Bone marrow as an alternative site for islet transplantation

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The liver is the current site for pancreatic islet transplantation, but has many drawbacks due to immunologic and nonimmunologic factors. We asked whether pancreatic islets could be engrafted in the bone marrow (BM), an easily accessible and widely distributed transplant site that may lack the limitations seen in the liver. Syngeneic islets engrafted efficiently in the BM of C57BL/6 mice rendered diabetic by streptozocin treatment. For more than 1 year after transplantation, these animals showed parameters of glucose metabolism that were similar to those of nondiabetic mice. Islets in BM had a higher probability to reach euglycemia than islets in liver (2.4-fold increase, P = .02), showed a compact morphology with a preserved ratio between α and β cells, and affected bone structure only very marginally. Islets in BM did not compromise hematopoietic activity, even when it was strongly induced in response to a BM aplasia-inducing infection with lymphocytic choriomeningitis virus. In conclusion, BM is an attractive and safe alternative site for pancreatic islet transplantation. The results of our study open a research line with potentially significant clinical impact, not only for the treatment of diabetes, but also for other diseases amenable to treatment with cellular transplantation. (Blood. 2009;114:4566-4574)

Introduction

The liver was suggested as an optimal site for islet transplantation by Lacy and colleagues, using a rat model of diabetes.1 By the 1980s, successful transplantation of islet autografts was reported in humans by using infusion of cells into the patient’s liver through the portal venous circulation.2–4 Subsequently, the publication of the first case of insulin independence in a diabetic patient after infusion of islets through the portal vein consecrated the liver as the site of choice for the islet transplantation in humans.5 The subsequent clinical experience of islet transplantation has been developed almost exclusively using the intrahepatic infusion through the portal vein.6

In the last years, however, it has become increasingly recognized that the liver may not be the optimal environment as a recipient site for pancreatic islets, owing not only to immunologic7,8 but also anatomic9,10 and physiologic factors that most likely contribute to the decline of islet mass after implantation.11–13 Intrahepatic islet infusion in humans is associated with an immediate blood-mediated inflammatory reaction, thrombosis, and hepatic tissue ischemia with elevated blood liver enzymes.12,14–19 Loss of as many as 50% to 75% of islets during engraftment in the liver has been suggested to be a prime factor necessitating the very large number of islets needed to achieve normoglycemia. Furthermore, the necessity for cannulation of the portal system to seed the islets produces an increase in portal pressure proportional to the islet mass administered by infusion,20 thus restricting the total mass that can be implanted. As a consequence, a highly purified suspension of islets is needed to transplant sufficient cells to achieve insulin independence. Because the purity of the suspension is inversely proportional to the islet yield per donor,21 fewer islets can be isolated from the already scarce donor pool, further limiting broad clinical applicability of pancreatic islet transplantation. The recognition of these problems has renewed the interest in the search for an alternative site for implantation, such as the intramuscular site and the omental pouch.22

Bone marrow (BM) may represent an ideal alternative site for pancreatic islet transplantation, thanks to its protected and extravascular (but well-vascularized) microenvironment.23,24 Because of its broad distribution and easy access, BM has the potential to overcome not only the physiologic loss of islets, but also the technical limitations and complications encountered with the intraportal infusion.20 To address the potential of BM as an alternative site for pancreatic islet transplantation, we implanted syngeneic pancreatic islet isografts (C57BL/6 islets to C57BL/6 mice) into BM of diabetic recipients and assessed short- and long-term graft survival, function, and safety in comparison with the liver site. The results show that the BM is a more suitable site than the liver for the implantation of islets in this model.

Methods

Islet isolation and culture

Pancreatic islets were isolated from C57BL/6 or BALB/c mice (9 weeks old, 20–22 g; Charles River Laboratories) by a collagenase digestion


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method. Briefly, 2 mL of cold Hanks buffer/collagenase type V solution (1 mg/mL; Sigma-Aldrich) was infused into the pancreatic duct in situ, and the removed pancreas was digested at 37°C for 15 minutes. Islets were purified on a discontinuous Ficoll gradient (Sigma-Aldrich). The islets (250 islets/mL) were cultured freely floating (37°C, 5% CO2) in medium RPMI 1640 (BioWhittaker) supplemented with 1-glutamine (Sigma-Aldrich), penicillin-streptomycin (1000 U/mL to 10 mg/mL, Sigma-Aldrich), and 10% (vol/vol) fetal calf serum (HyClone) for 20 to 24 hours before the transplantation. Islet purity was more than 90%.

**Mice**

Male C57BL/6 mice (9 weeks old, 20-22 g; from Charles River Laboratories) were used as recipients. Mice were made diabetic (nonfasting blood glucose levels between 400 and 600 mg/dL) with intravenous streptozocin (STZ) injection (175 to 200 mg/kg; Sigma-Aldrich) 1 week before transplantation. Blood glucose measurements were performed using a Glucometer Elite (Bayer). The animals had free access to tap water and pelleted food throughout the course of the study. The animal ethics committee of San Raffaele Scientific Institute approved all experiments.

