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Marc R. Hammerman

Washington University School of Medicine in St. Louis

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Organogenetic tolerance

Marc R. Hammerman
Renal Division; Departments of Medicine and Cell Biology and Physiology; Washington University School of Medicine; St. Louis, MO USA

Key words: beta cell, diabetes mellitus, non-human primates, transplantation, xenotransplantation

Abbreviations: E, embryonic day; ES, embryonic stem cell; GALT, gut associated lymphoid tissues; LEW, lewis; STZ, streptozotocin

Transplantation therapy for humans is limited by insufficient availability of donor organs and outcomes are complicated by the toxicity of immunosuppressive drugs. Xenotransplantation is a strategy to overcome supply problems. Implantation of tissue obtained early during embryogenesis is a way to reduce immunogenicity of transplants. Insulin-producing cells originating from embryonic pig pancreas obtained very early following initiation of organogenesis (embryonic day 28 (E28)) engraft long-term in non-immune suppressed diabetic rats or rhesus macaques. Recently, we demonstrated engraftment of morphologically similar cells originating from adult porcine islets of Langerhans (islets) in rats previously transplanted with E28 pig pancreatic primordia. Our findings are consistent with induction of tolerance to a cell component of porcine islets induced by previous transplantation of embryonic pig pancreas, a phenomenon we designate organogenetic tolerance. Induction of organogenetic tolerance to porcine islets in humans with diabetes mellitus would enable the use of pigs as islet donors with no host immune suppression requirement. Adaptation of methodology for transplanting embryonic organs other than pancreas so as to induce organogenetic tolerance would revolutionize transplantation therapy.

As reviewed previously in Organogenesis,1,2 transplantation of embryonic organ primordia to replace the function of diseased organs offers theoretical advantages relative to transplantation of either pluripotent ES cells, or of fully differentiated (adult) organs. (1) Unlike ES cells, organ primordia differentiate along defined organ-committed lines. There is no requirement to steer differentiation and no risk of teratoma formation. In the case of embryonic pancreas, the glucose sensing and insulin releasing functions of β cells that differentiate from primordia are functionally linked. (2) The growth potential of cells within embryonic organs is enhanced relative to those in terminally differentiated organs. (3) The cellular immune response to transplanted primordia obtained early during embryogenesis is attenuated relative to that directed against adult organs. (4) Early organ primordia are avascular. The ability of cellular primordia to attract a host vasculature renders them less susceptible to humoral rejection than are adult organs with donor blood vessels transplanted across a discordant xenogeneic barrier. And (5) organ primordia differentiate selectively. In the case of embryonic pancreas, exocrine pancreatic tissue does not differentiate following transplantation, obviating complications that can result from exocrine components such as the enzymatic autodigestion of host tissues. The finding that it is possible to transplant pig pancreatic primordia to non-human primates with no immunosuppression requirements3 is very important because it establishes the potential for transplantation in humans with no complications of immunosuppressive therapy.

The physiologies of several porcine organ systems are sufficiently similar to those in humans such that the pig has been suggested to be an organ donor for clinical transplantation.4 Pigs are plentiful, and can be bred in captivity under pathogen-free conditions.5 Porcine insulin works well in humans. Therefore transplanted porcine pancreatic endocrine tissue could be used to treat diabetes in patients. The severity of humoral rejection due to preexisting natural antibodies effectively precludes the use of non-transgenic swine as whole organ (pancreas) donors in primate hosts.6 However, isolated cells such as islets of Langerhans (islets) can be transplanted into humans7,8 or non-human primates7,8 without initiating humoral rejection. Recent experience with pig to primate islet,7 neonatal islet,8 or hCD46 transgenic pig islet9 transplantation shows that sustained insulin independence can be achieved, but unfortunately only through the use of immune suppressive agents that are not approved for humans or would result in an unacceptable level of morbidity.

