Quality of Isolated Pig Islets Is Improved Using Perfluorohexyloctane for Pancreas Storage in a Split Lobe Model

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Pancreas transportation between donor center and islet production facility is frequently associated with prolonged ischemia impairing islet isolation and transplantation outcomes. It is foreseeable that shipment of pig pancreases from distant centralized breeding facilities to institutes that have a long-term experience in porcine islet isolation is essentially required in future clinical islet xenotransplantation. Previously, we demonstrated that perfluorohexyloctane (F6H8) is significantly more efficient to protect rat and human pancreata from ischemically induced damage compared to perfluorodecalin (PFD). To evaluate the effect of F6H8 on long-term stored pig pancreases in a prospective study, we utilized the split lobe model to minimize donor variability. Retrieved pancreases were dissected into the connecting and splenic lobe, intraductally flushed with UW solution and immersed alternately in either preoxygenated F6H8 or PFD for 8–10 h. Prior to pancreas digestion, the intrapancreatic pO2 and the ratio of ATP-to-inorganic phosphate was compared utilizing 31P-NMR spectroscopy. Isolated islets were cultured for 2–3 days at 37°C and subjected to quality assessment. Pancreatic lobes stored in preoxygenated F6H8 had a significantly higher intrapancreatic pO2 compared to pancreata in oxygen-precharged PFD (10.11 ± 3.87 vs. 1.64 ± 1.13 mmHg, \( p < 0.05 \)). This correlated with a higher ATP-to-inorganic phosphate ratio (0.30 ± 0.04 vs. 0.14 ± 0.01). No effect was observed concerning yield and purity of freshly isolated islets. Nevertheless, a significantly improved glucose-stimulated insulin response, increased viability and postculture survival (57.2 ± 5.7 vs. 39.3 ± 6.4%, \( p < 0.01 \)) was measured in islets isolated from F6H8-preserved pancreata. The present data suggest that F6H8 does not increase islet yield but improves quality of pig islets isolated after prolonged cold ischemia.

Key words: Cold storage; Islet isolation; Organ preservation; Oxygenation; Pig pancreas

INTRODUCTION

Oxygen-precharged perfluorocarbons have been established in islet transplantation to reduce tissue damage induced by prolonged cold ischemia time (CIT) prior to isolation and subsequent transplantation (15,22,35). One member of this chemical group, perfluorodecalin (PFD), has been extensively investigated in pancreases retrieved from porcine (3,5) and human donors (28,34). The promising results that were initially reported could not be confirmed in larger trials involving more than 350 human islet isolations (9,21). Mathematical modeling on oxygen penetration depth demonstrated that, even at the most favorable conditions for completely saturated PFD, the oxygen penetration depth within a porcine or human average-sized pancreas at 8°C could be 1 mm at best, leaving the majority of the tissue anoxic (1,25). Isolation experiments in porcine pancreases also raised questions about the efficiency of PFD (4,18).

Comparative studies in rat and human pancreases suggested that pancreas preservation can be significantly improved replacing PFD by perfluorohexyloctane (F6H8), a semifluorinated alkane that is characterized...
by an increased lipophilicity and lower density facilitating the penetration of oxygen into deeper tissue layers of stored organs (17). This modification significantly improved islet isolation outcome after prolonged CIT in terms of islet yield, viability, and functional integrity in comparison to PFD (6,8).

In the present prospective study, we assessed the efficiency of F6H8 for long-term storage of pig pancreases procured for subsequent islet isolation utilizing the split lobe model in order to minimize donor-related variability (38). It is foreseeable that shipment of pig pancreases from distant centralized biosecure breeding facilities is essentially required in future clinical islet xenotransplantation (14).

Subjecting pig pancreas lobes to an estimated clinically relevant CIT of 8–10 h, the present study demonstrates that pig pancreas oxygenation by means of preoxygenated F6H8 improves islet quality.

MATERIALS AND METHODS

All animal studies were approved by local ethics committees.

Donor Animals and Pancreas Procurement

Donor sows were housed for piglet production by the German Federal Research Institute for Animal Health (Neustadt, Germany) until loss of fertility and were slaughtered at the same institute. All retired breeders (≤24 months) originated from the German Federal Hybrid Breeding Programme and were killed by exsanguination after carbon dioxide anesthesia. Ex situ dissection started after a primary warm ischemia time of 19±2 min. Retrieved pancreas were trimmed and dissected into the splenic and connecting lobe and cannulated according to the recommendations of Ferrer et al. (11). These lobes were particularly used because of the difficulty to preserve the integrity of the duodenal lobe without substantially extending the warm ischemia time. After a secondary warm ischemia time of 21±1 min, the pancreatic segments were intraductally flushed with cold University of Wisconsin solution (UWS, ViaSpan®, DuPont Pharmaceuticals Ltd., Letchworth Garden City, Hertfordshire, UK) at a ratio of 1 ml/g trimmed tissue. Total warm ischemia time accumulated to 40±1 min.