**Islet transplantation**

Islets were transplanted via the portal vein, as previously described.25 Islets were transplanted in the BM as follows: recipients were anesthetized with isoflurane, a 0.5-cm longitudinal incision was made in front of the right knee, and the plate of the femur was exposed and trepanized with a 3/32-inch shank carbide burr in the direction of the medullar channel. Islets packed in PE-50 polyethylene tubing (BD Biosciences) were then introduced into the medullar channel. When the tubing was withdrawn, the cluster of islets was left in the medullar channel by the vacuum effect. The skin was closed with 4-0 silk.

**Evaluation of graft function**

Blood sugar levels were measured 15, 30, and 60 minutes after the end of the surgical procedure, daily for the first week, and then every second day after transplantation. Surgical death was defined as death within the first 7 days after transplantation. Euglycemia was defined as nonfasting blood glucose levels less than 200 mg/dL for 2 consecutive measurements after islet transplantation. An intravenous glucose tolerance test (IVGTT) and an oral glucose tolerance test (OGTT) were performed at 1, 3, 6, and 12 months after transplantation to evaluate the function of the grafted islets. IVGTT was initiated after a 16-hour fast; mice were given glucose (0.5 g/kg) by tail vein injection. Blood samples were obtained at 0, 1, 5, 15, 20, 30, and 60 minutes after injection and were used to determine glucose concentrations. From the IVGTT, the glucose tolerance was quantified from the glucose elimination constant (KG; expressed as percentage of elimination of glucose per minute) as the reduction in circulating glucose between 1 and 15 minutes (KG1-15) after intravenous administration, following logarithmic transformation of the individual plasma glucose values.26 A similar estimation was performed for the total 1- to 60-minute glucose disappearance rate (KG1-60). This parameter indicates the rate of glucose disappearance during the whole test, when the delayed insulin effect is properly accounted for. OGTT was initiated after a 4-hour fast; mice were given glucose (1 g/kg) by oral gavage. Blood samples were obtained at 0, 10, 20, 30, 60, 90, and 120 minutes after glucose administration and were used to determine glucose concentrations. The area under the curve (AUC) for glucose during OGTT was calculated using the trapezoidal method (baseline = 0 minutes). Serum insulin and glucagon were determined by enzyme-linked immunosorbent multiple assay (Mouse Endocrine Lincolnplex Kit; Linco Research). The minimal modeling of insulin and glucose data from 4-hour fasting blood sample was used to assess the β-cell function. Homeostatic model assessment (HOMA) provided equations for estimating β-cell function and insulin resistance27 and was estimated using the Oxford HOMA calculator (http://www.dtu.ox.ac.uk).

**Evaluation of graft morphology**

For morphologic investigations, the recipients were killed 1 year after transplantation. Liver, pancreas, and femur were procured and fixed in 10% buffered formalin and processed routinely for histology. Femurs were placed in Fix Decal (Pro-EKo) overnight to decalcify the bones before embedding in paraffin. Histologic sections were stained with hematoxylin and eosin, anti-insulin guinea pig primary polyclonal antibody (DakoCytomation), anti-glucagon rabbit primary polyclonal antibody (NovoCastra Laboratories), and anti-Ki-67 rabbit primary monoclonal antibody (Thermo Fisher Scientific). The peroxidase-antiperoxidase immunohistochemistry method (PAP Kit; DakoCytomation) or AffiniPure anti–guinea pig tetramethyl-xylenol-diamine-5 (and 6)-isothiocyanate secondary antibodies (Jackson Immuno-Research Laboratories) were used for detection. For quantitative analysis, all immunoreactive cells were analyzed using a Leica DMIIRE2 microscope equipped with a color video camera connected to a computer (Hewlett-Packard) and quantified using Axiovision 4.4 (Carl Zeiss). Islet size was calculated with the image analysis of single measured islets and expressed as area in μm2. For the determination of the fractional β-cell area, each tissue section was imaged at ×40 magnification (×4 objective), and the ratio of the β-cell area/tissue area was digitally quantified. The insulin and glucagon cell numbers were normalized by the total islet cell number of corresponding islets and expressed as its percentage. β-cell size was calculated by dividing insulin-positive areas for the number of the nuclei. Ki-67+ insulin+ cells were reported as percentage of total insulin+ cell. Between 20 (minimum) and 30 islets per case were calculated.

