Alleviation of hyperglycemia in diabetic rats by intraportal injection of insulin-producing cells generated from surgically resected human pancreatic tissue

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Abstract

Although islet transplantation holds promise for the treatment of diabetes, the scarcity of donor tissue remains a major drawback. The aim of this study is to generate insulin-producing cells from adult human pancreatic cells isolated from surgically resected pancreatic tissue. To isolate pancreatic endocrine precursor cells from 57 surgically resected pancreases, the cells were cultured and propagated in conditioned medium after which they were differentiated in Matrigel. The resultant cells were characterized using morphology, immunofluorescent studies, expression of differentiated pancreatic islet-specific genes using quantitative reverse transcription-PCR, and glucose-induced insulin secretion through analysis of C-peptide secretion. The relationships between propagation of insulin-producing cells and clinical variables of the donor were also analyzed. Finally, insulin-producing cell function was examined in streptozotocin-induced diabetic rats. Pancreatic endocrine precursor cells were successfully cultured; insulin-producing cells cultured from soft pancreas parenchyma had a significantly higher success rate. Morphological examination revealed islet-like cluster formation upon transfer to Matrigel. The presence of the neural stem cell marker nestin, duct cell marker cytokeratin 19, and endocrine cell markers C-peptide and pancreatic and duodenal homeobox 1, was also observed. In addition, glucose-stimulated C-peptide release was significantly increased in the insulin-producing cells. Furthermore, in diabetic rats, transplantation of insulin-producing cells reduced hyperglycemia. Isolated pancreatic endocrine precursor cells from surgically resected pancreatic tissue differentiated into insulin-producing cells and showed characteristics of functional endocrine cells. Thus, surgically resected pancreatic tissue may represent an alternative source of functional insulin-producing cells.

Introduction

Type 1 diabetes is a chronic disease which is characterized by absolute insulin deficiency (Nerup et al. 1970, Atkinson 2005). Since the discovery of insulin more than 80 years ago and with the advent of glucose monitoring, considerable advances in insulin therapy have occurred. However, insulin replacement therapy remains only part of the ultimate solution for the treatment of patients who have type 1 diabetes. Therefore, novel therapies that address the shortcomings of insulin replacement therapy, leading to the eventual cure of type 1 diabetes, should be pursued.

Diverse therapeutic strategies for type 1 diabetes, including pancreas transplantation (Larsen 2004, Ryan et al. 2006), islet transplantation (Shapiro et al. 2000), regeneration therapy (Yamaoka 2002), and cell-based gene therapy (Samson & Chan 2006), have been reported. Because islet cell transplantation represents a less-invasive treatment strategy, it may be superior to whole pancreas transplantation. However, the paucity of donor pancreases and problematic harvest of sufficient islet cells from cadaverous donors remain the drawbacks to this therapeutic strategy. In previous studies, multiple donors as well as transplantations were required to achieve insulin independence, with a 1-year insulin-independent rate of 44% and only a 10% insulin-independent rate after a 5-year follow-up (Shapiro et al. 2003, 2006). Therefore, identification of new cell sources is necessary to develop future replacement strategies.
Regeneration therapy represents an approach that could potentially cure type 1 diabetes when combined with cell-based gene therapeutic strategies and immune system modulation (Sytwu et al. 2003, Sung et al. 2004). In ex vivo regeneration therapy, the patients’ own bone marrow stem cells are transiently removed and differentiated into β-cells in vitro (Lechner & Habener 2003, Moriscot et al. 2005) whereas in vivo regeneration therapy uses the patients’ own cells to regenerate impaired tissues (Zulewski et al. 2001, Gao et al. 2003). In vivo regeneration therapy is more cost-effective, has fewer side effects, and is more ethically and clinically acceptable; therefore, it may offer the greatest potential therapeutic value to diabetic patients, if effective protocols could be developed.

The β-cell mass within the adult pancreas possesses the ability to undergo limited regeneration following injury. Identifying the progenitor cells involved in this process and understanding the mechanisms leading to their maturation will open new avenues for the treatment of type 1 diabetes (Maria-Engler et al. 2004, Lin et al. 2006). Despite steady advances in determining the molecular mechanisms regulating early pancreatic development and β-cell regeneration, development of protocols and identification of determining factors to consistently produce successful cultures of pancreatic stem cells or β-cell progenitors have yet to be described (Dodge et al. 2009).

Although generation of functional neoislets has been reported from various adult human pancreatic tissues, including pancreatic islets (Lechner et al. 2005), nonislet pancreatic cells, discarded after islet isolation (Todorov et al. 2006), and exocrine cells (Rapoport et al. 2009), it remains to be determined whether these sources provide the amount of neoislets required for in vivo regeneration therapy for diabetic patients.

To identify and characterize a new cellular source for in vivo regeneration therapy in patients with type 1 diabetes, surgically resected human pancreatic tissue was obtained, and endocrine precursor cells were isolated, cultured, and characterized. Furthermore, donor characteristics were compared with insulin-producing cell culture success. Finally, insulin-producing cell function was examined in streptozotocin (STZ)-induced diabetic rats. Identifying a novel source of functional islet cells and determining the precise culture conditions required may represent a novel treatment strategy for patients with type 1 diabetes.