Pancreas Cold Storage

UWS-flushed pancreases were oxygenated utilizing the one-layer method as previously described (3). Briefly, the splenic and connecting lobes were immersed in 600 ml of either PFD or F6H8 (Novalig GmbH, Heidelberg, Germany) in an alternate manner to consider the asymmetric morphology of the porcine pancreas (23,36). Both oxygen carriers had been precharged for 15 min with 100% oxygen at a flow rate of 2,000 ml/min. Subsequently, the segments were shipped on ice to the isolation facility.

During preservation, the ratio of adenosine-triphosphate (ATP) to inorganic phosphate was assessed in the stored tissue by a noninvasive method utilizing 31P-nuclear magnetic resonance spectroscopy (NMR) in a small series of experiments (n=3), to measure 31P, a naturally abundant isotope, as previously described (30). Briefly, immediately after retrieval, pancreas lobes were intraductally injected with 1.0 ml/g histidine-tryptophan-ketoglutarate (HTK, Essential Pharmaceuticals, Newtown, PA, USA) solution instead of UWS to minimize background noise induced by phosphate dissolved in UWS. Afterwards, pancreatic segments were alternately incubated in either oxygen-precharged PFD or F6H8 and placed in a surface coil positioned in a 1.5-T magnet with a warm bore diameter of 680 mm and a gradient of 550 mm (Magnex Scientific, Abingdon, UK). The areas of the α-, β-, and γ-ATP peaks were visualized by means of a Tecmag Apollo spectrometer (Tecmag Inc., Houston, TX, USA) and were compared with the area of the inorganic phosphate peak to monitor the bioenergetic status of the organs.

Immediately after arrival in the laboratory, the intrapancreatic partial oxygen pressure (pO₂) was measured in both segments at a depth of 5–7 mm utilizing an optic fiber sensor housed in a cannula (Precision Sensing GmbH, Regensburg, Germany) that was introduced in the tissue as previously described (6). This procedure resulted in leakage during collagenase infusion and so the number of experiments was limited to six. Total CIT was defined as the interval between intraductal infusion of UWS and injection of the collagenase blend into the pancreatic duct.

Islet Isolation

After pO₂ measurement pancreas lobes were intraductally distended with collagenase NB-1 supplemented with neutral protease NB (Serva Electrophoresis, Uetersen, Germany). To provide similar conditions for each of the lobes, collagenase and neutral protease were adjusted to a concentration of 15 PZ-U and 1.3 DMC-U/g trimmed tissue weight and dissolved in 1 ml/g cold UWS, respectively. Isolation and purification was performed as previously described with minimal modifications regarding the digestion temperature kept in a range from 28°C to 32°C during recirculation and dilution phase (7). Purified islet fractions were washed twice in Hank’s balanced salt solution supplemented with 10% newborn calf serum and collected finally in 100 ml of bicarbonate-free CMRL 1066 supplemented with 25 mM HEPES, 1 mM pyruvate, 10 mM nicotinamide (all PAA, Pasching, Austria), 20% porcine serum, 2.5 mM l-glutamine, 100 U/µg/ml penicillin–streptomycin (all Gibco, Life Technologies, Paisley, Scotland, UK), and
20 µg/ml ciprofloxacin (Bayer, Leverkusen, Germany). Islet culture was performed in gas-permeable culture bags (Baxter Healthcare, Newbury, UK) incubated in humidified normal air for 2–3 days at 37°C.

**Islet Characterization**

Subsequent to purification and after culture islet yield and purity were determined in a standardized procedure converting islet yield to islet equivalents (IEQ) with an average diameter of 150 µm (26). Islet ATP content was measured in aliquots of freshly isolated islets utilizing the ATP Lite assay kit (Perkin Elmer, Seer Green, UK). ATP was normalized to islet DNA content determined by means of a fluorometric assay (Picogreen; Life Technologies, Paisley, Scotland, UK). Determination of fluorometric membrane integrity was performed utilizing 25 µmol/L of Syto-13 and 50 µmol/L of Sytox Orange (Molecular Probes, Life Technologies, Paisley, Scotland, UK) for staining of viable and dead cells, respectively (6). Islet in vitro function was assessed during static glucose (Sigma-Aldrich, AB., Stockholm, Sweden) incubation of islets precultured for 2–3 days at 37°C. Twenty hand-selected islets with an average diameter of 150–200 µm were sequentially incubated in duplicate first for 45 min in bicarbonate-free CMRL 1066 supplemented with 2 mmol/L glucose followed by 45 min of incubation at 20 mmol/L glucose followed finally by a second 45-min period at 2 mmol/L glucose. The glucose stimulation index was calculated by dividing the insulin release at 20 mmol/L glucose by the mean of the basal periods. After incubation islets were recovered and sonified in acid ethanol for subsequent determination of intracellular insulin content (6). Intracellular and released insulin was measured utilizing an enzyme immunoassay specific for pig insulin (Mercodia, Uppsala, Sweden) and normalized to islet DNA content.