**Evaluation of hematopoietic activity of BM**

Platelets, white blood cells (WBCs), red blood cells, hemoglobin, and hematocrit values were counted with an automated cell counter (HeCo VetC; SEAC); BM cells were isolated by flushing femurs of treated and control mice. For colony-forming cell (CFC) assay, 2.5 × 104 cells were suspended and grown in Methocult GM F-M3434 (StemCell Technologies) in triplicate, and colonies were scored at 10 to 14 days after plating. BM cell phenotyping was carried out by fluorescence-activated cell sorting analysis, after labeling with fluorochrome-conjugated antibodies against B220, CD3, Gr1, Ter119, CD71, c-Kit, Sca1, and Annexin V (BD Pharmingen).

**Lymphocytic choriomeningitis virus infection and related procedures**

The lymphocytic choriomeningitis virus (LCMV) strain Armstrong28 was used in this study. C57BL/6 mice were infected intravenously with 2 × 106 plaque-forming units of LCMV and weighted daily after infection. Whole blood was collected daily from the retro-orbital plexus, and, in addition to glycemia levels, platelets, WBCs, and hematocrit values were counted with an automated cell counter. Serum LCMV RNA was analyzed by quantitative real-time polymerase chain reaction, as described,26 using the following LCMV-specific primers: 5′-CTCTTTTCTCCCAAGAGAAGACTAAG-3′ and 5′-TCCATTGGGTCCAGGCAAATAC-3′. Single-cell suspensions were prepared from spleens harvested at day 8 postinfection, and intracellular interferon γ (IFN-γ) staining on CD8+ T cells was performed in the presence or absence of the H2Kb-restricted immunodominant glycoprotein 33 peptide, as previously described.26 Samples were analyzed with a FACScalibur flow cytometer, and the data were processed using the CellQuest software (BD Immunocytometry Systems).

**Evaluation of mouse bone: peripheral quantitative computed tomography analysis**

Peripheral quantitative computed tomography (pQCT) measurements were performed using a Stratec Research SA® pQCT scanner (Stratec Medizintechnik) with a voxel size of 70 μm and a scan speed of 3 mm/second. The graft-bearing femur and the contralateral femur of each mouse were scanned in the horizontal plane using 4 consecutive cross-sectional images at 3.75 mm, 4.25 mm, 5.5 mm, and 6.5 mm proximal to the distal end of the femur. These scanning sections were chosen as representative of graft implant site, and the relevant pQCT parameters included the measurement of the trabecular compartment only at the 2 most distal sections. The scans were analyzed with pQCT software 6.00 using contour mode 2 and peel mode 2 with a threshold of 350 mg/cm³ for the calculation of trabecular and total bone parameters at the metaphysis and with a threshold of 710 mg/cm³.
for cortical bone parameters at the diaphysis. The different thresholds of 350 and 710 mg/cm² for the metaphysis and diaphysis, respectively, were established to account for partial volume effect. The cortical bone density is lower at the metaphysis than at the diaphysis due to the thinner cortex. The threshold was therefore adjusted according to the cortical density to optimize accuracy. The polar strength strain index (SSI) was calculated by threshold was therefore adjusted according to the cortical density to establish for partial volume effect. The cortical bone density is therefore adjusted according to the cortical density to establish for changes in material properties, the section modulus was normalized for this value in pQCT software Version 2.7.

Statistical analysis

Data were generally expressed as mean standard error or median (Min value–Max value). Differences between parameters were evaluated using one-way analysis of variance (ANOVA) test, followed by Bonferroni post hoc test. Multivariate tests of repeated measures ANOVA were used to compare glycemia during the time. The χ² test was used for categorical variables. Kaplan-Meier analysis was used to compare normoglycemia gain. For all analyses, a 2-tailed P value of .05 was considered significant. Statistical analyses were performed using the Statistical Package for Social Science, Version 13.0 (SPSS).

Results

Islet transplantation in BM

Twenty-four STZ-treated severely diabetic C57BL/6 mice (nonfasting glycemia before transplantation: 509 ± 116 mg/dL) were transplanted with 500 equivalent syngeneic C57BL/6 islets (EI) in BM (Figure 1A). Transplants resulted in normalization of blood glycemia and promptly reverted to hyperglycemia when the graft-bearing BM was removed (Figure 1B). Histologic inspection of femur diaphysis revealed the presence of numerous insulin-immunoreactive cells at the site of implantation (Figure 1C), and dithizone staining of BM in the removed femur showed the presence of insulin-producing islets (Figure 1B). We confirmed these results with a second experiment in which 20 STZ-treated severely diabetic C57BL/6 mice (nonfasting glycemia before transplantation: 519 ± 97 mg/dL) were transplanted with 500 equivalent syngeneic C57BL/6 islets from the same isolation and therefore for changes in material properties, the section modulus was normalized for this value in pQCT software Version 2.7.

