of cPP cells, we injected these cells under the renal capsules of immunodeficient mice and immunostained markers of the three major pancreatic lineages after >23 weeks. We were able to identify large areas of cells expressing the β-cell marker C-peptide as well as the duct marker keratin 19 (KRT19), but we were unable to find trypsin+ acinar cells or glucagon+ endocrine cells (Figure 5L). However, trypsin+ cells were also observed rarely by Rezania et al. (2014) following transplantation of pancreatic progenitors, possibly because acinar cells cannot survive in the absence of ducts to carry away the digestive enzymes they secrete. The absence of cells expressing glucagon was surprising, but likely reflects generation of C-peptide+ cells by default in the absence of inductive signals required to form glucagon+ α cells.

The C-peptide+ cells did not form classical islet-like structures, but instead formed a series of interconnected cystic structures, as others have observed previously (Rostovskaya et al., 2015). Furthermore, we did not observe expansion of the progenitor population once transplanted, suggesting cPP cells differentiate rapidly into less proliferative cells.
in vivo. Accordingly, none of the 12 mice we assessed exhibited teratoma formation, despite transplanting >3 million cells into each mouse. These observations demonstrate that cPP cells retain the ability to differentiate into endocrine and duct cells in vivo, although it remains to be seen whether they are capable of forming acinar cells. Furthermore, the absence of teratoma formation suggests cPP cells may represent a safer alternative for transplants than cells differentiated directly from pluripotent stem cells.

**DISCUSSION**

Pluripotent stem cells have been proposed as an unlimited source of β cells for modeling and treating diabetes. However, the routine generation of functional β cells from diverse patient-derived hiPSC remains a challenge, partly because of the variability inherent in long, multi-step-directed differentiation protocols. Here, we describe a platform for long-term culture of self-renewing pancreatic progenitor cells derived from human pluripotent stem cells. These cPP cells are capable of rapid and prolonged expansion, thereby offering a convenient alternative source of β cells. Furthermore, cPP cells can be stored and transported as frozen stocks, and to date we have cultured cPP cells for up to 25 passages with no loss of proliferation. We observed that cPP cells express markers of pancreatic endocrine, duct, and acinar cells when differentiated in vitro, thereby demonstrating their multipotency, and we were able to generate up to ~20% C-peptide+ cells using a modified version of the β cell differentiation protocol described by Russ et al. (2015). The definitive test of developmental potency is whether a cell can differentiate into a particular lineage in vivo, and cPP cells indeed generate significant numbers of keratin-19+ duct cells and C-peptide+ β-like cells when transplanted under the renal capsule of an immunodeficient mouse, although it is unclear whether they retain the ability to form acinar cells in vivo.

Cells differentiating in vitro typically do so in an unsynchronized manner, causing cultures to become progressively more heterochronic with time and reducing the efficiency with which cells can be directed toward particular lineages. Therefore, the ability to capture and synchronize differentiating progenitors is essential for developing robust protocols for generating functional β cells from diverse genetic backgrounds. Extensive molecular characterization revealed that cPP cultures generated from both hESC and hiPSC represent stable populations of cells that express early pancreatic transcription factors consistently over time. The cPP transcriptome is closely related to that of the progenitor cells of the CS16-18 pancreas. However, comparison with human embryos at different stages of development suggests that cPP cells most closely resemble cells of the pancreatic bud between CS12 and CS13, based on robust expression of PDX1, SOX9, FOXA2, and GATA4/6 and the absence of NKX6-1 and SOX17 (Jennings et al., 2013, 2015).