### Materials and Methods

**Pancreatic endocrine precursor cell isolation and culture**

From June of 2004 to September of 2007, pancreatic tissue was obtained from patients undergoing pancreatic resection for pancreatic head cancer \((n=20, 35.1\%)\), Ampulla of Vater cancer \((n=15, 26.3\%)\), common bile duct cancer \((n=6, 10.5\%)\), duodenal cancer \((n=2, 3.5\%)\), chronic pancreatitis \((n=3, 5.3\%)\), and other lesions \((n=22, 35.1\%)\) (Table 1). Among the 57 donors, 41 males and 16 females were enrolled in this study; the average donor age was 64 years \((\pm 15.4\text{ years})\) (Table 1). After the resection, normal pancreatic tissue, which consists largely of the exocrine acini, interlobular ducts, and islets of Langerhans, was identified as tissue without cancer cell infiltration by a pathologist at the Taipei Veterans General Hospital.

The human pancreatic tissue was immediately immersed in solution D \((0.137 \text{ M NaCl, 5.38 mM KCl, 0.19 mM Na}_2\text{HPO}_4, 0.205 \text{ mM K}_2\text{HPO}_4, 5.49 \text{ mM glucose, 0.058 M sucrose, 1% penicillin/streptomycin, and 0.12% fungizone (Invitrogen)) to prevent degradation. The average size of

| Table 1 Donor clinical characteristics with respect to insulin-producing cell culture success \((n=57)\). Data are shown as mean ± s.d. for donor age, and \(n\ (%)\) for categorical data which includes age above or below 65 years old, sex, pancreas parenchyma consistency, and diagnosis of primary lesion |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| **Characteristics** | **Total \((n=57)\)** | **Success \((n=20)^a\)** | **Failure \((n=37)\)** | **P value** |
| Age (years) | | | | |
| ≤ 65 years | 64.0 ± 15.4 | 64.6 ± 14.6 | 63.7 ± 15.9 | 0.532 (C) |
| > 65 years | 54.4 ± 16.7 | 50.0 ± 14.9 | 54.1 ± 17.4 | 0.704 (C) |
| Sex, females (%) | 18.8 | 25.0 | 12.8 | 0.010* (F) |
| Parenchyma consistency, \(n\ (%)\) | | | | |
| Soft | 61.4 | 85.0 | 48.6 | 0.070 (F) |
| Hard | 38.6 | 15.0 | 51.4 | |
| Diagnosis of primary lesion, \(n\ (%)\) | | | | |
| Pancreatic head cancer | 20 (35.1) | 5 (25.0) | 15 (40.5) | |
| Ampulla of Vater cancer | 15 (26.3) | 6 (30.0) | 9 (24.3) | |
| Common bile duct cancer | 6 (10.5) | 3 (15.0) | 3 (8.1) | |
| Duodenal cancer | 2 (3.5) | 0 (0) | 2 (5.4) | |
| Chronic pancreatitis | 3 (5.3) | 1 (5.0) | 2 (5.4) | |
| Others | 11 (19.3) | 5 (25.0) | 6 (16.2) | |

\(^a^P<0.05. \text{ C, using the } \chi^2 \text{ test; F, using Fisher’s exact test.}\)

\(^a^\text{Success was determined by formation of islet-like cell cluster and expression of PDX1 and C-peptide.}\)
pancreatic tissue obtained was 0.5×1×2 cm, weighing 0.5–2 g. After the isolated tissue was minced, it was digested with 2 mg/ml Type V Collagenase (Sigma–Aldrich) for 30 min at 37 °C. The digested sample was then washed three times with cold DMEM/F12 (Invitrogen). Pancreatic duct cells, islets, and stem cells were isolated by centrifugation at 1200 g for 20 min at 4 °C in Histopaque (1.077 mg/ml; Sigma–Aldrich):DMEM/F12 gradients. The cells were aspirated from the Histopaque:DMEM interface and then washed with DMEM/F12. The isolated cells were then cultured with CMRL 1066 medium (5.5 mM glucose, Invitrogen) containing 10% fetal bovine serum, 1% penicillin/streptomycin, 100 ng/ml nerve growth factor (NGF, R&D Systems, Minneapolis, MN, USA), 10 mM nicotinamide (Sigma), and 25 ng/ml epidermal growth factor (EGF, Invitrogen). Thereafter, the cells were cultured for 7–10 days until reaching confluence. During the expansion phase, the culture medium was changed every 3 days. Cells intended for immunofluorescent labeling were cultured on glass coverslips.

This study followed the tenets and regulations described in the Declaration of Helsinki. This study was reviewed by the Institutional Review Committee of the Taipei Veterans General Hospital.

**Cell proliferation analysis**

Cell growth assays were performed using the CellTiter96 Aqueous One Solution Cell Proliferation Assay kit (Promega) following the manufacturer’s instructions. Isolated pancreatic endocrine precursor cells were plated at concentrations of 10^4 cells/well in 96-well plates. The growth rate was analyzed at days 3, 5, and 7 in the expansion phase. Absorbance was measured at a wavelength of 490 nm, and the absorbance values of growth factor and serum-treated cells are presented as a percentage of the absorbance of untreated cells at day 3.