We have shown that glucose tolerance can be normalized in streptozotocin (STZ)—diabetic (type 1) LEW70-12 rats or ZDF13 (type 2) diabetic rats within 4 weeks following transplantation in mesentery of pig pancreatic primordia obtained very early during embryogenesis [on embryonic day 28 (E28)—just after the organ differentiates and prior to the time dorsal and ventral anlagen fuse] without host immune-suppression. Rats are rendered permanently independent of a requirement for exogenous insulin to maintain normoglycemia. No circulating rat insulin can be detected in STZ-treated rats. Porcine insulin circulates post-transplantation of E28 pig pancreatic primordia (embryonic pancreas) and levels increase after a glucose load. Cells expressing insulin and porcine proinsulin mRNA with β-cell morphology

Correspondence to: Marc R. Hammerman; Email: mhammerm@dom.wustl.edu
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Glucose (mg/dl)

Time (minutes)

Pre-STZ
Post-STZ
Post-TX

0 15 30 60

Figure 1. Intravenous glucose tolerance in rhesus macaques. Glucose in peripheral venous blood was measured prior to intravenous infusion of 0.5 g/kg over 30 seconds as 50% dextrose (Time 0) and at several times after infusion in three fasted rhesus macaques either prior to administration of 60–140 mg/kg intravenous STZ (Pre-STZ); 5 days following administration of STZ (Post-STZ); or 3 months following transplantation of 20–40 E28 pig pancreatic primordia in mesentery of STZ diabetic macaques (Post-TX). Levels measured at 60 minutes after intravenous glucose administration were not different from levels measured at Time 0 in Pre-STZ or Post-TX groups [Bonferroni Multiple Comparisons Test, p < 0.05, two tailed analysis (GraphPad Instat 3, San Diego CA)]. In contrast, levels measured at 60 minutes after intravenous glucose infusion were elevated relative to those measured at Time 0 in the Post-STZ group. Data are shown as mean ± SE (3 macaques).

Figure 2. Photograph of kidney from a STZ-treated rat transplanted previously with embryonic pig pancreas in mesentery. The photo was taken 4 weeks after implantation of pig islets in kidney. The whitish well-demarcated graft (white arrow) and the origin of a venous blood vessel (black arrow) are shown. Reproduced with permission from the American Society for Investigative Pathology.12

Intact porcine islets do not engraft following renal subcapsular implantation. However, a population of cells originating from donor islets with β-cell morphology that express insulin and porcine proinsulin mRNA engraft in kidneys of rats transplanted previously with E28 pig embryonic pancreas. Our observations are consistent with induction of tolerance to a cell component of adult porcine islets by previous transplantation of E28 pig pancreatic primordia in rats.12

Figure 3 shows sections from a kidney from a STZ-diabetic rat that had been transplanted with embryonic pig pancreas in mesentery, taken 4 weeks after implantation of islets in kidney. A distinct, whitish well-demarcated graft (white arrow) and the origin of a venous blood vessels that radiate from the graft out into the renal parenchyma (black arrow).

In lieu of increasing the number of transplanted primordia in diabetic rhesus macaques, we embarked on a series of experiments to determine whether porcine islets, a more easily obtainable and possibly more robust source of insulin-producing cells, could be substituted in animals rendered tolerant to embryonic pig pancreas. Our first step was to determine using rats, whether engraftment of cells originating from E28 pig pancreatic primordia renders hosts tolerant to the same or similar cell component present in porcine islets from adult swine (adult islets). To this end, we implanted adult porcine islets beneath the renal capsule of rats that previously had been transplanted with E28 pig pancreatic primordia in mesentery.12
probes. Hybridization occurs with cells hybridized to the former (arrows), but not the latter probe.