Figure 1. C57BL/6 syngeneic islet transplantation in BM. (A) Nonfasting glycemia of STZ-treated severely diabetic C57BL/6 mice transplanted with 500 equivalent islets into the BM (n = 24). (B) Left, Nonfasting glycemia of STZ-treated severely diabetic C57BL/6 mice transplanted in BM with 500 equivalent islets. BM of right femur was removed 9 months after transplantation (arrow; n = 6). Values shown are means ± SEM. Right, Dithizone staining of right femur BM contents after removal at 30 days after transplantation. (C) Histologic appearance (<10) of islets in BM 3 months after transplantation. Immunofluorescent images show insulin staining (red). Scale bar 200 μm. (D) Nonfasting glycemia reduction in the 6 weeks after transplantation with 500 equivalent islets according to transplant site. Intra-BM: islets infused directly into the BM, n = 10. Data are expressed as box plots of mean glycemia of the single weeks. Statistical analysis was performed by tests of repeated measures ANOVA.

Next, we performed a histologic time-course study of BM after syngeneic islet transplantation (supplemental Figures 1 and 2, available on the Blood website; see the Supplemental Materials link at the top of the online article). Immediately after infusion, islets appeared to be healthy, stained strongly for insulin, and never induced the formation of thrombi. Absence of pathologic changes in islets and surrounding BM tissue was still evident by 24 hours after transplants, a time point in which necrotic areas appear in the central region of islets infused into liver and ischemic necrosis of liver tissue is already evident. Note also that BM tissue-remodeling activity was detected at this and later time points exclusively in proximity of the bone hole performed for the islet infusion. After 24 to 72 hours, no evidence of leukocyte infiltration around islets was present (something that occurs in syngeneic livers in response to ischemic necrosis), and the situation remained unchanged until at least day 7.

Next, we transplanted smaller numbers of C57BL/6 islets either in the BM or liver of C57BL/6 mice to best compare their functional performance at these sites. Ninety-one STZ-treated severely diabetic C57BL/6 mice (mean nonfasting glycemia before transplantation: 519 ± 97 mg/dL) were transplanted with 500 equivalent syngeneic C57BL/6 islets from the same isolation and therefore for changes in material properties, the section modulus was normalized for this value in pQCT software Version 2.7. Statistical analysis was performed using the Statistical Package for Social Science, Version 13.0 (SPSS).
The probability and the median time to reach euglycemia (<200 mg/dL) in our marginal islet mass transplantation were as follows: 52% and 85 days for transplants of 125 EI into the BM and 21% and less than 180 days for transplants of 125 EI into the liver (P = .05, log rank test in Kaplan-Meier analysis); 76% and 14 days for transplants of 250 EI into the BM and 43% and 57 days for transplants of 250 EI into the liver (P = .04, log rank statistic in Kaplan-Meier analysis). Multivariate Cox Regression Analysis, including site, islet number, and recipient pretransplant glycemia,29 showed that significant independent factors for reaching euglycemia were as follows: (1) the BM versus liver as recipient site (odds ratio, 2.5; 95% confidence interval, 1.2-5.2; P = .02); (2) the infusion of 250 EI versus 125 EI (odds ratio, 2.2; 95% confidence interval, 1.1-4.3; P = .02); and (3) the pretransplant glycemia (odds ratio, 0.996; 95% confidence interval, 0.993-0.999; P = .01). Finally, mouse survival was significantly higher in mice that received islets in the BM site (P = .05, log rank statistic adjusted for islet number in Kaplan-Meier analysis; Figure 2B).

Because impure preparations of islets are often transplanted in humans, we also examined outcome for semipure syngeneic islet grafts in the BM site (n = 19 diabetic mice receiving 200 pure equivalent islets versus 19 diabetic mice receiving 200 equivalent islets mixed with acinar tissue at a 1:1 ratio; supplemental Figure 3). None of the mice died from the surgical procedure. The proportion of mice achieving normoglycemia was similar in both groups, suggesting that the BM site was likely to be suitable also for impure islet preparations.

Graft survival and glucose metabolism after syngeneic islet transplantation in BM and in liver

The probability to reach normoglycemia was improved when islets were transplanted in the BM compared with the liver in a minimal mass model. Nevertheless, it is also important that the quality of normoglycemia when reached is at least comparable in both sites. In mice that reached euglycemia (28 of 47 for intra-BM and 10 of 32 for intraliver), islet transplant gradually decreased nonfasting glycemia to normal level within 6 months and maintained function thereafter (1 year) with glucose levels comparable with normal nondiabetic control mice (Figure 3A). In mice that reached eugly-

![Figure 2. Comparison of outcomes after islet transplantation into the BM and liver in minimal mass models. C57BL/6 recipients were transplanted with 125 or 250 syngeneic equivalent islets either into the portal vein (intraliver: 125 EI n = 20, 250 EI n = 21) or into the BM (intra-BM: 125 EI n = 31, 250 EI n = 25). Mice that died due to the surgical procedure (4 of 56 and 8 of 41, respectively, for intra-BM and intraliver) were excluded from the analysis. (A) Kaplan-Meier analysis for the gain of normoglycemia (< 200 mg/dL). (B) Kaplan-Meier survival analysis. Differences between the intra-BM and intraliver models were tested using the log rank statistic adjusted for islet number.