**Pancreatic differentiation in Matrigel**

Cultured cells were trypsinized with 0.05% trypsin/EDTA (Invitrogen), washed with serum-free DMEM/F12 (17.5 mmol/l glucose), and seeded into 6-well culture dishes coated with Matrigel (BD Bioscience, Bedford, MA, USA) diluted 1:10 at a concentration of 1×10^6 cells/well for differentiation. The culture medium contained insulin, transferrin, sodium selenite + linoleic acid (ITS +1, Sigma), 2 g/l BSA, and 10 ng/ml basic fibroblastic growth factor (bFGF, Invitrogen). Cells aggregated from monolayers to cell clusters in Matrigel and the expression of pancreatic and duodenal homeobox 1 (PDX1) and C-peptide during the differentiation and maturation phases.

**Immunofluorescent staining and microscope analysis**

The cells were fixed in PBS containing 4% paraformaldehyde, 2 mM EGTA and 400 mM sucrose at room temperature for 15 min. Cells were then permeabilized using 0.1% Triton X-100 in PBS at room temperature for 1 h, and blocked with 5 mg/ml BSA and 10% goat serum in PBS for 1 h. Cells were then incubated in blocking solution with primary antibody overnight at 4 °C after which they were washed in PBS for five times for 10 min each. Primary antibodies and dilutions were as follows: nestin mouse monoclonal antibody, 1:200 (Chemicon International, Temecula, CA, USA); Pdx1 rabbit polyclonal antibody, 1:500 (Chemicon International); C-peptide rabbit polyclonal antibody, 1:100 (Linco Research, St Charles, MO, USA); cytokeratin 19 (CK19) mouse monoclonal antibody, 1:200 (Dako, Carpinteria, CA, USA); Ki-67 mouse monoclonal antibody, 1:200 (Dako); and human nuclei mouse monoclonal antibody, 1:400 (Chemicon International). For detection of primary antibodies, Alexa-Fluor-488-conjugated (green, Molecular Probes, Inc., Eugene, OR, USA) or Cy-3–conjugated (red, Jackson Immunoresearch Laboratories, West Grove, PA, USA) secondary antibodies were used according to the manufacturer’s instructions. Briefly, cells were incubated with secondary antibodies diluted 1:200 in blocking solution for 1 h at room temperature. Cell nuclei were visualized using TOTO-3 di 5X00 (Molecular Probes) in blocking solution for 10 min at room temperature. The samples were mounted using an antiphotobleaching medium containing 20 mM n-propyl-gallate (Sigma) in 80% glycerol/20% PBS and then observed under a microscope equipped with phase contrast and epifluorescence light paths (Leica DMR.E2, Heidelberg, Germany). For confocal imaging, a Zeiss confocal microscope LSM 510 (Zeiss, Göttingen, Germany) was used for data acquisition.

