Does Islet Size Really Influence Graft Function After Clinical Islet Transplantation?

Stephen J. Hughes, PhD,1 Paul A. Bateman, PhD,1 Sarah E. Cross, PhD,1 Daniel Brandhorst, PhD,1 Heide Brandhorst, PhD,1 Ioannis Spiliotis, MD,2 Chitrabhanu Ballav, MD,2 Miranda Rosenthal, PhD,3 Martin K. Rutter, MD,4,5 James Shaw, PhD,6 Stephen Gough, PhD,2 and Paul R.V. Johnson, MD1

Background. It has been proposed that islet transplants comprised primarily of small rather than large islets may provide better graft function, due to their lower susceptibility to hypoxic damage. Our aim was to determine whether islet size correlated with in vivo graft function in islet transplant recipients with C-peptide-negative type 1 diabetes when islets have undergone pretransplant islet culture. Methods. Human pancreatic islets were isolated, cultured for 24 hours and infused by standardized protocols. Ninety-minute stimulated C-peptide concentrations were determined during a standard meal tolerance test 3 months posttransplant. The islet isolation index (IEq/islet number) was determined immediately after isolation and again before transplantation (after tissue culture). This was correlated with patient insulin requirement or stimulated C-peptide. Results. Changes in insulin requirement did not significantly correlate with islet isolation index. Stimulated C-peptide correlated weakly with IEq at isolation (P = 0.40) and significantly with IEq at transplantation (P = 0.018). Stimulated C-peptide correlated with islet number at isolation (P = 0.013) and more strongly with the islet number at transplantation (P = 0.001). In contrast, the correlation of stimulated C-peptide and islet isolation index was weaker (P = 0.018), and this was poorer at transplantation (P = 0.034). Using linear regression, the strongest association with graft function was islet number (r = 0.722, P = 0.001). Islet size was not related to graft function after adjusting for islet volume or number. Conclusions. These data show no clear correlation between islet isolation index and graft function; both small and large islets are suitable for transplantation, provided the islets have survived a short culture period postisolation.

(Transplantation 2018;102: 1857–1863)
best in leading clinical centers worldwide, only 50% of islet preparations are of sufficient quality and high enough yield for transplantation.\textsuperscript{2,3,8} In addition, early posttransplant islet destruction, variable engraftment, the nature of the immunosuppression used, and subsequent graft deterioration over time all contribute to variable long-term islet function.\textsuperscript{9-12} Indeed, it is widely accepted that less than 50% of transplanted islets have superior function compared with larger ones.\textsuperscript{20-24} Consequently, there has been considerable debate about the optimal composition of islet preparations used for transplantation. It has long been established that total islet graft volume and islet number are critical for transplantation success,\textsuperscript{7-19} but more recently, it has also been suggested that grafts comprising predominantly of small islets may be preferential due to their reduced susceptibility to hypoxia and central necrosis. Indeed, a number of in vitro studies have shown that smaller islets have superior function compared with larger ones.\textsuperscript{20-24} However, data confirming this in clinical studies are more limited. Lehmann et al\textsuperscript{15} acknowledged the importance of the factors discussed above for islet success and transplantation outcomes but also reported that better function was associated with grafts comprising of smaller islets in simultaneous islet-kidney transplant recipients; however, there were only 7 recipients in the study, all in end-stage renal failure. Importantly, the islets used within that study were transplanted “fresh” after islet isolation, rather than having the benefit of undergoing a period of islet culture. A larger study in autologous islet recipients after pancreatectomy also supported this conclusion, although in this study, grafts of marginal islet mass were often transplanted.\textsuperscript{25} Our own observation,\textsuperscript{26} however, is that provided larger islets survive a period of pretransplant islet culture, they can confer advantages over smaller islets in terms of graft function and graft longevity. To investigate this discrepancy, we aimed to correlate islet size (preculture and postculture) with in vivo measures of graft function in all islet transplant recipients receiving their first transplant in our islet transplant program over a 6-year period.