Islet in vivo function was assessed in NMRI nude mice (Scanbur, Sollentuna, Sweden) rendered diabetic by a single intravenous injection of 90 µg/g alloxan (Sigma-Aldrich, AB., Stockholm, Sweden) 3 days prior to transplantation of an allotment of 1,750 IEQ beneath the kidney capsule. Blood samples were taken from the recipients’ tail vein and analyzed utilizing a one-touch glucoseometer (Bayer, Leverkusen, Germany). Pretransplant, the nonfasting serum glucose levels did not vary significantly after utilization of F6H8 or PFD (Table 1). No significant differences were observed between F6H8 and PFD with respect to total islet yield (260,300 ± 30,180 vs. 232,500 ± 42,930 IEQ) and corresponding insulin content (911 ± 212 vs. 953 ± 204 µg/lobe).

**Table 1. Islet Isolation Variables**

<table>
<thead>
<tr>
<th>Oxygen Carrier</th>
<th>n</th>
<th>Digestion Time (min)</th>
<th>Undigested Tissue (%)</th>
<th>Packed Tissue (µL/g)</th>
<th>Yield</th>
<th>Insulin Content</th>
</tr>
</thead>
<tbody>
<tr>
<td>F6H8</td>
<td>10</td>
<td>24.5 ± 1.2</td>
<td>12.7 ± 2.2</td>
<td>0.35 ± 0.04</td>
<td>260,300 ± 30,180</td>
<td>911 ± 212</td>
</tr>
<tr>
<td>PFD</td>
<td>10</td>
<td>24.3 ± 1.4</td>
<td>10.9 ± 1.8</td>
<td>0.33 ± 0.04</td>
<td>232,500 ± 42,930</td>
<td>953 ± 204</td>
</tr>
</tbody>
</table>

F6H8, perfluorohexyloctane; PFD, perfluorodecalin; IEQ, islet equivalent.

**RESULTS**

**Pancreas Storage Variables**

Measurement of intrapancreatic pO2 (n=6) revealed that storage of pancreatic segments in preoxygenated F6H8 resulted in a significantly higher pO2 (10.11 ± 3.87 mmHg) compared to lobes preserved in oxygen-charged PFD (1.64 ± 1.13, n=6, p<0.05). This observation corresponded with a higher ATP-to-inorganic phosphate ratio in F6H8-stored lobes (0.30 ± 0.04 vs. 0.14 ± 0.01) as measured by 31P-NMR during 8 h of cold storage in a small series of experiments (n=3). In agreement, the ATP content in freshly isolated pig islets recovered from pancreatic lobes that were preserved in preoxygenated F6H8 was significantly increased compared to lobes stored in precharged PFD [290 ± 60 vs. 203 ± 31 ATP (pg/ng DNA), p<0.01]. No significant difference was found between PFD (413 ± 23 mmHg) and F6H8 (392 ± 43 mmHg, NS) with respect to media pO2 after having been used for storage of porcine pancreatic segments for 9.8 ± 1.5 and 9.6 ± 1.5 h of cold ischemia, respectively.

**Islet Isolation Outcome**

Isolation variables such as digestion time, percentage of undigested tissue, and packed volume of digested tissue did not vary significantly after utilization of F6H8 or PFD (Table 1). No significant differences were observed between F6H8 and PFD with respect to total islet yield (260,300 ± 30,180 vs. 232,500 ± 42,930 IEQ) and corresponding insulin content (911 ± 212 vs. 953 ± 204 µg/lobe)
Likewise, islet yield per gram trimmed pancreatic tissue (3,020 ± 330 vs. 2,940 ± 270 IEQ/g) and final islet purity (85.0 ± 4.3 vs. 83.3 ± 5.3%) (Table 1) did not differ as well. However, islet quality assessment revealed that significantly more pig islets survived during culture at 37°C when isolated from pancreases stored in preoxygenated F6H8 (57.2 ± 5.7% vs. 39.3 ± 6.4%, p < 0.01) as shown in Table 2. Higher survival of cultured islets isolated from F6H8-preserved organs was associated with increased islet viability (81.8 ± 2.6% vs. 75.9 ± 2.6%, p < 0.05). Pancreas oxygenation influenced also in vitro function determined as glucose stimulation index during static incubation (2.28 ± 0.35 vs. 1.90 ± 0.42, p < 0.05). Moreover, islets isolated from PFD-preserved pancreases were not capable of downregulating insulin release after switching from 20 to 2 mmol/L glucose concentration as demonstrated in Figure 1. Intracellular insulin content was not altered by the preservation method used during cold storage (Table 2).