**Quantitative real-time PCR**

Total RNA was extracted from pancreatic tissue or cultured cells using Trizol reagent (Invitrogen) and was treated with DNase (Invitrogen). RNA (500 ng) was reverse transcribed to cDNA in a 25 μl solution containing 5 μl 5 x first strand buffer (Invitrogen), 10 mmol/l dithiothreitol (Invitrogen), 1 mmol/l dNTPs (New England Biolabs, Ipswich, MA, USA), 2 mmol/l dNTPs (New England Biolabs, Ipswich, MA, USA), 400 μg/ml oligo(dT) (Invitrogen), 200 μg/ml RNase inhibitor (Invitrogen), and 200 units/ml Moloney murine leukemia virus (M-MLV) reverse transcriptase (Invitrogen). Oligonucleotide primers were designed using Primer Express software (Applied Biosystems) and used according to the manufacturer’s instructions. Real-time PCR was performed using the Applied Biosystems 7500 Fast Real-Time PCR System. The reaction was performed using a 20-μl reaction mixture containing 10 μl of SYBR Green PCR Master Mix (Applied Biosystems) with 2 μl cDNA. PCR amplification was performed with 40 cycles of 95 °C for 15 s, 60 °C for 1 min, and 72 °C for 1 min. Melting curve analysis was performed with a melting temperature (Tm) of 95 °C for 15 s, 60 °C for 15 s, and 95 °C for 15 s. The specificity of the PCR products was confirmed by gel electrophoresis and melting curve analysis. Relative quantitation was performed using the 2^(-ΔΔCt) method (Livak and Schmittgen 2001).
50 ng random hexamers (Invitrogen), and 200 units SuperScript II Rnase H reverse transcriptase (Invitrogen). Reverse transcription reactions were incubated for 10 min at 25 °C, 60 min at 42 °C, and 10 min at 95 °C. PCR analyses were carried out in duplicate using 1/10th of the cDNA per reaction and 400 nmol/l forward and reverse primers with Smart Quant Green Master Mix (Protech Technology Enterprise, Taipei, Taiwan). The expression of human glyceraldehyde-3-phosphate dehydrogenase (GAPDH) was used as a housekeeping gene for normalization of gene expression levels. The following primer pair sequences (with accession numbers and product sizes) were used in this study: Nestin (NM_006617, 131 bp) F: 5'-CTGGCCGGCTACTGAAAGTT-3' and R: 5'-AGGCTAGGACATCTGGAG-3'; neurogenin 3 (NGN3, NM_020999, 293 bp) F: 5'-TGGCCCGGTTAGAGATGAC-3' and R: 5'-GCTGGCCGTGGTCTGCTTCTTCTTC-3'; Paired box gene 6 (PAX6, NM_001604, 303 bp) F: 5'-GAATACAGAGACAGGAGCA-3' and R: 5'-GTTGAGGTATCATATACTCCG-3'; NeuroD (NM_002500, 384 bp) F: 5'-AACACTCATCTGGGCTCTGTCG-3' and R: 5'-GGTCCATCAAAGG-3'; GLUT2 (NM_000340, 211 bp) F: 5'-TGCGCCGGTAGAAAGGATGAC-3' and R: 5'-ACTGCGGGCTACT-3'; and GAPDH (NM_000209, 139 bp) F: 5'-GGAGCCGGAGGAGACAG-3' and R: 5'-TCTGGTCAAGTTCAACATGACAG-3'; glucose transporter 2 (GLUT2, NM_000340, 211 bp) F: 5'-GGTTTGTAACTGGTTTCACAC-3' and R: 5'-GGCTAGTTCTATTGCCAAG-3'; and Insulin (NM_000207, 115 bp) F: 5'-ACCAGCATCTGCTCCCTCTA-3' and R: 5'-GTTCAAGGCTTTGTTCTTT-3'; and GAPDH (NM_000204, 372 bp) F: 5'-CACACATCTTTCCAGGAGCAG-3' and R: 5'-TCA-GGCCACAGTTCTCCCAGGA-3'. Real-time PCR was performed using the ABI 7500 System (Applied Biosystems, Foster City, CA, USA) with the following cycling program: 50 °C for 2 min, 95 °C for 10 min, and 35 cycles at 95 °C for 15 s and 60 °C for 1 min. Melting curves were obtained at 60 °C. The number of PCR cycles was titrated in order to remain in the linear range of amplification. The resultant amplification products (10 μl) were separated using 2% agarose gel electrophoresis and were visualized with ethidium bromide that validated the specificity of the real-time PCRs. The data were analyzed using the 2-ΔΔCt method (Livak & Schmittgen 2001). Gene expression in the insulin-producing cell cultures at various stages (expansion, differentiation, and maturation) was expressed relative to that observed in the insulin-producing cell of the pancreatic tissue stage.

### Table 2

Average yield of insulin-producing cells and clusters at each phase of pancreatic endocrine precursor cell culture. Data are shown as mean ± s.d.

<table>
<thead>
<tr>
<th>Cell number</th>
<th>Pancreatic tissue (×10^7/g)</th>
<th>Endocrine precursors (×10^6)</th>
<th>Expansion phase (×10^6)</th>
<th>Differentiation phase (×10^6)</th>
<th>Maturation phase (×10^6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Islet-like cell cluster (&gt; 50 m)</td>
<td>2.49 (0.27)</td>
<td>1.28 (0.21)</td>
<td>3.58 (0.25)</td>
<td>2.98 (0.23)</td>
<td>3.21 (0.20)</td>
</tr>
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ND, not determined.

**Insulin and DNA content**

The cells were washed twice with PBS, resuspended in 300 μl of distilled cold water, and homogenized by sonication on ice. An aliquot of the homogenates was analyzed fluorometrically for DNA content in duplicate, and another aliquot was extracted with acid ethanol overnight and measured for insulin content using an ELISA kit (Merckodia, Uppsala, Sweden).

**Glucose challenge test**

Islet-like cell clusters in the maturation phase were washed three times with assay buffer (111 mmol/l NaCl, 27 mmol/l NaHCO3, 5 mmol/l KCl, 1 mmol/l CaCl2, 1 mmol/l MgCl2, 0.3 mmol/l MgSO4, 1.18 mmol/l Na2HPO4, 0.29 mmol/l KH2PO4, 0.2% BSA, and 2.8 mmol/l glucose) after which they were incubated for 30 min in 5-6 mmol/l glucose. Fresh assay buffer (500 μl) containing either 25 mmol/l glucose or additional KCl (30 mmol/l) was added, and the cells were incubated for an additional 30 min at 37 °C. Following the incubation, 300 μl of the assay buffer was collected, and C-peptide concentrations were measured using an ELISA assay kit as follows.