**METHODS**

Human pancreases were retrieved with appropriate consent and ethical approval from 25 deceased multiorgan donors (15 women and 10 men). Median (range) donor age was 49 years (34-60 years) and median donor BMI of 29 kg/m\textsuperscript{2} (23-37 kg/m\textsuperscript{2}). After standardized procurement, the pancreas was transported in University of Wisconsin solution at 4°C to the Diabetes Research & Wellness Foundation Human Islet Isolation Facility in Oxford. Median (range) cold ischemia time was 6.7 hours (4-10.5 hours). Pancreatic islets were isolated using a standard protocol as described previously.\textsuperscript{27} After infusion and digestion with collagenase NB 1 and neutral protease NB enzyme blend (Serva, Heidelberg, Germany), islets were purified using a Ficoll-based continuous density gradient and quality-assessed (for sterility, viability, purity, and yield) as previously described.\textsuperscript{28-30} Islet number and size were determined in dithizone stained islet samples by visual microscopic inspection by comparing the stained islet-particles in multiple representative samples against a size graticule ranging from 50 to greater than 400 μM in size. Each preparation was resuspended in 150- to 200-mL CMRL-based culture media and multiple samples collected whilst in continuous suspension. The islets in the samples were visualized by dithizone staining under a light microscope and the size of each islet in the sample determined by comparison with a calibrated graticule in the microscope objective; the total number of islets in each sample and their sizes were recorded. Islet preparations were then cultured for a minimum of 24 hours in the CMRL medium in a humidified atmosphere at 37°C before a full reassessment. Islets were then transplanted either locally or at a satellite transplant center. If allocated to a recipient at a satellite transplant center,\textsuperscript{31} the islet preparation was transported by road (maximum journey time, 8 hours) in cooled standard 500 mL blood transfusion bags with a temperature monitor.\textsuperscript{27,28} After confirmation of satisfactory temperature maintenance during transport together with confirmation of maintained islet integrity and viability from a side-arm islet sample, islets were transplanted directly from the transport bag.

Only recipients undergoing their first islet transplant were included in this study (n = 25). In all cases, islets were transplanted by percutaneous transhepatic delivery into the portal vein under radiological visualization. All patients received a minimum of 5000 Islet Equivalents (IEq) per kg body weight (median, 5500 IEq/kg). Peritransplant, recipients were placed on an intravenous glucose/insulin sliding scale and received a heparin infusion according to published protocols.\textsuperscript{32}

Each patient received a standard immunosuppression protocol comprising alemuzumab (Campath) induction 30 mg before and on day 1 after transplant; tacrolimus at 0.05 mg/kg per day titrated to a serum trough level of 8 to 12 ng/mL; mycophenolate mofetil (500 mg BD) and total daily insulin requirements were monitored at routine intervals. Intensified insulin regimens were continued in all postdischarge, with the goal being maintenance of optimal glycemic control.

In a subgroup of the recipients (n = 18), 90-minute stimulated C-peptide concentrations were determined during a standard meal tolerance test 3 months posttransplant. Grafts with primary nonfunction as defined by a stimulated C-peptide level less than 50 pmol/L were excluded from all subsequent analysis. Assessment of stimulated C-peptide was made at 3 months posttransplantation as an indicator of early graft function. It was also done at this time to avoid any complicating effects of a subsequent second transplant which is carried out in the UK islet transplant program.\textsuperscript{31} The beta score was also determined for each transplant recipient at 3 months postransplant as described by Ryan et al.\textsuperscript{33}

For each islet preparation transplanted, the islet isolation index (IEq/islet number) was calculated; the total IEq of the preparation is the internationally agreed standard for the total volume of the islet graft. The islet isolation index therefore is a measure of the average size of each islet within the graft, with an index of 1 indicating an average islet size of 150 μM in diameter. Islet isolation index was related to 3 variables of graft function in transplant recipients: (a) the change in insulin requirement (before vs 3 months after transplant), (b) the 90-minute stimulated C-peptide level taken 3 months postransplant, and (c) the beta score at 3 months postransplant.\textsuperscript{35}