Posttransplant function of pig islets cultured for 3 days was assessed after transplantation beneath the kidney capsule of diabetic nude mice. A gradual decrease in hyperglycemia was observed until alloxan-injected recipients became normoglycemic through transplanted pig islets (Fig. 2). Altogether, 9 of 10 (90.0%) and 7 of 8 (87.5%) diabetic recipients were cured by islets from F6H8- or PFD-oxygenated pig pancreases (NS by Fisher’s exact test), respectively.

**DISCUSSION**

Several studies previously demonstrated that storage of ischemic pig pancreases in oxygenated PFD improves islet isolation outcome in comparison to cold storage in UWS (5,12). The present study was initiated to investigate whether PFD can be successfully replaced in a large animal model utilizing an oxygen carrier with different chemical characteristics. In order to provide similar conditions for both experimental groups, the pancreas split lobe model was utilized (38). Because of the asymmetric morphology of the porcine pancreas, it was not possible to provide perfectly identical conditions for the paired experiments particularly concerning the potential native islet yield within the connecting and splenic lobe. With respect to this parameter, previous histological observations are not consistent and seem to depend on the pig breed used (23,36). Regardless of these concerns, we did not expect any bias for the present study, since the different lobes were symmetrically allocated (five splenic and five connecting lobes) for both PFD- and F6H8-preserved organs and were digested with the same concentration of enzymes adjusted according to trimmed tissue weight.

The present report shows that pig islet quality can be even more enhanced after 8–10 h of cold storage when PFD is replaced by the new oxygen carrier F6H8. These findings confirm previous experiments in rat and human pancreases, suggesting that F6H8 improves pancreas preservation during 24-h storage prior to islet isolation (6,8). The specific structure of F6H8, belonging to the group of semifluorinated alkanes, results in a lower density and 20-fold higher lipophilicity compared to PFD (17,24). These characteristics seem to favor the penetration of the oxygen carrier into deeper tissue layers resulting in a still limited but nevertheless enhanced oxygenation of the stored tissue as demonstrated by intrapancreatic pO2 determination and

<table>
<thead>
<tr>
<th>Oxygen Carrier</th>
<th>n</th>
<th>Postculture Survival (%)</th>
<th>Stimulation Index</th>
<th>Insulin Content (ng/ng DNA)</th>
<th>Viability (%)</th>
<th>ATP (pg/ng DNA)</th>
</tr>
</thead>
<tbody>
<tr>
<td>F6H8</td>
<td>9</td>
<td>57.4 ± 5.7</td>
<td>2.28 ± 0.35</td>
<td>6.0 ± 2.2</td>
<td>81.8 ± 2.6</td>
<td>290 ± 60</td>
</tr>
<tr>
<td>PFD</td>
<td>9</td>
<td>39.3 ± 6.4</td>
<td>1.90 ± 0.42</td>
<td>5.5 ± 2.1</td>
<td>75.9 ± 2.6</td>
<td>203 ± 31</td>
</tr>
<tr>
<td>p value</td>
<td></td>
<td>&lt;0.01</td>
<td>&lt;0.05</td>
<td>&gt;0.05</td>
<td>&lt;0.05</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

F6H8, perfluorohexyloctane; PFD, perfluorodecalin.

**Figure 1.** Glucose-stimulated insulin release. Isolated pig islets precultured for 2 days at 37°C were sequentially incubated for 45 min in 2 (white bars), 20 (gray bars), and again 2 mmol/L of glucose (n=9). Insulin release was normalized to islet DNA content. Wilcoxon test revealed *p < 0.05 and ***p < 0.001 as indicated. F6H8, perfluorohexyloctane; PFD, perfluorodecalin.
well during culture (33). It was shown that culture at 37°C removes all predamaged islets thereby leveling potential differences with regard to morphological integrity and improving functional capacity (10,27). A critical point of the present study is therefore that the diabetic nude mice were not treated with a marginal islet mass.

In summary, the present study demonstrates that the quality of pig islets isolated after long-term storage is significantly improved when pancreases are oxygenated utilizing oxygen-precharged F6H8 in comparison to PFD. This new oxygen carrier has lipophilic characteristics and improves the oxygenation of deeper pancreatic tissue layers, thereby increasing the ATP synthesis in islets, which seems to be important for enhanced islet survival after culture. As previously shown, techniques are existing that have a high efficiency to optimize intrapancreatic oxygenation (31). Nevertheless, these advanced techniques are still at the investigational stage, whereas our approach could be immediately translated to the clinic and is essentially as simple as the static cold storage methods currently used.

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