![Figure 3. Graft survival and long-term glucose metabolism after islet transplantation in BM and liver. (A) Blood glucose profile of normal C57BL/6 mice (Ctrl, white; n = 11), and mice that achieved normoglycemia after receiving islets intra-BM (dark gray; n = 47) or intraliver (gray; n = 32). Data are expressed as mean (line) and ±1 SD (area). *Statistical analysis was performed by tests of repeated measures ANOVA considering intervals of 2 months. **Statistical analysis was performed by tests of repeated measures ANOVA adjusted for islet number considering intervals of 2 months. * P < .05; ** P < .001; ns = not significant. (B) Fasting glycemia (mg/dL), insulin (pM), glucagon (pM), β-cell function (HOMA2-B, %), and insulin resistance (HOMA-IR) in normal C57BL/6 mice (Ctrl, white), and mice that achieved normoglycemia after receiving islets intra-BM (dark gray; n = 47) or intraliver (gray; n = 32). Assays were performed 6 to 9 months after islet infusion. Data are expressed as box plots. Statistical analysis was performed using 1-way ANOVA test, followed by Bonferroni post hoc t test. * P < .05; ** P < .001.
cemia, glycemia, insulin, glucagon, HOMA-B, and HOMA-IR evaluated after 4 hours of fasting at 6 to 9 months after transplantation were also similar in mice that achieved normoglycemia via islets into BM or islets into liver (Figure 3B). No differences were observed between mice reaching normoglycemia in the intra-BM and intraliver models. Mice that did not reach euglycemia showed glucose levels constantly higher than control, and mice eventually died as a result of uncontrolled diabetes (data not shown).

The time course of the effects of islet transplantation on glucose tolerance after IVGTT and OGTT is illustrated in Figures 4 and 5. No differences were observed in the euglycemic mice when liver versus bone marrow site was compared for both IVGTT and OGTT parameters. After 1 month, glucose elimination constant between 1 and 15 minutes (KG1-15) and 1 and 60 minutes (KG1-60) was significantly reduced in intra-BM and intraliver transplanted mice compared with control mice (Figure 4). Subsequently, in mice reaching euglycemia, the KG1-60 gradually improved in both intra-BM and intraliver models, reaching control values at 3 months after transplantation. The KG1-15 in both intra-BM and intraliver also improved, but remained significantly lower than in control mice during all the period of observation. OGTT at 1, 3, 6, and 12 months after transplantation in BM and liver. Glucose tolerance in an OGTT was measured at 1, 3, 6, and 12 months after islet transplantation into BM or liver. Shown are the OGTT glycemic profile and AUC for glucose in mice that achieved euglycemia (28 of 47 for intra-BM and 10 of 32 for intraliver). The yellow shaded area represents the range of values in healthy control mice (Ctrl, n = 11). Data are expressed as mean for glycemic profile and boxplots for glucose tolerance. Statistical analysis was performed using 1-way ANOVA test, followed by Bonferroni post hoc t test. *P < .01 versus control.

Morphologic study of islets 1 year after transplantation

Islet morphology at 1 year after transplantation was evaluated in BM and liver in normoglycemic mice (6 mice per group) and in the pancreas of 6 control mice (Figure 6). Islets were detected in all femurs analyzed from the intra-BM group. Morphologically intact islets were present in femur diaphysis embedded in the cellular component of the BM (Figure 6A). For the intraliver model, several islets were present within the liver parenchyma and close to blood vessels of the portal tract. Generally, the islets in the liver did not maintain their original shape and appeared spread in the tissue (Figure 6A). As expected, the fractional β-cell area was significantly different between tissues, as follows: 14.2% in BM, 0.7% in liver, and 1.4% in control (P < .01 BM vs control and BM vs liver). Intra-BM transplants were characterized by approximately 4.5-fold increase in islet size compared with pancreatic islets (87 055 μm² vs 18 596 μm², P < .001, intra-BM vs control). An increased islet size in the liver (approximately 3.5-fold) was also observed (67 929 mm², P < .05 vs control). Islets in the BM also had an increased density of β cells: BM, 8.9 ± 0.4 × 10⁻³ nuclei/μm²; liver, 6.7 ± 0.5 × 10⁻³ nuclei/μm²; control, 7.1 ± 1.2 × 10⁻³ nuclei/μm² (P = .025, intra-BM vs control; P < .01, intra-BM vs liver transplantation). The frequency of β-cell replication (Ki67/insulin cells) at the 12-month time point was low in all groups.
Figure 6. Graft morphology. (A) Histologic appearance (top, ×5, scale bar, 400 μm; bottom, ×20; scale bar, 100 μm) of islets in control pancreas, BM, and liver 1 year after transplantation. (B) Fractional β-cell area, islet size, islet composition, β-cell proliferation, and nucleus density of insulin-positive area in mice reaching normoglycemia after islet transplantation in BM (n = 6) and liver (n = 6) 1 year after islet infusion, and in the pancreas of control mice (n = 6). Data are expressed as mean ± SD. Statistical analysis was performed using 1-way ANOVA test, followed by Bonferroni post hoc test. *P < .01 versus control; $P < .05 versus intra-liver.