For statistical analysis, data were tested for normality and are presented as means ± standard deviations (SD) or median (range) depending on data distribution. Groups were compared using paired Student t test when appropriate. Correlations were assessed using Pearson correlation coefficient.
Linear regression was performed with preculture and postculture islet isolation variables (age, BMI, CIT, viability, purity, IEQ/kg, Islet number/kg, and islet isolation index) as exposure variables and change in insulin dose or stimulated C-peptide as outcome variables. Multiple regression was conducted to assess the impact of islet isolation outcomes on stimulated C-peptide, adjusting for donor age and BMI, or on beta score, adjusting for donor age and prep purity. Final multiple regression models were fitted using backward stepwise elimination of candidate variables. \( P \) less than 0.05 was considered statistically significant. Finally, a power calculation was carried out assuming \( P \) less than 0.05, a sample size of 25, and \( \alpha = 0.05 \). To obtain an \( R \) value of 0.5, the power was 0.73; \( R \) value of 0.6, power was 0.9; and an \( R \) value of 0.7, the power was 0.98. Data analysis was performed using SPSS, Version 20 & 24 (2011) (IBM Corp, Armonk, NY).

**RESULTS**

Donor and islet preparation characteristics are summarized in Table 1. Immediately postisolation, the islet yield (mean ± SD) was 454800 ± 190900 IEq with a mean islet number of 232320 ± 114060 islets, purity ranged from 50% to 90% and viability was 75% or greater in all cases. After 24 hours in culture, the islet number had significantly reduced to 197300 ± 91200 islets \((P = 0.013, \text{ paired } t\text{ test})\) whereas the small reduction in IEq to 408600 ± 126700 was not significant. Before transplantation, patients required a mean of 31.5 ± 13.4 U insulin per day \((0.48 ± 0.18 \text{ U/kg body wt per day})\), which was reduced by 17.1 ± 9.9 U/day to 15.3 ± 14.0 U/day \((0.22 ± 0.18 \text{ U/kg body wt per day})\) at 3 months posttransplant. The mean stimulated C-peptide concentration during a meal tolerance test at 3 months posttransplant was 624 ± 524 pmol/L (Table 1).

The change in insulin requirement before and 3 months after transplant was correlated with islet size as assessed by the islet isolation index of the transplanted islet preparation (Figure 1). There was no significant correlation with changing islet isolation index of the transplanted islets and graft function; either absolute change in insulin requirement \((r = -0.005, P = 0.49)\), percentage change in insulin requirement \((r = -0.27, P = 0.099)\), or insulin change per kg body weight of the recipient \((r = -0.15, P = 0.24)\). The change in insulin requirement also correlated poorly with IEQ \((r = 0.11, P = 0.49)\) or IEQ/kg body weight \((r = 0.05, P = 0.4)\) (data not shown).

In contrast, stimulated C-peptide levels at 90 minutes during a meal tolerance test were significantly correlated with the number of islets in the graft \((P = 0.004, r = 0.605)\), and the strongest correlation determined was with the number if islets in the graft per kg body weight \((r = 0.722, P < 0.001, \text{ Figure 2B})\). In contrast, the correlations with IEQ \((r = 0.17, P = 0.25)\) or IEQ per kg recipient body weight \((r = 0.494, P = 0.018)\) were poorer (Figure 2A). When plotted against the

---

**TABLE 1.**

| Donor characteristics (N = 25) |  |
|---|---|
| Age, y | 49 (34-60) |
| BMI, kg/m² | 29 (23-37) |
| CIT, h | 6.5 (4-10.5) |

| Islet preparation characteristics (N = 25) |  |
|---|---|
| Isolation yield, IEq | 454800 ± 190900 |
| Isolation number (islet particles) | 232320 ± 114060 |
| Isolation purity | 74 ± 12 |
| Isolation Viability | 84.6 ± 8.8 |
| Transplant yield, IEq | 408600 ± 126700 |
| Transplant number (islet particles) | 197300 ± 91200 |
| Transplant purity | 74 ± 12 |
| Transplant viability | 85.6 ± 8.4 |

