Partial Pancreatectomy in the Rat and Subsequent Defect in Glucose-induced Insulin Release

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Abstract To define the consequences of a known reduction of B cell mass in rats, 90% partial pancreatectomies were performed. For the 6 wk following surgery moderate hyperglycemia was maintained in the fed state but there were no differences in body weight nor plasma insulin concentrations compared with sham-pancreatectomized controls. 8–10 wk following surgery regeneration of the remnant was evident with remnant weight being 26%, B cell mass being 42%, and non-B cell mass being 47% of values found for control whole pancreas. There were comparable increases in the remnant content of insulin, glucagon, and somatostatin. Following meal challenges, intraperitoneal and intravenous glucose tolerance tests and intravenous arginine challenge given 6–7 wk after surgery, the insulin responses to glucose were blunted or absent but the responses following the meals or arginine were intact. Similarly, when the pancreatic remnant was perfused in vitro, insulin release after challenge with 300 mg/dl glucose was markedly reduced whereas intact responsiveness to 10 mM arginine was retained. These data suggest that the chronic stimulation of a reduced B cell mass can lead to a selective loss of glucose-induced insulin secretion.

Introduction

Noninsulin-dependent diabetes mellitus (NIDDM) in man is characterized by a defect in glucose-stimulated insulin release and retained sensitivity to such secretagogues as arginine, isoproterenol, and secretin (1–3). Recently we described a rat model produced by a neonatal injection of the B cell toxin streptozotocin, which was found to have a similar abnormality of insulin secretion (4, 5). It was hypothesized that this lesion may have resulted from chronic stimulation of a reduced B cell mass or from a long-term effect of streptozotocin. To explore the consequences of a reduction of B cell mass exclusive of the effects of streptozotocin, the classic model of partial pancreatectomy (6–10) was studied using the newer methods of radioimmunoassay (RIA), organ perfusion, immunochemistry, and quantitative morphometry.

Methods

Partial pancreatectomy. 4–5-wk-old male Sprague-Dawley rats (95–125 g) were anesthetized with sodium amytal and 85–90% of their pancreas was removed by the technique of Folgia (8). The remnant (residual pancreas) was anatomically well defined, being the tissue within 2 mm of the common bile duct and extending from the duct to the first part of the duodenum. This remnant is the upper portion of the head of the pancreas thought to be embryologically of dorsal anlagen origin. Control animals were laparotomized and received a sham pancreatectomy that consisted of disengaging the pancreas from the mesentery and gently rubbing it between the fingers. Following surgery the animals were allowed to feed ad lib. on standard laboratory chow.

For the first week postoperatively body weight and plasma glucose values were followed daily. Then these values as well as the plasma insulin concentrations were followed at weekly intervals for 6 wk. Blood was collected in a heparinized glass tube by snipping the tail of a fed restrained rat and then centrifuged. For glucose measurements, 10 μl plasma was withdrawn, diluted with saline, and assayed using a Beckman Glucose Analyzer II (Beckman Instruments, Inc., Fullerton, CA). Pending assay for insulin, the plasma was frozen at −20°C.

Tissue extraction. 8–10 wk following surgery, pancreases were dissected from decapitated rats, cleared of gut and major lymph nodes, rinsed in saline, blotted, weighed, and kept on ice until homogenization. Pancreases from sham
animals were divided into the remnant equivalent (that portion of the pancreas with the same anatomical boundaries as the remnant left in the partial pancreatectomized animals as described above) and the rest. Data are presented for the remnant equivalent and the whole pancreas, which is the sum of the two portions. The tissue was homogenized in cold acid ethanol using an Ultra-Turrax SDT homogenizer (Tekmar Co., Cincinnati, OH). The samples were heated for 5 min in a 70°C water bath and then were adjusted in volume to 7 ml for the major pancreatic portion of sham animals and 3 ml for either the pancreatectomy remnant or the remnant equivalent and were stored at −20°C until assayed.