**C-peptide secretion**

For analysis of C-peptide secretion, medium recovered from the glucose challenge test was analyzed using an ELISA kit (Merckodia) according to the manufacturer’s instructions. Briefly, using the supplied 96-well plate, 25 μl of sample was added to each well containing 50 μl assay buffer and incubated for 1 h at 18–25 °C on a shaker. After washing six times with wash buffer (0.5% Tween 20 in PBS), 100 μl enzyme conjugate was dispensed into each well, and then incubated at 18–25 °C on a shaker. After a final wash, 200 μl tetramethylbenzidine substrate was added to each well and incubated for 15 min after which 50 μl stop solution was added quickly to the wells. The optical density was determined using a 450 nm wavelength light in an Automated Immuno and Chemical Analyzer (ChemWell, Taipei, Taiwan).

**Intrahepatic islet-like cell injection in STZ-induced diabetic rats**

Twenty-five male Sprague–Dawley rats maintained as a closed colony (body weight 180–200 g) were used in this study.
The experiment was performed with the approval of the Laboratory Animal Center of the National Defense Medical Center in Taipei, Taiwan. Hyperglycemia was induced in 16 rats through i.p. injection of 60 mg/kg STZ. Blood glucose levels were determined using the Roche ACCU-CHEK glucose meter (Roche Diagnostics) and blood isolated from the tail vein. Stable hyperglycemia (defined as blood glucose levels ranging from 16.7 to 33.3 mmol/l) developed in 14 rats after 1 week. Nonfasted, recipient diabetic rats were anesthetized with pentobarbital (40 mg/kg, i.p.), and implantation of an indwelling vascular catheter was performed as previously described (Strubbe & Steffens 1977). Briefly, after midline laparotomy, a small s.c. pocket in the left groin was created for placement of the vascular access device (Port-A catheter), which was filled with heparinized saline (10 U/ml). The catheter, connected to the Port-A, was s.c. inserted into the peritoneal cavity through a small puncture in the anterior abdominal wall. The free end of the catheter was then placed into the gastroduodenal vein, at the junction with the portal vein. For islet infusion, the Port-A was percutaneously punctured with a 21 gauge needle. After injecting 1 ml heparinized saline, aliquots of $5 \times 10^6$ insulin-producing cells or an equal volume of physiological saline were transplanted into the liver according to the study protocol. Blood glucose levels were monitored under nonfasting condition every 2 days after transplantation for a total of 63 days. To measure the human C-peptide, blood samples were analyzed using an ultrasensitive human C-peptide ELISA kit (Mercodia).

**Immunohistochemistry**

The rats were killed 9 weeks after transplantation and perfused with 4% formaldehyde (Ferak, Berlin, Germany). The livers were dissected and cut into 0.5–1.0 cm$^3$ sections. The samples were dehydrated and embedded in OCT (Sakura Finetek USA, Inc., Torrance, CA, USA) in liquid nitrogen. The cryosections (5 µm) were washed twice with PBS and incubated overnight at 4°C with mouse anti-human nuclei (1:400; Chemicon) and rabbit anti-human C-peptide (1:100; Santa Cruz Biotechnology, Santa Cruz, CA, USA) antibodies. After three washes with PBS, slides were incubated for 1 h at room temperature with FITC-labeled goat anti-mouse IgG (1:200; Chemicon) and rhodamine-conjugated goat anti-rabbit IgG (1:500; Chemicon) antibodies. The sections were mounted with mounting medium (Vector Laboratories, Burlingame, CA, USA) and viewed with a fluorescence microscope using appropriate filters.

**Statistical analysis**

Statistical analyses were performed using SPSS 15.0 statistics software (SPSS, Inc., Chicago, IL, USA). Donor age and yield of insulin-producing cell data are shown as mean ± S.D. and n (%) for categorical variables. The Pearson $\chi^2$ test was performed to compare the success rate of insulin-producing cell culture relative to donor characteristics. In addition, Fisher’s exact test was performed if any one cell number was less than five. Furthermore, one-way ANOVA was used to compare the measurements for greater than three conditions.

**Figure 1** Morphological analysis of cultured human pancreatic insulin-producing cells. (A) Phase contrast microscopy of isolated pancreatic cells shows a round shape with a large nucleus and abundant cytoplasm and some spindle-shaped cells. (B) In the expansion phase, isolated pancreatic stem cells grew rapidly in conditioned culture medium. (C) Upon culture in serum-free medium with Matrigel (the differentiation phase), some cells formed network structures. (D) In the maturation phase, dithizone staining was used to detect zinc-containing islet cell clusters by light microscopy. Bar=60 µm (A and B). Bar=20 µm (C and D).

**Figure 2** Endocrine precursor cell proliferation assay. Proliferation is expressed relative to absorbance of cells not treated with serum or growth factor on day 3. No serum or growth factor, black bars; serum addition, white bars; serum and growth factor addition, gray bars. Data represent the means ± S.E.M. of six separate experiments with cells from different donors. *P<0.001 versus cells not treated with serum and growth factor. $^*P<0.001$ versus cells on day 3.
A repeated ANOVA was also performed to identify differences in glucose changes among the treatment groups. The Bonferroni adjustment was performed for pair-wise comparisons after ANOVA. For most analyses, a $P$ value $<0.05$ was considered statistically significant; an adjusted significance level of $0.0167$ ($0.05/3$) was used for the Bonferroni test.