The contralateral (nontransplanted) kidney from a STZ-diabetic rat transplanted previously with embryonic pig pancreas in mesentery and subsequently with pig islets in the ipsilateral kidney is shown in Figure 4A–D. Sections are stained using anti-insulin antibodies (Fig. 4A and C) or control serum (Fig. 4B and D). A low magnification view of the contralateral kidney shows normal renal morphology with no evidence of engrafted tissue (Fig. 4A). A higher power view confirms that, in contrast to what is observed in the transplanted kidney (Fig. 3A) there is no expansion of the subcapsular space in the contralateral kidney (Fig. 4C, arrowheads) and no cells are present with β-cell morphology as shown in Figure 3C. Figure 4E and F shows the subcapsular space of a kidney from a rat implanted with porcine islets four weeks previously with no prior transplantation of E28 pig pancreatic primordia. Sections are stained using anti-insulin antibodies (Fig. 4E) or control serum (Fig. 4F). Insulin staining of proximal tubules (PT) is evident in Figure 4E. The subcapsular space (Fig. 4E arrowheads) is expanded relative to that depicted in Figure 4A and C. However there are no cells with β-cell morphology that stain for insulin (Fig. 4E).

To provide additional evidence that pig cells are present in kidneys and mesenteric lymph nodes of rats transplanted with pig pancreatic primordia and subsequently with porcine islets, we performed fluorescent in-situ hybridization using a probe specific for the pig X chromosome. Shown in Figure 5A (arrows) are pig X chromosomes in nuclei of cells from a normal porcine pancreas (positive control). Figure 5 (arrows) shows pig X chromosomes in the nuclei of cells in the renal subcapsular space [between a tubule (T) and the renal capsule membrane (RC) in Fig. 5B]; and outside of a germinal center (GC) of a mesenteric lymph node (Fig. 5C). There are no cells containing pig X chromosomes in renal cortex from the transplanted kidney (Fig. 5D), consistent with the subcapsular localization of the insulin positive and porcine proinsulin mRNA containing cells (Fig. 3), and with species specificity of the pig X chromosome probe (host rats are females).

We have shown previously using electron microscopy that cells with β-cell morphology containing granules, some of which have a crystalline core surrounded by a clear space, are present in mesentery of rats following transplantation of E28 pig pancreatic primordia. To ascertain whether similar cells are present in kidneys of rats transplanted with E28 pig pancreatic primordia and subsequently with porcine islets, we performed electron microscopy. Shown in Figure 6A (arrow) is a cell in the renal subcapsular space with β-cell morphology [containing granules (250–400 nm) which have a crystalline core surrounded by a clear space]. A renal tubule (T) is labeled. A mononuclear cell infiltrate in the renal subcapsular space consists predominantly of macrophages. Two are labeled (arrowheads). Figure 6B is a high-power view of two others. Figure 6C shows a high power view of the granules with a crystalline core surrounded by a clear space.

To our knowledge, ours is the first report describing sustained survival of β cells following transplantation of porcine islets in non-immune suppressed immune sufficient rodents. In previous studies that did not employ transplantation of E28 pig pancreatic primordia prior to implantation of islets or islet cell clusters complete destruction of grafts was evident within two weeks. Pre-immunization of rats by subcutaneous injection of porcine islet cell clusters accelerates the rejection of islet...
cell component that differentiates into insulin-producing cells) of adult porcine islets implanted in LEW rats by previous transplantation of E28 pig pancreatic primordia.

Though not observed following xenotransplantation under all conditions, host tolerance to early stage pancreatic progenitors has been reported twice previously. Eloy et al. described normalization of glucose post-transplantation of E15, but not E18 embryonic chick pancreas into non-immune suppressed STZ-diabetic immune competent rats. Abraham et al. described successful xenograftment in multiple organs of human pancreatic islet-derived progenitor cells infused in non-immunosuppressed immune competent mice. It is possible that xenotransplantation of fetal pancreas is particularly suited to induction of tolerance. However, neither Eloy et al. nor Abraham et al. nor we define an immunological mechanism.

Although the antigenicity of fetal tissues may be less than that of corresponding adult tissues, animal data suggest the reduction is not enough by itself to ensure permanent graft survival. Thus, the use of embryonic tissue (pancreas) per se cannot explain our findings. Host immune suppression is required for successful engraftment of embryonic pig pancreas in rodents or non-human primates carried out using methodology different from ours. Therefore, it is likely that one or more differential factors in the methodology we employ is critical for engraftment of embryonic pig pancreas without an immune suppression requirement. Such factors could include the developmental stage of embryos from which primordia are obtained, the number of pancreatic primordia transplanted, the manner in which embryonic pancreas organs are incubated in vitro prior to implantation, the diabetic status of the host, the transplantation site and methodology for securing the implants in place and the stringency by which glucose levels in diabetic hosts are controlled post-transplantation.