Quantitative morphometry. Pancreases after perfusion (see below) were dissected, cleared of lymph nodes, fixed in Bouin's fixative, and embedded in paraffin. Adjacent sections of blocks of either remnant or remnant equivalent were stained by immunoperoxidase (4, 11) for either B cells using guinea pig anti-porcine insulin antibody (the gift of Dr. Peter Wright, Indiana University Medical School, lot 607/27) or for the non-B cells using a slurry of rabbit antisyntetic somatostatin (our D-10), rabbit anti-bovine pancreatic polypeptide (the gift of Dr. R. Chance, Eli Lilly & Co., Indianapolis, IN) and rabbit anti-porcine glucagon (the gift of Dr. M. Appel, University of Massachusetts Medical School). Control incubations using excess antigen added to immune sera, nonimmune sera as primary antibody, and omission of the primary antibody resulted in the absence of hormone-specific staining. The relative volumes of B and non-B cells were quantitated by the point counting method of Weibel (12).

At a magnification of ×400 starting at a random point at one corner of the section, every other field was scored using a 25-point ocular grid. An unbiased but systematic selection of fields was easily accomplished using the markings of the stage micrometer. Intercepts over blood vessels, fat, ducts, lymph nodes, or interlobular space were substracted to give the total pancreatic counts. The mean total counts for the pancreatectomized animals (mean = 7.263±640, n = 7) reflected the 1.7-fold greater amount of tissue than in the remnant equivalent (mean = 4.686±673, n = 4).

In vivo challenges. 6 wk after pancreatectomy, animals were given different in vivo challenges. Some were given a meal challenge followed 6 d later by an intraperitoneal glucose tolerance test (IPGTT); others had an IPGTT followed 4 d later by an intravenous glucose tolerance test (IVGTT) and 4 d later by an intravenous arginine (IV ARG) challenge. For the meal challenge, animals were fasted 19 h, bled at 0 time, given food, and then bled at 30 and 60 min. All animals began eating within 10 min of receiving food. For the IPGTT, fasted (19 h) animals were bled at 0 time, injected intraperitoneally with enough 20% glucose solution to give a glucose load of 2 g/kg body wt and then bled at 10, 30, and 50 min. 2–3 d before the intravenous challenges (13, 14) indwelling jugular catheters were put in place. Animals were fed but food was removed 1 h before the intravenous challenges. For the IVGTT, enough 60% glucose solution to give 1 g/kg was injected into the cannula, which was then flushed with 1 ml of heparinized saline. Samples were collected over the next 60 min. The same animals 4 d later were given in the same manner enough arginine hydrochloride solution (180 mg/ml, pH 7.4) to give a dose of 300 mg/kg arginine and samples were taken at 0, 2, 5, and 15 min.

In vitro challenge: pancreas perfusion. Insulin secretion was studied in three groups of fed rats 8–11 wk following surgery using the in situ isolated perfused pancreas technique (10), with the same surgical approach used in each. Two of the groups included rats that had received either pancreatectomies or sham pancreatectomies, as has been described herein. The third group included rats of the same body weight in which perfusate flow to the pancreas was acutely restricted to the remnant area defined above. This restriction was accomplished with additional surgical ties and verified by blanching of the area once the perfusion was initiated. In all perfusions there was a 20-min equilibration period that followed the end of surgery and during this period the perfusate glucose concentration was 120 mg/dl. The flow rate for rats with pancreatectomies averaged 2.2 ml/min, for rats with sham operations was 2.7 ml/min, and for the group with acute restrictive surgery was 2.0 ml/min. Each group received the same protocol, which consisted of a base-line perfusate glucose concentration of 120 mg/dl and then 10-min challenges with first glucose at a concentration of 300 mg/dl, which was infused via a side-arm syringe, and then arginine (10 mM), delivered from a second reservoir. The exact timing of these perfusions is depicted in Fig. 3. Each sample was collected over a 60-s time period and Fig. 3 indicates the only samples that were obtained. Total insulin output over a given period of time was calculated by multiplying concentration of hormone in the perfusate by flow rate. In those time periods in which no sample was taken, values were calculated by extrapolation.

RIA. Insulin in the perfusate and extracts was measured by RIA using charcoal separation (16), with rat insulin (Novo Research Institute, Copenhagen, Denmark) used for standards. Glucagon was measured by RIA (15) using porcine glucagon as standard and antiseraum 30K, the latter purchased from Dr. Roger H. Unger, University of Texas, southwestern Medical School. Somatostatin was measured by a modification (17) of the method of Patel and Reichlin (18) using our antisemur D-6. Plasma insulin was measured with a double-antibody RIA (19).