To determine whether the presence of islets in the BM could affect hematopoietic activity, single-cell suspensions were prepared from BM by flushing the right femur (site of islet transplantation) and left femur (sham surgery) of 6 normoglycemic mice 1 year after islet transplantation. The cellularity of the right femur did not differ significantly from that of the left (7.9 ± 1.3 × 10^6 vs 7.8 ± 1.6 × 10^6 cells, P = .9). Analysis of BM cell subpopulations showed the same percentage of B220^−, CD3^+^, Gr1^−^, Ter119^−^, c-Kit^+^, Sca1^+^, and Annexin V^+^ cells (supplemental Figure 4A). By CFC assay, the frequency of hematopoietic progenitors was 22.3 ± 7.3 × 10^−4 BM cells and 25.5 ± 8.1 × 10^−4 BM cells, respectively, for the right and left femur (P = .37). The types of hematopoietic colony-forming units (burst-forming unit–erythrocyte, colony-forming unit–granulocyte macrophage, colony-forming unit–granulocyte erythroid macrophage megakaryocyte) were equally represented (supplemental Figure 4A). We also evaluated peripheral blood cellularity 1 year after islet infusion in normoglycemic mice receiving islets in BM (n = 17), liver (n = 9), and in control mice (n = 11; supplemental Figure 4B). WBC and platelet counts were similar among the 3 groups. A moderate reduction in red blood cell counts, hemoglobin, and hematocrit was observed in mice receiving islets in the BM and in the liver, suggesting an association with the previous condition of diabetes rather than with the site of islet transplantation.

Next, we tested whether the presence of islets affects the capacity of BM to respond to virus-induced aplasia. At 1 year after transplantation, 6 of the normoglycemic mice that had received islet transplants in the BM were acutely infected with LCMV (BM-Tx-LCMV). LCMV is a noncytopathic mouse pathogen causing a systemic infection that targets most organs and tissues (including BM), but not pancreatic β cells. Acute LCMV infection is associated with an IFN-α/β–dependent BM aplasia that produces a transient pancytopenic state. The initial lymphopenia is rapidly reversed, such that within 1 week of LCMV exposure, the mice display lymphocytosis and clear the infection through a vigorous virus-specific cytotoxic T-lymphocyte (CTL) response. Results obtained in BM-Tx-LCMV were compared with those of age-matched, nontransplanted C57BL/6 mice that were either infected (LCMV) or not (control) (n = 6).

As shown in Figure 7A, all animals remained normoglycemic throughout the duration of the experiment (8 days). Weight loss was moderate and comparable between BM-Tx-LCMV and LCMV mice. Also comparable between the 2 groups of animals were the changes in platelet and WBC counts, which reached the expected nadir at day 4 postinfection and rebounded thereafter. In keeping with these results, there was no change in the hematopoietic activity of BM obtained from the femurs of BM-Tx-LCMV and LCMV mice, as can be inferred by the similar percentages of BM precursors detected at day 8 postinfection either by flow cytometry (Figure 7B) or CFC assay (data not shown). Note also that this was true when we compared the islet-containing right femur of BM-Tx-LCMV mice with the contralateral one in which islets were not present (Figure 7B). Importantly, BM-Tx-LCMV and LCMV mice cleared the virus with similar kinetics and by day 8 postinfection mounted a LCMV-specific CTL response that was quantitatively identical (Figure 7C).

pQCT measurements: cross-sectional in vivo evaluation of graft-bearing and sham-treated contralateral femurs