Results

Effects of clinical variables on insulin-producing cell culture success

To determine donor characteristics that may influence culture success, success of insulin-producing cell generation for each characteristic was determined. As shown in Table 1, the overall culture success rate was 35% for this donor group. Neither donor age nor gender was significantly associated with culture success. Parenchyma consistency was also analyzed; normal pancreatic tissue has soft parenchyma, which by definition is without fibrotic change; hard pancreatic tissue contains fibrotic changes as determined by pathological examination (Haber et al. 1999). Significant differences were observed in culture success and parenchyma consistency; the success rate in insulin-producing cell culture was significantly higher in tissue with soft parenchyma as compared to hard (48.6% vs 13.6% respectively, $P=0.010$). Furthermore, the primary-type lesion was not significantly associated with insulin-producing cell generation. Thus, pancreatic parenchyma consistency was the only donor characteristic associated with culture success.

Generation of islet-like clusters from adult human pancreas

To increase the efficiency of insulin-producing cell generation, our culture procedure was developed by modifying the previous methods. The average yield of insulin-producing cell clusters isolated at each culture phase (pancreatic tissue, endocrine precursors, expansion phase, differentiation phase, and maturation phase) was determined using phase contrast micromicroscopy (Table 2). There was no difference in the amount of insulin-producing cells generated from patients undergoing pancreatic head cancer resection versus duodenal cancer resection (data not shown).

Figure 3 Immunofluorescent analysis of protein expression in insulin-producing cells cultured in the expansion and maturation phases. Confocal microscopy was used to analyze the expression of pancreatic markers. (A and B) Cells in expansion phase were labeled with anti-Nestin (green) and anti-PDX1 (red) primary antibodies. Cultured pancreatic cells were double labeled with (C) anti-Ki-67 antibody (red) and TOTO-3 nuclear stain (blue) or (D) anti-CK19 antibody (red) and TOTO-3 nuclear stain (blue). Cultured pancreatic cells in the expansion phase (E) and within islet-like cell cluster (F) in the maturation phase were labeled with anti-C-peptide antibody (green). Bar = 20 μm.
In addition to tracking cell and cluster numbers, cell morphology was analyzed throughout the culture period (Fig. 1). Upon culture initiation, the cells began to proliferate and form small colonies (Fig. 1A). After 10–14 days, the cells reached confluence (Fig. 1B). In the expansion period, a 280-fold increase in cell number was estimated during this period (Table 2). As determined by a proliferation assay, a significant increase in cell growth was observed at days 5 and 7 in the expansion phase (Fig. 2). During the differentiation phase, which extended from 5 to 7 days, both spindle-shaped cells and islet-like cluster formation were observed (Fig. 1C). Finally, in the maturation phase, which lasted 5 days, a red color was observed within the cell clusters upon staining with dithizone, indicating the presence of zinc-containing islet cells (Fig. 1D). The maturation phase increased both the number and size of cell clusters (Table 2).

Cell marker expression in cultured insulin-producing cells within the expansion and maturation phases

To characterize the cultured cells at both the expansion and maturation phases, immunofluorescent staining with confocal microscopy was employed. Most cells cultured in the expansion phase were immunopositive for both nestin (Fig. 3A) and PDX1 (Fig. 3B), which are the markers for pancreatic endocrine stem cells. Although cell numbers increased in the expansion phase (Table 2), few (<5%) were immunopositive for Ki-67 (Fig. 3C), a proliferation marker. Compared with the cells in the expansion phase, final harvested cellular insulin/DNA ratio increased by 40-fold in the differentiation and maturation phases (Fig. 4). In addition, the majority of cells (>80%) were immunopositive for CK19, suggesting that they were of pancreatic duct cell origin (Fig. 3D). Approximately 5–10% of the cells in the expansion phase were immunopositive for C-peptide (Fig. 3E); C-peptide expression was also observed in cell clusters within the maturation phase (Fig. 3F).

Islet-like cell clusters expressed genes characteristic of stem and mature pancreatic cells

The phenotype of cultured pancreatic cells was analyzed by quantitative real-time PCR (Fig. 5). The expression of nestin, a neural stem cell marker, was detected in both surgically resected pancreatic tissue (pancreatic cells) and cells in the expansion phase of culture. The expression of nestin decreased considerably in the differentiation phase and was barely detectable by the maturation phase. During differentiation and maturation, the expression of the transcription factor, NGN3, which is transiently expressed in endocrine-specific precursor cells, was observed; NGN3 expression was also detected within the pancreatic cells (Rajagopal et al. 2003). In the expansion phase, the level of insulin mRNA in the cultured cells had markedly decreased compared to the starting material; however, in the differentiation and maturation phases, upregulation of insulin gene expression (up to sixfold) was observed in the majority of cultures. In successful cultures that expressed high levels of insulin, the expression levels of NEUROD, PDX1, and GLUT2 were upregulated in the differentiation and maturation phases as compared to the expansion phase. Furthermore, PAX6 expression also decreased in the expansion phase as compared to the pancreatic cells and increased in the differentiation and maturation phases of culture. Thus, cells within the maturation phase expressed genes associated with mature pancreatic cells to a similar degree as surgically resected pancreatic tissue.