Statistics. Results are presented as mean±standard error of the mean. Two-tailed unpaired Student's t test was used throughout this study.

**RESULTS**

Quantification of the extent of pancreatectomy. The remnant of the pancreas was anatomically defined as described above. At the time of pancreatectomy (4–5 wk of age) the remnant was 12.1±0.5% of the total pancreatic weight and contained 9.7±1.0% of the total insulin content (n = 6, data not shown). 8–10 wk after surgery (12–14 wk of age) the same portion of the sham-pancreatectomized controls, called the remnant equivalent, was 10.5±1.0% of the total pancreatic weight and had 10.3±0.9% of the total pancreatic insulin content (n = 6; Table 1).

Longitudinal study after partial pancreatectomy. Animals bled daily for the first week after pancreatectomy showed significant increases in the fed plasma glucose concentrations by day 4 (Fig. 1). This moderate hyperglycemia was maintained for the following 6 wk as shown in Fig. 2. During this time period there were no significant differences in body weight nor in plasma insulin concentration. Although a fed plasma glucose >200 mg/dl was occasionally found for any animal at any weekly bleed, only one animal (of 23) so followed had values consistently this high; data from this animal was not included in the results.

Hormone content and tissue weight. The hormone

**Insulin Secretion Following Pancreatectomy**

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TABLE I

Hormone Content of Remnant from Partially Pancreatectomized (Px) Rats, and of Remnant Equivalent and of Whole Pancreas from Sham-pancreatectomized (Sham Px) Rats

<table>
<thead>
<tr>
<th>Animals (n)</th>
<th>Tissue weight</th>
<th>Insulin Content</th>
<th>Glucagon Content</th>
<th>Somatostatin Content</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>g</td>
<td>µg</td>
<td>µg/g</td>
<td>µg</td>
</tr>
<tr>
<td>Px remnant (8)</td>
<td>0.303±0.036</td>
<td>20.7±2.6</td>
<td>71.4±7.85</td>
<td>1.77±0.14</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sham Px (6)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Remnant equivalent</td>
<td>0.125±0.012</td>
<td>6.40±1.30</td>
<td>54.4±12.8</td>
<td>0.416±0.047</td>
</tr>
<tr>
<td>Whole pancreas</td>
<td>1.190±0.070</td>
<td>64.0±11.7</td>
<td>54.6±9.6</td>
<td>5.83±0.62</td>
</tr>
</tbody>
</table>

* Pancreatic specimens were excised 8–10 wk after surgery (12–14 wk of age). The remnant equivalent is that portion of the pancreas contained roughly within the anatomical boundaries of the bile duct and duodenum as defined in Methods.
† Differs from Px remnant (2P < 0.001) and from whole pancreas (2P < 0.05).

content and concentrations of insulin, glucagon, and somatostatin for the pancreatectomy remnant, the remnant equivalent, and the whole sham pancreatectomized pancreas 8–10 wk following surgery are presented in Table I. At this time the remnant equivalent was still ~10% of the pancreatic weight and had 10% of the content of insulin and somatostatin. However, the remnant of the pancreatectomized animals showed increased growth and was 26% of the total pancreatic weight with 32% of the total insulin content. It is of interest that with the enhanced growth of the pancreatectomy remnant the hormone concentration of all three hormones is maintained at the level of the whole pancreas. It should be noted that the glucagon concentration of the remnant, equivalent was significantly lower than that of the whole pancreas.

FED PLASMA GLUCOSE VALUES IN FIRST DAYS AFTER PANCREATECTOMY (Px)

FIGURE 1 Effect of 90% partial pancreatectomy (Px) on the fed plasma glucose concentration during the first week following surgery. Px (------), n = 14; sham (------), n = 8. *2P ≤ 0.05.

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Morphology and morphometrics. The islet tissue in the pancreatocctomy remnant 10 wk after surgery was more heterogenous than controls when examined on immunoperoxidase-stained sections. There was great variation among animals but there were four distinct configurations found in most animals (Fig. 3): (a) normal appearing islets, which are the predominate population; (b) grossly larger islets with a seemingly lowered ratio of non-B-to-B cells; (c) small islets with extremely few non-B cells; and (d) fibrotic islets that show a disorganization of the usual mantle-core configuration. The fibrosis was restricted to the islet and perinsular region; there was no fibrosis between the exocrine acini.