Right femur (site of islet transplantation) and left femur (sham surgery) of 11 normoglycemic mice were analyzed 9 months after islet transplantation. pQCT analysis showed a significant decrease (P < .01) of the cortical density that was localized at all cross-sectional scanning sites (supplemental Figure 5). Cortical thickness was found slightly, but significantly decreased only at the most distal section, that is, the closest section to graft implant site. None.
reaction–based assay with a detection limit of 10^3 copies of LCMV RNA (genome
tation and monitored for the relative content of BM precursors and the presence of
intra-BM islet-transplanted LCMV-treated mice and from controls (age-matched
counts were measured either daily or every other day. (B) Femurs collected from
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that the same technique could be used for islet infusion.
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hematopoietic activity in intra-BM islet-infused mice after infection
with LCMV. (A) Mice receiving islets intra-BM were infected at approximately 1 year
of age with 2 x 10^6 plaque-forming units of LCMV. Glycemia, platelet, and WBC
counts were measured either daily or every other day. (B) Femurs collected from
intra-BM islet-transplanted LCMV-treated mice and from controls (age-matched
mice either infected with LCMV or not) were harvested on day 8 postinfec-
tion and monitored for the relative content of BM precursors and the presence of
insulin-producing β cells. (C) Viral titers in blood harvested at the indicated time
points after infection were measured by a reverse transcription–polymerase chain
reaction–based assay with a detection limit of 10^3 copies of LCMV RNA (genome equivalents [GE]) per mL of total blood. Results are expressed as means of triplicate
measurements ± SD. Assessment by intracellular IFN-γ staining of the percentage
of glycoprotein 33-specific CD8^+ T cells recovered from spleens harvested at day 8
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Figure 7. Hematopoietic activity in intra-BM islet-infused mice after infection
with LCMV. (A) Mice receiving islets intra-BM were infected at approximately 1 year
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Discussion

The liver is the current site for pancreatic islet transplantation, but
presents important limitations. The potential solution to this
problem is to establish an alternative site for clinical islet transplan-
tation. We asked whether pancreatic islets could be engrafted in the
BM, an easily accessible and widely distributed transplant site that
may lack the limitations seen in the liver. The results show that
pancreatic islets can be efficiently engrafted into BM. This is the
first report of BM as a successful site for seeding pancreatic islets
in diabetic recipients. A single work previously described injection
of islet in BM of nondiabetic rats with the detection of positive
staining for insulin up to 21 days after infusion without any
functional evaluation.\textsuperscript{32}

In our study, both the percentage and the timing in reversal of
hyperglycemia were superior after BM infusion compared with
intrahepatic infusion using the minimal mass model. Moreover, with
the exception of a small delay in gaining normal glucose tolerance after OGTT, the quality of glucose metabolism in mice
that reached normoglycemia via intra-BM islet infusion was
similar to that achieved by islet transplant into the liver for all the
parameters evaluated (fasting and not fasting glycemia, blood
insulin, HOMA-B, and glucose tolerance after IVGTT). Based on
our results, we can conclude that the BM site for islet transplantation
has a higher probability to reach euglycemia (2.5-fold increase in a multivariate analysis) than the liver without compromising the
quality of glucose metabolism. This is relevant because the process
of intrahepatic infusion was traditionally considered optimal due to
the supposition that insulin is delivered more physiologically after
intraportal transplantation.\textsuperscript{32-33} However, this argument has recently
been challenged by experimental studies showing that
intraportally transplanted islets respond to glucose stimulation only
when perfused via the hepatic artery; no response is observed after
challenge via the portal vein.\textsuperscript{36} There are also reports on alterations
in islet function after intraportal islet transplantation, such as
a defective glucagon response to hypoglycemia\textsuperscript{37,39} and defective
insulin-glucagon release.\textsuperscript{40}

In relation to the site of engraftment, islet morphology and
and cellular composition showed profound changes after transplantation.
Islets transplanted in BM showed an increased size with a
compact morphology. The most likely explanation for the increased
size lies in the method of cell isolation and infusion. Before the
introduction into BM, islets are packed, and this could allow the
formation of aggregates. This was also supported by the fact that
the increase in islet size was not associated with increased cell
volume. Nevertheless, without time course studies, we cannot
exclude an increased replication of cells within the transplanted
islets leading to the increased size observed at 12 months after
transplantation. The fraction of α and β cells in islets engrafted in
BM was similar to control islets. In contrast, the fraction of α cells
was markedly decreased in the intraportally transplanted islets. The
reason for this selective decrease of the α-cell fraction is unknown,
but is consistent with the observed decreased glucagon response to
hypoglycemia seen in humans after islet transplantation.\textsuperscript{37}

The process of intrahepatic infusion is currently considered
safe, although there is a low risk of portal vein thrombosis and
and elevated portal pressures, in addition to bleeding from the percuta-
nous hepatic puncture site.\textsuperscript{41-44} From a clinical viewpoint, BM
seeding has the potential of being a less invasive, ambulatory
procedure with almost unlimited opportunities for repeated implan-
tation, as well as a low-risk, easy-access site for graft biopsies. This
potential was recently confirmed by the clinical experience of
direct intrabone cord-blood transplantation. Frassoni et al recently
reported a cohort of 32 patients receiving infusion of 20 mL cell
suspension in superior-posterior iliac crest.\textsuperscript{45} The infusion was
performed easily without resistance. Moreover, no side effects,
such as pain, hemorrhage, or infections, were recorded, suggesting
that the same technique could be used for islet infusion.