Secretion of C-peptide by insulin-producing cells upon glucose stimulation

To test whether cell clusters within the maturation phase generated in vitro showed functional characteristics of pancreatic islets, the release of C-peptide was examined (Fig. 6). Because the cells were cultured in media supplemented with insulin, C-peptide was chosen instead of insulin to measure islet function (Hansson et al. 2004). Insulin-producing cells were stimulated with 5.6 or 25 mM glucose or 25 mM glucose + KCl after which C-peptide secretion was measured. As compared to insulin-producing cells stimulated with 5.6 mM glucose, those receiving 25 mM glucose or 25 mM glucose + KCl secreted significantly higher levels of C-peptide (1.68 vs 101.71 and 86.54 pmol/DNA per 30 min respectively; *P<0.01). Thus, the insulin-producing cells displayed characteristics associated with pancreatic islet function.

Insulin-producing cells reduced hyperglycemia after transplantation into the diabetic rat model

To better characterize the functional properties of the insulin-producing cells, they were transplanted into the liver of STZ-induced diabetic rats, and secretion of insulin in
response to high glucose was examined. Hyperglycemia was observed in the untreated STZ-induced diabetic rats as compared to the control group ($P < 0.001$). Significant reductions (30–40%, $P < 0.001$) in blood glucose levels were observed in rats that received insulin-producing cell transplantation as compared to untreated STZ-induced rats (Fig. 7A). Specifically, after week 3, reduced glucose levels were detected in the transplant group as compared to the untreated STZ group. The average survival time was 63, 52.3, and 63 days in the control, STZ, and STZ insulin-producing cell groups respectively.

To demonstrate that the glucose responses shown in Fig. 7A were associated with insulin-producing cell transplantation, human C-peptide levels were measured in the sera of each group (Fig. 7B). The levels of human C-peptide were highest in the STZ+ insulin-producing cell group ($P < 0.0167$); human C-peptide levels were undetectable in the control and STZ groups. Furthermore, human C-peptide-positive cells were detected within the liver tissue of the STZ+ insulin-producing cell group 9 weeks following islet transplantation (Fig. 8A). Specifically, at least ten C-peptide-positive cell aggregates were detected in the liver of each animal transplanted with insulin-producing cells; human C-peptide-positive cells were absent in the control and untreated STZ rats. The presence of human nuclei confirmed the presence of human cells within the rat liver (Fig. 8B). These results indicate that the transplanted insulin-producing cells are not only functional but also capable of long-term survival upon transplantation.

**Discussion**

In this study, functional insulin-producing cells were generated from surgically resected adult human pancreatic tissue. To our knowledge, this is the first study reporting...
the generation of functional insulin-producing cells from this donor source. In addition, culture success was compared with donor characteristics. While not all cultures were successful, the insulin-producing cells expressed NGN3, insulin, PDX1, PAX6, and NEUROD, and GLUT2. In addition, glucose-stimulated C-peptide release was significantly increased in the insulin-producing cells. Furthermore, transplantation of insulin-producing cells reduced hyperglycemia in STZ-induced diabetic rats. Few studies have determined the factors associated with culture outcomes of insulin-producing cells isolated from different human subjects. In this study, insulin-producing cell culture success was not affected by donor age (<65 vs >65 years) or gender, which is inconsistent with the previous report (Ihm et al. 2006). Using human islets from cadaveric donors, Ihm et al. (2006) reported that islet preparations from younger donors (<40 years) may improve the success rate of single-donor transplantation. Differences in success rate were observed in insulin-producing cell cultures from pancreatic tissue with soft versus hard parenchyma. It is possible that progenitor cells were more difficult to obtain from pancreatic tissue with hard parenchyma due to the high fibrotic tissue content, which is consistent with Mitnala et al. (2010).

The generation of enough functional islets represents an important step in regeneration therapy for diabetic patients. Previous studies have generated functional islets from islet cells themselves (Gershengorn et al. 2004, Russ et al. 2008, 2009, Joglekar et al. 2009a,b) as well as exocrine and duct cells from adult human pancreas (Bonner-Weir et al. 2000, Todorov et al. 2006) from either cadavers or living organ donors. Indeed, bone marrow stem cells can also be subcultured in vitro to more passages, but reduced levels of functional cells were obtained (Wu et al. 2009).

Despite the enormous efforts directed at understanding the factors that control growth and development of β-cells, the existence of adult pancreatic stem or progenitor cells for β-cell replacement in patients with diabetes remains unknown (Levine & Mercola 2004, Hao et al. 2006). Because islets comprise <2% of cells within the adult human pancreas (Bouwens & Rooman 2005), whole pancreatic tissue instead of islets alone was used as the starting material. Surgically resected pancreatic tissue consists of endocrine cells as well as interstitial and adherent exocrine acinar and ductal cells, blood vessels, stroma, and potentially yet unknown stem cells. We reasoned if resected pancreatic tissue contained endocrine progenitors, the amount of insulin-producing cells available for transplantation may be increased. Approximately 3000 islet-like cell clusters were generated from 1 g of pancreatic tissue. Because only 0.5–2 g pancreatic tissue were used as the starting material in this study, the scale of insulin-producing cell cultures could be readily expanded to meet the needs of transplantation in a clinical setting.