Morphometric quantification dispelled the subjective impression that there was a change in the non-B-to-B cell ratio. The relative volumes (Table II) showed the same non-B-to-B cell ratio (remnant: 0.787±0.102% B cell/0.195±0.030% non-B cell = 4.04, n = 7 vs. remnant equivalent: 0.462±0.038% B cells/0.103±0.014% non-B cells = 4.07, n = 4). However, when the mass of B and non-B cells was calculated (with the assumption that the volume is equivalent to mass, i.e., 1 ml = 1 g tissue), it was clear that there had been considerable proliferation of B and non-B cells after the partial pancreatectomy. The mass of both the B and the non-B cells were fourfold greater than those of the remnant equivalent of the same aged animal. To look at this another way, making the assumption that the remnant equivalent is truly representative of the whole pancreas, then the mass of B and non-B cells in the whole pancreas can be calculated (Table II). The comparison of these values with those of the pancreatectomy remnant, showed that 8–10 wk after partial pancreatectomy the remnant had 42% of the total B cell mass and 47% of the total non-B cell mass. Thus, the reduction of islet tissue did not remain at the 90% level of the pancreatectomy. Further calculations from these data give values for insulin content per milligram B cell tissue (Table II); there was no significant difference in this concentration even though there was a tendency for the remnant values to be decreased.

In vivo challenges. Animals studied 6–7 wk after partial pancreatectomy had normal fasting glucose and insulin values but were hyperglycemic after either a
meal challenge or an IPGTT (Table 3). 30 min after
the meal, the plasma insulin concentrations increased
to a comparable degree in both the pancreatectomized
and the sham-pancreatectomized (control) animals.
However, in response to intraperitoneal administered
glucose, the pancreatectomized animals did not show
an increase in plasma insulin values whereas the con-
trols did (Table III).

The data for the IVGTT and the IV ARG challenge,
as presented in Table IV, show that the pancreatec-
tomized animals have a dulled insulin response to glu-
cose and a relatively better response to arginine. The
difference in the insulin responses to these intravenous
challenges is emphasized by comparing the increase
above base line at 2 min in both groups. Taking into
consideration the finding that the B cell mass of the
remnant in 42% of the sham control whole pancreas
(Table II, which contains data from a comparable
group of animals), it is apparent that the arginine re-
sponses are comparable in the two groups of rats, but
that the responses to glucose are very different.

In vitro challenge with pancreatic perfusion. When
the whole pancreases from the sham control rats were
challenged with glucose (300 mg/dl), the expected
biphasic response occurred; a clear response to argi-
nine (10 mM) was seen as well (Fig. 4, Table V). In
contrast, the pancreatic remnants from the rats that
had received a partial pancreatectomy had a markedly
impaired response to the high glucose challenge, but
retained their ability to respond to arginine. As an
additional control, the same challenges were given to
seven pancreases in which perfusate flow was re-
stricted to the remnant area. The responses, when ex-
pressed as percent rise from base line, were compa-
rable to the results found with the whole pancreas.
Thus the mean percent increase in response to high
glucose for the whole pancreas was 612±101%, for the
restricted remnant area was 386±57%, (not significant
as compared to whole pancreas) and for the actual
remnant was 159±21% (2P ≤ 0.001 as compared with
whole pancreas). The responses to arginine were for
the whole pancreas 1,316±355%, for the restricted
remnant 1,259±215%, and for the actual remnant
1,462±482%.

It was of interest to compare the insulin responses
of the remnant and sham-pancreatectomized whole

\[ \frac{1548}{S.\; Bonner-Weir,\; D.\; F.\; Trent,\; and\; G.\; C.\; Weir} \]
TABLE II
B Cell and Non-B Cell Volumes and Insulin Concentration (as Function of B Cell Mass) in Remnant from Partially Pancreatectomised (Px) Rats, and in Remnant Equivalent and in Whole Pancreas from Sham-pancreatectomised (Sham Px) Rats