| Recipient characteristics (N = 25) |  |
|---|---|
| Weight, kg | 66 ± 11.2 |
| IEQ transplanted/kg body weight | 6340 ± 2323 |
| Insulin requirement before TX, U/d | 31.5 ± 13.4 |
| Insulin requirement after TX, U/d | 15.3 ± 14.0 |
| Stimulated C-peptide, pmol/L | 624 ± 524 |
islet isolation index of the transplanted islets, the correlation with stimulated C-peptide remained significant ($r = -0.416$, $P = 0.043$) as did stimulated C-peptide per IEq per kg body weight of the recipient ($r = -0.439$, $P = 0.034$, Figures 2C and D). As the islet yield and number had changed during culture from the values measured at isolation, we also determined correlations for stimulated C-peptide with parameters measured immediately after isolation. The correlation of stimulated C-peptide with islet number was significant (Figure 3B, $r = 0.526$, $P = 0.013$) but poorer than that determined at transplantation, whereas the correlation with IEq at isolation was weak (Figure 3A, $r = -0.056$, $P = 0.411$). The correlation of stimulated C-peptide per IEq per kg body weight, however, was more significant with the islet isolation index of the preparations determined immediately after isolation (Figure 3C, $r = -0.495$, $P = 0.018$) than that determined at transplantation.

We assessed the strength of univariable- and multivariable-adjusted associations between islet isolation yield parameters, purity, viability, donor variables (age, BMI, CIT), and 90-minute stimulated C-peptide levels using linear regression. These data confirmed the observation that the factor most strongly associated with graft function in univariable analysis was the number of islets in the graft ($P = 0.001$; Table 2). In a model including both total islet volume (IEq/kg) and islet size (Isolation index; IEq/Islet number), total islet volume and donor age were related to 90-minute stimulated C-peptide levels (islet volume, $\beta = 0.101$, $P = 0.019$; donor

**FIGURE 2.** Correlation of stimulated C-peptide with the number IEQ (A), islets (islet particles) (B), or islet isolation index (C) or stimulate C-peptide/kg wt (D) at transplantation.

**FIGURE 3.** Correlation of stimulated C-peptide with the number of IEQ (A), number of islets (islet particles) (B) or islet isolation index (C) at isolation.
In a model including both islet number (number/kg) and islet size, only the islet number was related to the 90-minute stimulated C-peptide levels ($\beta=0.19$, $P=0.001$; Table 2). Islet size was not related to graft function after adjusting for total islet volume or islet number or any of the other variables analyzed.

Islet isolation index, islet volume, and islet number at transplantation were also correlated with the beta score as another measure of graft function at 3 months posttransplant (data not shown). The correlation of the beta score with islet isolation index at transplantation ($r = -0.584$, $P=0.011$) was poorer than that with islet number ($r = 0.784$, $P=0.001$). Univariate linear regression with the following factors: islet isolation variables and donor variables with beta score showed that islet number/kg ($P=0.026$), islet isolation index ($P=0.022$), and islet purity significantly ($P=0.027$) affected this measure of graft function. Further multiple regression analysis, however, indicated that both islet number/kg ($P=0.032$) and islet purity ($P=0.38$) but not islet isolation index had a significant effect in the final model of the analysis.

**DISCUSSION**

In this study, we tested the existing hypothesis that islet grafts comprising predominantly smaller islets have superior function compared with those composed of large islets. We have done this by correlating islet size within transplanted grafts with the posttransplant metabolic outcomes of the recipients. Several studies using experimental animal models have previously shown that smaller islets are potentially more beneficial as islet grafts. Smaller islets have been shown to have improved nutrient supply, with larger islets depleted of both oxygen and glucose at the core. Small islets also exhibit improved insulin secretory function in vitro, exhibit a higher vascular density, and function preferentially in transplant models. Studies with human islets in vitro also reproduce some of these results; small human islets were...
shown to be less susceptible to hypoxia and had improved secretory function compared with large islets.\textsuperscript{15,36}