Because it was suggested that hyperinsulinemia might contrib-
ute to cancer development through the growth-promoting effect
of elevated levels of insulin, it is possible that intra-BM islet
transplantation could increase the risk of proliferative disease. For
this reason, we evaluated the impact of islets on hematopoietic
activity of BM. One year after islet infusion, the cellularity, the histologic appearance, the analysis of cell subpopulation, and the progenitor cell frequency were unaffected by the presence of islets in the BM. It should also be underlined that a potential tumorigenic risk from local hyperinsulinemia is also present after intrahepatic islet infusion. Indeed, in diabetic bio breeding rats receiving intraportal islet transplants with long-term partial function, there was a remarkably high incidence of adenomas, and even hepatocellular carcinoma. The initial stages in the process leading to these pathologic conditions were hepatocyte steatosis and glycogen deposits in the vicinity of the transplanted islets, findings that are analogous to those reported in patients from several transplant centers.

We also took into consideration the consequences of BM islet infusion on the capacity to respond to virus-induced aplasia and the bone structure. Islets in BM of LCMV-infected mice did not affect hematopoietic activity consequent to aplasia nor CTL-mediated viral clearance. These results also suggest that islets in BM are capable of sustaining those metabolic changes that are likely to occur during the rapid expansion of a very robust adaptive immune response (ie, by day 8 postinfection, secondary lymphoid organs of LCMV-infected mice are much larger in size, and approximately 50% to 70% of all CD8+ T cells are LCMV specific). The islet graft did cause minor alterations of the bone microenvironment with a subsequent volumetric bone density loss at the cortical compartment facing the grafted pancreatic islets. Because bone development predicts bone mass accrual as a consequence of a modeling process during growth, it is conceivable that the decreased cortical density is the outcome of enhanced osteoclastogenesis rather than a reduced osteoblastogenesis, as the latter should have resulted in a reduced outward shift of the bone cortex. Regardless of the mechanism, the issue is whether bone resorption in the cortical compartment is a potential contraindication to the use of BM as a site of engraftment. From a quantitative point of view, the bone loss was between 4 and 6% compared with the contralateral femur after 9 months. Moreover, it should be considered that in an eventual clinical application, the site of infusion would be the iliac crest. Because iliac crest is not a weight-bearing skeletal site, a limited loss of its cortical density is not expected to have important clinical consequences. Moreover, bone per se might constitute the critical microenvironment for pancreatic islet function and survival. It has been in fact shown that picomolar amounts of osteocalcin, the osteoblast-specific secreted molecule, are sufficient to enhance the expression of the insulin genes and β-cell proliferation markers in cell-based assays using isolated pancreatic islets. For all safety considerations and indeed for evaluation of the suitability of the BM site in humans, it should be considered that islet transplantation is usually performed with allogenic islets and under immunosuppression. Islet function after transplantation is affected by immunosuppression, and our preliminary results suggest that this is also the case when syngeneic islets are transplanted in the BM (L.P., unpublished observations).

In conclusion, we show that pancreatic islets can be engrafted into the BM, thus opening a research line with potentially significant clinical impact not only for the treatment of diabetes, but for other diseases amenable to treatment with cellular transplantation. Because the BM as a site for pancreatic islet grafts can be clinically applicable and, in theory, can solve many of the problems encountered with the intrahepatic location, further research is warranted by the initial findings presented in this study to determine whether the results can be reproduced in large animals and eventually in humans.

Acknowledgments

We thank Prof Claudio Doglioni (Department of Pathology, San Raffaele Scientific Institute) and Dr Fabio Ciceri (Unit of Hematology and Bone Marrow Transplantation, Department of Oncology, San Raffaele Scientific Institute) for the useful discussion of these experimental results.

This work was supported by the European Union (DIAPREPP Project, HEALTH-F2-2008-202013), Telethon Italy and the Juvenile Diabetes Research Foundation (J011Y01), Italian Telethon Foundation (Telethon Institute for Gene Therapy grant) and CARIPLO Foundation, and the Deutsche Forschungsgemeinschaft-Center for Regenerative Therapies Dresden, Cluster of Excellence at Dresden University of Technology. V.S. is enrolled as a PhD student at the Ludwig-Maximilians University of Munich.

Authorship

Contribution: E.C. performed research; R.M. performed research; A.M. performed research; V.S. analyzed data; G.F. designed research; C.W.L. performed research; E.M. performed research; A.R. designed research; M.P. analyzed data; G.S. performed research; L.G.G. designed research and wrote the paper; E.B. designed research and wrote the paper; and L.P. designed research, performed research, and wrote the paper.

Conflict-of-interest disclosure: The authors declare no competing financial interests.

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References

36. Lauer J, Jansson L, Carlsson PO. Islet transplants implanted intraportally into the liver are stimulated to insulin and glucagon release exclusively through the hepatic artery. Am J Transplant. 2006;6:967-975.
Bone marrow as an alternative site for islet transplantation

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