Korsgren has proposed that rejection of an islet cell xenograft is dependent on two different cellular mechanisms. The first is recognition of pig MHC molecules by cytotoxic lymphocytes via both direct and indirect pathways of antigen recognition, as occurs following rejection of an allograft. The second is an immune response characterized by T cell dependent infiltration of macrophages with histopathological characteristics of delayed-type hypersensitivity. In non-immune suppressed rodents transplanted with porcine islets, the second mechanism predominates. In non-immune suppressed non-human primates, rejection is dominated by the first. However the first process is inhibited in immune suppressed non-human primates and the second, less sensitive to immune suppression, is revealed.

The presence of macrophages beneath the renal capsule of rats transplanted with porcine islets subsequent to E28 pig pancreatic primordia (Fig. 6) may reflect the second cellular mechanism described by Korsgren in the context of engraftment of a cell component of porcine islets to which rats have been rendered tolerant.

Clusters transplanted subsequently beneath the renal capsule. In contrast, prior transplantation of E28 pig pancreatic primordia in mesentery enables survival of an insulin-expressing cell component of porcine islets implanted subsequently in rat kidneys (Fig. 3).

Schroeder et al. define transplantation tolerance as immune unresponsiveness to the transplanted organ, but not to other antigens, in the absence of ongoing immunosuppression. LEW rats transplanted with E28 pig pancreatic primordia retain reactivity to other porcine xenoantigens (E28 pig renal primordia are rejected). Thus, our findings are consistent with induction of specific tolerance to a cell component (either β cells or a stem

Figure 4. Photomicrographs of the contralateral kidney from a diabetic rat into which embryonic pig pancreas had been transplanted in mesentery and pig islets had been implanted subsequently in the other kidney (A–D) or of a kidney from a diabetic rat in which pig islets had been implanted without prior transplantation of E28 pig pancreatic primordia (E and F) stained using anti-insulin antibody (A, C and E) or control antibody (B, D and F). Arrowheads delineate a normal sized subcapsular space (A and C) or expanded subcapsular space (E). PT, proximal tubule (A, B, E and F). M, medulla (A and B). Scale bars 100 μm (A and B), 10 μm (C and D) or 40 μm (E and F). Reproduced with permission from the American Society for Investigative Pathology.
convergence of endocrine and intestinal contents at this site may explain the link between an autoimmune pathogenesis for type 1 diabetes and environmental provocation.\textsuperscript{31,32} Low doses of orally administered antigen induce antigen-specific peripheral tolerance through active suppression of T cells and induction of clonal anergy. High doses induce tolerance by extrathymic deletion of antigen-reactive T cells.\textsuperscript{33} It was proposed tolerant by prior transplantation of E28 pig pancreatic primordia. Alternatively, since porcine islets are rejected by immune competent rats without prior transplantation of E28 pig pancreatic primordia\textsuperscript{12} the macrophages may represent a population of M\textsubscript{2} phenotype\textsuperscript{24} partially responsible for the tolerance we observe.

We have proposed\textsuperscript{25} that transplantation of E28 pig pancreatic primordia in the mesentery and migration of cells to mesenteric lymph nodes and liver recapitulates events that occur during induction of oral tolerance,\textsuperscript{26-28} the induction of which is dependent on antigen transport via afferent lymphatics into the draining mesenteric lymph nodes.\textsuperscript{26} In effect, we suggest that heterotopic introduction of embryonic pig pancreas in rat or primate mesentery co-opts the function of the gut associated lymphoid tissues (GALT) a complex, redundant\textsuperscript{26-28} and phylogenetically ancient system\textsuperscript{29,30} of which embryonic pancreas is a part,\textsuperscript{31} that under normal conditions induces peripheral tolerance to ingested antigens in jawed vertebrates and their descendants.