In recent years, several groups reported methods for culturing human endodermal derivatives. Two separate reports demonstrated that hESC-derived definitive endoderm can be serially passaged and expanded if cultured on a feeder layer in the presence of appropriate mitogenic signals (Cheng et al., 2012; Sneddon et al., 2012). Subsequently, another group showed that foregut progenitor cells can be cultured in feeder-free conditions (Hannan et al., 2013). However, slow growth and variable gene expression between different lines have limited their utility. More recently, it was shown that pancreatic progenitors derived from reprogrammed endodermal cells could...
be expanded and passaged (Zhu et al., 2015). However, these cultures are highly heterogeneous, and it is not clear whether the minimal combination of signaling molecules and inhibitors used is sufficient to culture cells from different genetic backgrounds. Therefore, the culture system described here is the first to enable long-term self-renewal of multipotent pancreatic progenitors derived from genetically diverse hESC and hiPSC.

Intriguingly, 3T3-J2 feeders have been used to culture diverse cell types, including epidermal keratinocytes (Rheinwald and Green, 1975a, 1975b), corneal epithelium (Osei-Bempong et al., 2009; Rama et al., 2010), and intestinal stem cells (Wang et al., 2015), among others. Nonetheless, the mechanism(s) by which 3T3-J2 feeders stabilize cultured progenitors is unknown. An obvious candidate is signaling through the WNT/LGR5 pathway, which is required for the maintenance of organoid cultures generated from a variety of endoderm-derived adult tissues, including the pancreas (Barker et al., 2007, 2010; Boj et al., 2015; Huch et al., 2013). However, cPP cells do not express LGR5 and are unaffected by inhibition of endogenous WNT secretion (data not shown). Therefore, identification of the factor(s) produced by 3T3-J2 cells could lead to the discovery of a common signaling axis required to support self-renewal of progenitors from a wide variety of tissue types. Finally, it will be interesting to investigate whether adapted versions of the culture system described here are capable of capturing progenitors from other endoderm-derived tissues, such as the liver, stomach, and lung, or progenitors resident in the adult pancreas.

EXPERIMENTAL PROCEDURES

Human Pluripotent Stem Cell Culture and Differentiation

Human pluripotent cell lines were obtained as described in the Supplemental Experimental Procedures. Pluripotent stem cells were maintained on tissue culture plastic coated with Matrigel in mTeSR1 medium as described previously (Ludwig et al., 2006), and differentiated into pancreatic progenitors using the STEMdiff Pancreatic Progenitor kit (STEMCELL Technologies, 05120) according to the manufacturer’s instructions with the following modifications: (1) cells were initially seeded into 12-well plates (Corning, 353043) at a density of 10⁶ cells/well, and (2) stage 1 was extended to 3 days by repeating the final day’s treatment. All tissue culture was carried out in 5% CO₂ at 37°C.

Passaging and Maintenance of cPP Cells

Gentle cell dissociation reagent (STEMCELL Technologies, 07174) was used to passage cPP cells as aggregates that were then seeded at a 1:6 split ratio onto a layer of 3T3-J2 feeders (0.5 × 10⁶ to 1 × 10⁶ cells/cm²) in medium composed of advanced DMEM/F12 (Thermo Fisher Scientific, 21634010), 2 mM L-glutamine (Thermo Fisher Scientific, 25030), 100 U/mL penicillin/streptomycin (Thermo Fisher Scientific, 15140122), 1× N2 supplement (Thermo Fisher Scientific, 17502-048), 1× B27 supplement (Thermo Fisher Scientific, 17504-044), 30 nM dexamethasone (STEMCELL Technologies, 72092), 50 ng/mL EGF (R&D Systems, 236-EG-200), 50 ng/mL FGF10 (Source Bioscience, ABC144), 3 μM RA (Sigma, R2625), 10 μM SB431542 (Calbiochem, 616464), and 1 μM DAPT (Sigma, D5942). If plating single cPP cells, complete medium was supplemented with 10 μM Y27632 for the first 48 hr (Sigma, Y0503). Medium was completely replenished every 2–3 days. See Supplemental Experimental Procedures for details of 3T3-J2 cell culture.