The expression of NGN3 during the differentiation phase supports the occurrence of endocrine cell neogenesis, as NGN3 is essential for the development of the endocrine cell lineages in the pancreas during embryogenesis (Gradwohl et al. 2000). In this study, NGN3 was detected in freshly isolated adult human pancreatic tissue. Thus, the presence of NGN3 could reflect ongoing endocrine cell neogenesis in

Figure 6 Secretion of C-peptide by cultured human pancreas insulin-producing cells in response to glucose stimulation. Insulin-producing cells cultured in the maturation phase were stimulated as indicated, and C-peptide in the medium was analyzed using ELISA. Measurements were normalized for the DNA content of each sample. Three samples per condition were measured. Similar results were obtained in at least three independent experiments. *P<0.001 compared to low-glucose stimulation.

Figure 7 Cultured human pancreas insulin-producing cells reduced hyperglycemia after transplantation into streptozotocin-induced diabetic rats. (A) Cultured insulin-producing cells or PBS were transplanted into the portal vein of streptozotocin-induced diabetic rats (STZ+insulin-producing cell and STZ groups respectively). Control (closed circle), STZ (open circle), and STZ+insulin-producing cell (filled down triangle). (B) Human C-peptide levels in the sera of control, STZ, and STZ+insulin-producing cell groups. Day 0 indicates the day of cell transplantation. *P<0.01.
adult islets in vivo or induced neogenesis due to the stress of pancreatic tissue isolation. However, the expression of NGN3 itself does not always support neogenesis mature cells may be dedifferentiated into NGN3-expressing cells (Kodama et al. 2005, Baeyens et al. 2006). Furthermore, a recent study demonstrated the clonal isolation of nestin-positive, endocrine progenitor cells from the adult mouse pancreas (Seaberg et al. 2004). Nestin has been previously suggested as a marker of pancreatic stem/progenitor cells (Lumelsky et al. 2001); however, this issue remains controversial (Selander & Edlund 2002, Treutelaar et al. 2003, Kushner et al. 2010). Lineage tracing studies found no contribution of stem/progenitor cells to β-cell renewal in adult mice, questioning the role of such cells in vivo (Gershengorn et al. 2004, Russ et al. 2008, 2009, Joglekar et al. 2009a, b). Other reports have indicated that CK19-expressing cells, believed to be of ductal origin, are the in vitro and in vivo precursors of β-cells (Gao et al. 2003, Hao et al. 2006). Lineage tracing studies of pancreatic ductal cells revealed that they are the in vivo progenitor, producing new islets after injury (Bonner-Weir et al. 2008). However, Solar et al. (2009) reported no such contribution of pancreatic ductal epithelium in postnatal endocrine or acinar cell development. In this study, expansion cultures consisted of nestin- as well as CK19-expressing cells, but the lineage relationships between these cell types and endocrine cells remain to be determined.

The dedifferentiation of pancreatic β-cells during in vitro expansion represents a major problem in using cultured cells for transplantation (Beattie et al. 1999). Maintenance of islet architecture by expansion in a three-dimensional gel matrix may limit this problem (Beattie et al. 2002). In this study, formation of three-dimensional cell clusters along with growth arrest in serum-free medium may be a critical factor for successful endocrine differentiation. Dedifferentiation and redifferentiation of cells in vitro have also been observed in other systems, including neural (Kondo & Raff 2000) and skin progenitor cells (Li et al. 2004). Further studies are required to differentiate between β-cell dedifferentiation/redifferentiation and the presence of preexisting stem/progenitor cells in the adult pancreas. However, from a therapeutic point of view, any approach that generates more β-cells for transplantation is important.

This study has limitations that should be considered. The surgically resected donor pancreatic tissues were isolated from patients with various pancreatic diseases. However, normal tissue within the donor tissue was pathologically identified and used for subsequent insulin-producing cell culture generation. Functional islets were generated from 35% of the donor tissue. Thus, the identification of a new source of tissue from which to generate neoislets is beneficial for the overall field of transplantation therapy (Wang et al. 2010). In addition, the precise feature of the culture procedure associated with neoislet proliferation and differentiation was not assessed in this study. For example, use of mixed cell cultures with a three-dimensional structure may have increased neoislet proliferation and differentiation. In addition, supplementation of the culture medium with NGF, EGF, and nicotinamide was essential to induce the proliferation of endocrine precursors. Furthermore, use of bFGF, ITS + l serum-free medium, and high-glucose culture medium was also important to promote islet differentiation and increase insulin content. Further studies are necessary to determine the association of these factors with proliferation and differentiation of insulin-producing cells.

In conclusion, this study describes the isolation of pancreatic endocrine precursor cells from adult human pancreatic tissue and the redifferentiation of insulin-secreting β-like cells from in vitro-cultured insulin-producing cell precursors. Further studies to optimize the in vitro differentiation and augmentation of cell proliferation are necessary.

Declaration of interest

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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