Interestingly, GALT may have served similarly to prevent an immune response to insulin-producing cells scattered originally in the gut epithelium of primitive vertebrates\textsuperscript{29,30} and has been proposed to induce tolerance or immune suppression towards islet cell antigens during normal embryonic development.\textsuperscript{31} Developmentally controlled lymphogenesis establishes a preferential trafficking route from the gut to pancreatic lymph nodes, a GALT component, in which T cells can be activated by antigens drained from the peritoneum and the gastrointestinal tract. Intestinal stress modifies the presentation of pancreatic self-antigens in pancreatic lymph nodes. The convergence of endocrine and intestinal contents at this site may explain the link between an autoimmune pathogenesis for type 1 diabetes and environmental provocation.\textsuperscript{31,32}

Low doses of orally administered antigen induce antigen-specific peripheral tolerance through active suppression of T cells and induction of clonal anergy. High doses induce tolerance by extrathymic deletion of antigen-reactive T cells.\textsuperscript{33} It was proposed

\textbf{Figure 5.} Fluorescence microscopy of tissue sections originating from (A) a normal porcine pancreas or (B–D) a diabetic rat that had been transplanted with embryonic pig pancreas in mesentery and subsequently with porcine islets in kidney: (B) a subcapsular section from kidney, T tubule, RC Renal Capsule; (C) a section of mesenteric lymph node, GC, germinal center, INSET enlargement; and (D) renal cortex, T, tubule. Arrows (A–C) delineate pig X chromosomes. Scale bar 10 um (D). Reproduced with permission from the American Society for Investigative Pathology.\textsuperscript{12}

\textbf{Figure 6.} Electron micrographs of rat kidney following sequential transplantation of E28 pig pancreatic primordia in mesentery and implantation of porcine islets in kidney. (A) Subcapsular space. T, renal tubule; Cell containing granules with a crystalline core surrounded by a clear space is delineated by an arrow; Macrophages are delineated by arrowheads. (B) Enlargement of macrophages. (C) Enlargement of granules with a crystalline core surrounded by a clear space. Scale bar 5 um (A). Reproduced with permission from the American Society for Investigative Pathology.\textsuperscript{12}
originally, that oral tolerance depends exclusively on antigen uptake by cells within intestinal Peyer’s Patches.28 However, recently it has been shown that high dose oral tolerance can be induced in the absence of Peyer’s Patches, so long as mesenteric lymph nodes are present.28,34

Given similarities between the host immune responses in rodents and non-human primates to porcine xenografts,23,35 studies are warranted employing non-immune suppressed rhesus macaques transplanted with E28 pig pancreatic primordia3 as hosts for porcine islets. Such experiments are currently underway in our laboratory. If engraftment of porcine islets takes place in non immune suppressed macaques rendered tolerant by prior transplantation of embryonic pig pancreas, it is likely that comparable engraftment will occur in humans with diabetes mellitus. The ability to employ porcine islet transplants to normalize glucose tolerance in non-immune suppressed patients would widen the applicability for and reduce the toxicity of transplantation therapy for diabetes mellitus.

We have employed embryonic and adult endocrine pancreas to demonstrate a phenomenon, the immunologic mechanism for which remains undefined, that we term organogenetic tolerance. Induction of organogenetic tolerance for endocrine pancreas is critically dependent on the methodology employed to transplant pig pancreatic primordia,3,10-13,18,19 The phenomenon may be applicable to other organs with appropriate modifications in techniques used for implantation. For example, adaptations in methodology used to xenotransplant embryonic pig kidneys1 so as to result in engraftment that recapitulates the pattern observed following xenotransplantation of embryonic pancreas (i.e., renal cells engrafted in mesenteric lymph nodes) could render hosts tolerant to the embryonic kidney tissue and also to cells originating from adult kidneys from the same donor species. Successful induction of organogenetic tolerance to pig organs other than endocrine pancreas would increase the donor pool and, by virtue of eliminating the need for immune suppression, reduce the risk of transplantation so as to revolutionize organ replacement therapy.

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