RNA-Seq Analysis of Gene Expression

RNA was isolated from samples harvested from cPP and PPD15 cultures using an RNeasy mini kit (Qiagen, cat. no. 74104). Feeder removal microbeads (Miltenyi Biotec, 130-095-531) were used to deplete cPP cells of 3T3 feeders prior to RNA extraction. All RNA samples had an RNA integrity number >9. RNA-seq libraries were generated using the NEBNext Ultra RNA Library Prep Kit (NEB, E7530L) and sequenced on an Illumina HiSeq 2500 system generating single-end reads of 100 bp. Table S1 contains metadata for these and public datasets used for the RNA-seq gene expression analysis. Full details of how RNA-seq reads were aligned and analyzed can be found in the Supplemental Experimental Procedures.

Multilineage Differentiation

Monolayer differentiation cultures were established as described in Supplemental Experimental Procedures. Basal differentiation medium consists of advanced DMEM/F12 (Thermo Fisher Scientific, 21634010), 2.5 g/30 mL BSA (Sigma, A9418), 2 mM L-glutamine (Thermo Fisher Scientific, 25030), 100 U/mL penicillin/streptomycin (Thermo Fisher Scientific, 15140122), and 1× B27 supplement (Thermo Fisher Scientific, 17504-044). Supplements were added as follows: days 1–3 (3 μM RA [Sigma, R2625], 1 μM DAPT [Sigma, D5942], 100 μM BNZ [Sigma, B4560]) and days 4, 7, and 10 (3 μM RA, 167 ng/mL KAAD-cyclopamine [Calbiochem, 239807]).

β Cell Differentiation

Differentiation sphere cultures were established as described in Supplemental Experimental Procedures. Basal differentiation medium consists of DMEM high glucose, 2 mM L-glutamine, and 100 U/mL penicillin/streptomycin. Supplements were added as follows: days 1–4 (1× B27 supplement, 50 ng/mL EGF, 1 μM RA [days 1–2 only], 50 ng/mL FGF7 [days 3–4 only]); days 5–10 (1× B27 supplement, 500 mM LDN-193189 [STEMCELL Technologies, 72142], 30 nM TPB [EMD Millipore, S65740], 1 μM RepSox [STEMCELL Technologies, 72392], 25 ng/mL FGF7); and days 11–17 (DMEM low glucose [Thermo Fisher Scientific, 12320-032], 2 mM L-glutamine, 1× MEM non-essential amino acids [Thermo Fisher Scientific, 11140-050]).

Transplantation Assays

cPP cells were grown to confluence to displace and eliminate feeder cells, then treated with gentle cell dissociation reagent to generate single cells. Approximately 3 × 10⁶ to 5 × 10⁶ cells were
resuspended in 50 μL of undiluted Matrigel and injected under the kidney capsule of 8- to 12-week-old immunocompromised (NOD/SCID) mice. After 23–27 weeks, transplanted mice were euthanized and their kidneys cryopreserved prior to sectioning and immunostaining. The study protocol was approved by the National University of Singapore Institutional Review Board (NUS IRB 12–181) and Biomedical Research Council IACUC committee (151040).

**ACCESSION NUMBERS**

Primary RNA-seq datasets generated here are available at ArrayExpress under accession number ArrayExpress: E-MTAB-5731.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures, five figures, and three tables and can be found with this article online at http://dx.doi.org/10.1016/j.stemcr.2017.05.019.

**ACKNOWLEDGMENTS**

We thank Dr. Aya Wada and Prof. Huck Hui Ng for advice and guidance on cell culture techniques. We would also like to thank Dr. Michael Riedel, Dr. Jenna Moccia, and Dr. Charis Segeritz-Walko of STEMCELL Technologies for providing differentiation reagents, and Dr. Kim Robinson, Prof. Birgit Lane, and Prof. Yann Barrandon for supplying 3T3-J2 feeder cells. This work was funded by an EDB Singapore Childhood Undiagnosed Diseases Program grant and an A*STAR Strategic Positioning Fund (SPF) Genetic Orphan Diseases Adopted: Fostering Innovation Therapy (GODAFIT) grant.

Received: March 14, 2017
Revised: May 15, 2017
Accepted: May 15, 2017
Published: June 6, 2017

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