However, studies investigating whether the hypothesis is born out in allotransplant recipients of islet grafts have been limited. Lehmann et al\textsuperscript{15} correlated islet size with stimulated C-peptide in 7 simultaneous islet-kidney transplant recipients with type 1 diabetes. This study showed that increasing the number of islets in the graft significantly correlated with graft function and critically, the most significant correlation was found between islet isolation index and stimulated C-peptide per kg body weight of the recipient. The authors argued that correction for islet size proved to be the best predictor of graft function with 89\% of the variability (as indicated by the correlation $R^2$) being accounted for by the islet isolation index parameter. However, the authors\textsuperscript{15} recognized that their findings should be confirmed in a larger series of islet transplants. The present study was undertaken to do this. Here, we report on 2 parameters of islet function in a larger cohort of islet transplant recipients: reduction in insulin requirement per day and stimulated C-peptide levels during a meal tolerance test. Neither of the parameters analyzed show convincing correlation with the islet isolation index. There was no significant correlation with changes in patient insulin requirement in absolute terms or per kg body weight or when expressed as % change. Nor did the present study reproduce the findings of Lehmann et al,\textsuperscript{15} as the most significant correlation we found was between the number of islets in the graft and graft function as measured by stimulated C-peptide. When the correction for islet size was included, the correlation was poorer and $R^2$ reduced with only 19.3\% of the variability accounted for by this parameter.

Two factors may contribute to the different findings in our study compared with those of Lehmann et al.\textsuperscript{15} First, their study was in patients undergoing simultaneous islet kidney transplants, whereas all our patients received islet transplants alone. Second, their observations were made using islets that had not undergone the benefit of a period of pretransplant islet culture.\textsuperscript{37} In the UK islet transplant program, all islet preparations considered for transplantation undergo a minimum of 24-hour culture before transplant.\textsuperscript{37} Using this protocol, any marginal grafts susceptible to hypoxic damage and leading to declining viability are screened out and therefore do not proceed to transplantation. When islets are transplanted immediately after isolation or where only the immediate postisolation characteristics (of yield and viability) are used to decide suitability for transplantation, declining or marginal grafts are potentially transplanted. Because large islets are particularly susceptible to central necrosis, it can be postulated that when islets are transplanted immediately after isolation, it is the large islets that are particularly vulnerable to destruction, whereas large islets that are transplanted after surviving a period of islet culture are primarily robust and fully viable. Furthermore, the culture process itself may promote an adaptive or remodeling process in the islet preparation in which susceptible islets are selectively lost. Changes in islet morphology (rounding up) during culture and the significant reduction in islet number after culture in the present study is evidence of just such a process. We have also recently shown that short-term tissue culture ameliorates the destruction of human islets by instant blood-mediated inflammatory reaction.\textsuperscript{38} Thus it is possible that islets cultured for short periods before transplant are effectively preconditioned and any correlation between islet size and function in grafts is lost or reduced. In the present study, the correlation of islet isolation index and graft function was stronger in freshly isolated islets compared with islets cultured for a minimum of 24 hours at transplant.

In addition, it may be the case that there are too many other confounding donor and recipient factors which override any effect of islet size in determining graft function.\textsuperscript{25} Suszynski et al\textsuperscript{25} argued that islet autotransplant patients offered a good model for studying the effect of the size of islets in the graft as these patients are uncompromised by factors, such as autoimmunity, prolonged diabetic environment,\textsuperscript{39} and possibly IBMIR.\textsuperscript{16,40} In a cohort of 58 patients receiving islet autotransplants after pancreatectomy for chronic pancreatitis, the islet isolation index correlated with the change in insulin requirement and insulin independence rates in the patients.\textsuperscript{25}

Although the number of observations we present here is significantly (threefold) greater than those reported by Lehmann et al,\textsuperscript{15} our analysis indicates that this retrospective study is still underpowered. Unfortunately, the number of patients available to study has been limited by the practice in the UK islet transplant program to retransplant islet recipients within 6 months of receipt of the first graft.\textsuperscript{31} The effect of a second islet graft to potentially change the mean islet size transplanted in the recipient makes the analysis carried out in the present study impossible.

Another theoretical limitation of the present study is that the method used to determine islet size in the graft samples may be less accurate than computational methods which may capture more of the population of smaller islets during analysis. In the present study, islet sizes were determined by inspection and comparison using a graticule after microscopic visualization. Our methodology has been agreed and validated between the participating islet isolation laboratories operating within the UK islet transplant program. However, there is no evidence of a significant difference in the size distribution of islets in our study compared with the study by Lehmann et al because the range of islet isolation indices here (0.8-4.2) overlap almost completely with those which they reported (0.75-3.3).\textsuperscript{15}

Finally, it is important to note that changes in insulin requirement as a measure of graft function are less robust than stimulated C-peptide measurements. The latter is a positive output directly measuring graft function, whereas changes in insulin requirement are indirect and may not be necessarily linearly responsive to graft size. This less objective parameter, however, has been used as the indicator of islet graft function in previous studies on islet transplant recipients.\textsuperscript{25,41}

In summary, our study found no clear correlation between islet isolation index and islet graft function in recipients receiving islets that had been cultured before implantation. These data therefore do not support the hypothesis that smaller islets have better function when used in clinical islet transplantation; large islets are equally suitable provided they have undergone and survived a short period of culture postisolation.

REFERENCES

1. Lakey JR, Warnock GL, Rajotte RV, et al. Variables in organ donors that affect the recovery of human islets of Langerhans. Transplantation. 1996;61:1047–1053.

2. O’Gorman D, Kin T, Murdoch T, et al. The standardization of pancreatic donors for islet isolation. Transplantation. 2005;80:801–806.
3. Ponte GM, Pfeffer A, Messinger S, et al. Toward maximizing the success rates of human islet isolation: influence of donor and isolation factors. Cell Transplant. 2007;16:595–607.

4. Hanley SC, Paraskevas S, Rosenberg L. Donor and isolation variables predicting human islet isolation success. Transplantation. 2008;85:950–955.

5. Niclauss N, Bosco D, Morel P, et al. Influence of donor age on islet isolation and transplantation outcome. Transplantation. 2011;91:360–366.

6. Wang LJ, Kim T, O’Gorman D, et al. A multicenter study: North American islet donor score in donor pancreas selection for human islet isolation for transplantation. Cell Transplant. 2018;25:1515–1523.

7. Ihm SH, Matsumoto I, Sawaeda T, et al. Effect of donor age on function of isolated human islets. Diabetes. 2006;55:1361–1368.

8. Johnson PR, Jones KE. Pancreatic islet transplantation. Semin Pediatr Surg. 2012;21:272–280.

9. Friberg AS, Lundgren T, Malm H, et al. Transplanted functional islet mass: donor, islet preparation, and recipient factors influence early graft function in islet-after-kidney patients. Transplantation. 2012;93:632–638.

10. Gerber PA, Pavlivcik V, Demartines N, et al. Simultaneous islet-kidney vs pancreas-kidney transplantation in type 1 diabetes mellitus: a 5 year single centre follow-up. Diabetologia. 2009;51:110–119.

11. Ryan EA, Paty BW, Senior PA, et al. Five-year follow-up after clinical islet transplantation. Diabetes. 2005;54:2060–2069.

12. Smith RM, Gale EA. Survival of the fittest? Natural selection in islet transplantation. Transplantation. 2005;79:1301–1303.

13. Davalli AM, Ogawa Y, Ricordi C, et al. A selective decrease in the beta cell mass of human islets transplanted into diabetic nude mice. Transplantation. 1995;59:817–820.

14. Eriksson O, Eich T, Sundin A, et al. Islet size index as a predictor of outcomes in clinical islet autotransplantation. Transplantation. 2001;50:710–719.

15. Lehmann R, Dressler RA, Kugelmeier P, et al. Superiority of small islets in human islet transplantation. Diabetes. 2007;56:94–963.

16. Lehmann R, Zuellig RA, Kugelmeier P, et al. Superiority of small islets in human islet transplantation. Diabetes. 2007;56:394–403.

17. Nilsen B, Ekdahl KN, Korsgren O. Control of instant blood-mediated inflammatory reaction to improve islets of Langerhans engraftment. Curr Opin Organ Transplant. 2011;16:620–626.

18. Shapiro AM, Lackey JR, Ryan EA, et al. Islet transplantation in seven patients with type 1 diabetes mellitus using a glucocorticoid-free immunosuppressive regimen. N Engl J Med. 2000;343:230–238.

19. Ryan EA, Lackey JR, Rajotte RV, et al. Clinical outcomes and insulin secretion after islet transplantation with the Edmonton protocol. Diabetes. 2001;50:710–719.

20. Benhamou PY, Oberholzer J, Toso C, et al. Human islet transplantation network for the treatment of type 1 diabetes: first data from the Swiss-French GRAGIL consortium (1999–2000). Groupe de Recherche Rhin Rhône Alpes Genève pour la transplantation d’îlots de Langerhans. Diabetologia. 2001;44:859–864.

21. Kauri LM, Dickson RC, Constantinius I, et al. Effects of oxygen on metabolic and secretory activities of beta TC3 cells. Biochim Biophys Acta. 1996;1291:163–166.

22. Fujita Y, Taikita M, Shimoda M, et al. Large human islets secrete less insulin per islet equivalent than smaller islets in vitro. Islets. 2011;3:1–5.

23. Zorzzi D, Phan T, Sequi M, et al. Impact of islet size on pancreatic islet transplantation and potential interventions to improve outcome. Cell Transplant. 2015;24:11–23.

24. Nam KH, Yong W, Harvat T, et al. Size-based separation and collection of mouse pancreatic islets for functional analysis. Biomed Microdevices. 2010;12:985–987.

25. Suszynski TM, Wilhelm JJ, Radosевич DM, et al. Islet size index as a predictor of outcomes in clinical islet autotransplantation. Transplantation. 2014;97:1286–1291.

26. Hughes SJ. Islet size does not influence graft function in islet transplant recipients. 5th EPITA Winter Symposium; 2015: OP17.

27. Brooks AM, Walker N, Aldibbit A, et al. Attainment of metabolic goals in the integrated UK islet transplant program with locally isolated and transported preparations. Am J Transplant. 2013;13:3236–3243.

28. Aldibbit A, Huang GC, Zhao M, et al. Validation of islet transport from a geographically distant isolation center enabling equitable access and National Health Service Funding of a clinical islet transplant program for England. Cell Med. 2011;2:97–104.

29. Gray DW, Morris PJ. Developments in isolated pancreatic islet transplantation. Transplantation. 1987;43:321–331.

30. Wang LJ, Kin T, O’Gorman D, et al. A multicenter study: North American islet donor score in donor pancreas selection for human islet isolation for transplantation. Cell Transplant. 2001;10:2512–2518.

31. Hudson A, Bradbury L, Johnson R, et al. The UK pancreas allocation scheme for whole organ and islet transplantation. Am J Transplant. 2015;15:2443–2455.

32. Kohn A, Senior P, Salam A, et al. Insulin-heparin infusions peritransplant substantially improve single-donor clinical islet transplant success. Transplantation. 2010;89:465–471.

33. Ryan EA, Paty BW, Senior PA, et al. Beta-score: an assessment of beta-cell function after islet transplantation. Diabetes Care. 2005;28:343–347.

34. Kauri LM, Jung SK, Kennedy RT. Direct measurement of glucose gradients and mass transport within islets of Langerhans. Biochim Biophys Res Commun. 2003;304:371–377.

35. Papas KK, Long RC Jr, Constantinius I, et al. Effects of oxygen on metabolic and secretory activities of beta TC3 cells. Biochim Biophys Acta. 1996;1291:163–166.

36. Fujita Y, Taikita M, Shimoda M, et al. Large human islets secrete less insulin per islet equivalent than smaller islets in vitro. Islets. 2011;3:1–5.

37. Lehmann R, Weber M, Barthold P, et al. Successful simultaneous islet-kidney transplantation using a steroid-free immunosuppression: two-year follow-up. Am J Transplant. 2004;4:1117–1123.

38. Ramnath RD, Maillard E, Jones K, et al. In vitro assessment of human islet vulnerability to instant blood-mediated inflammatory reaction (IBMIR) and its use to demonstrate a beneficial effect of tissue culture. Cell Transplant. 2015;24:2505–2512.

39. Sutherland DE, Gruessner AC, Carlson AM, et al. Islet autotransplantation outcomes after total pancreatectomy: a contrast to islet allograft outcomes. Transplantation. 2008;86:1790–1802.

40. Bennet W, Sundberg B, Groth CG, et al. Incompatibility between human blood and isolated islets of Langerhans: a finding with implications for clinical intraportal islet transplantation?. Diabetes. 1999;48:1907–1914.

41. Andres A, Kin T, O’Gorman D, et al. Clinical islet isolation and transplantation outcomes with deceased cardiac death donors are similar to neurological determination of death donors. Transplant Int. 2016;29:34–40.