<table>
<thead>
<tr>
<th>Animal (n)</th>
<th>Relative volume*</th>
<th>Mass</th>
<th>Insulin content per unit of B cell mass</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>B cell</td>
<td>Non-B cell</td>
<td>B cell</td>
</tr>
<tr>
<td></td>
<td>%</td>
<td>mg</td>
<td>µg/mg</td>
</tr>
<tr>
<td>Px remnant (7)</td>
<td>0.78±0.102</td>
<td>0.195±0.030</td>
<td>2.33±0.25</td>
</tr>
<tr>
<td>Sham Px (4)</td>
<td>0.462±0.038</td>
<td>0.103±0.014</td>
<td>0.576±0.555</td>
</tr>
<tr>
<td>Whole pancreas§</td>
<td>—</td>
<td>—</td>
<td>5.50±0.32</td>
</tr>
</tbody>
</table>

* Using the Weibel (12) point counting method on immunoperoxidase-stained sections, the relative volume was determined by dividing the number of intercepts over specifically stained cells by the number of total intercepts over pancreatic tissue.

† With the assumption that volume is the equivalent of the mass of a tissue, i.e., 1 ml = 1 g tissue, the mass is determined by multiplying the relative volume by the mean pancreatic weight as given in Table I.

§ With the assumption that the remnant equivalent is representative of the whole pancreas, one can make calculations for the whole pancreas for comparisons.

pancreas in terms of either B cell mass or insulin content (Table V). In both cases the insulin release elicited by 120 mg/dl glucose or 10 mM arginine was comparable, but the response by the remnant to 300 mg/dl glucose was clearly reduced. The secretion data for the whole pancreas experiments were used for these calculations instead of those from the acutely restricted remnants, because the latter were presumed to have more variability from acute surgical trauma.

DISCUSSION

Partial pancreatectomy resulted in discernible regeneration of the remnant and additionally in defective glucose-induced insulin release, which was shown by both in vivo and in vitro techniques. These findings add further weight to the hypothesis that excessive glucose stimulation of a reduced B cell mass leads to functional abnormalities of the B cell. Since similar

TABLE III
Meal Challenge and IPGTT*

<table>
<thead>
<tr>
<th>Animal (n)</th>
<th>Time (min)</th>
<th>Meal challenge</th>
<th>IPGTT</th>
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<tbody>
<tr>
<td></td>
<td>0</td>
<td>30</td>
<td>60</td>
</tr>
<tr>
<td>Pancreatectomy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Plasma glucose (mg/dl)</td>
<td>96±5</td>
<td>184±6†</td>
<td>181±7†</td>
</tr>
<tr>
<td>(n)</td>
<td>(14)</td>
<td>(14)</td>
<td>(14)</td>
</tr>
<tr>
<td>Plasma insulin (ng/ml)</td>
<td>0.34±0.03</td>
<td>1.0±0.12</td>
<td>—</td>
</tr>
<tr>
<td>(n)</td>
<td>(7)</td>
<td>(14)</td>
<td></td>
</tr>
<tr>
<td>Sham pancreatectomy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Plasma glucose (mg/dl)</td>
<td>87±4</td>
<td>136±9</td>
<td>137±5</td>
</tr>
<tr>
<td>(n)</td>
<td>(6)</td>
<td>(6)</td>
<td>(6)</td>
</tr>
<tr>
<td>Plasma insulin (ng/ml)</td>
<td>0.58±0.16</td>
<td>1.42±0.34</td>
<td>—</td>
</tr>
<tr>
<td>(n)</td>
<td>(4)</td>
<td>(6)</td>
<td></td>
</tr>
</tbody>
</table>

* Animals were fasted (19 h) and given the meal challenge that consisted of receiving standard laboratory chow. 6 d later the same animals were fasted (19 h) and given an IPGTT (2 g/kg glucose).
† 2P ≤ 0.001, as compared with sham pancreatectomy.
Pancreatectomy
Plasma glucose (mg/dl) 171±4\$ 148±20 403±20 315±14 211±15 (n) (10) (10) (10) (10) (10)
Plasma insulin (ng/ml) 1.6±0.3 2.7±0.4 2.5±0.4 2.0±0.3 1.4±0.2 (n) (10) (10) (9) (10) (9)
Sham pancreatectomy
Plasma glucose (mg/dl) 120±7 422±23 330±25 237±11 112±7 (n) (10) (10) (10) (10) (10)
Plasma insulin (mg/dl) 2.0±0.4 11.7±0.2 7.4±1.1 4.7±0.9 1.3±0.2 (n) (10) (8) (9) (9) (9)

* 2-3 d before intravenous challenges, indwelling jugular catheters were put in place. Glucose (60% solution, 1 g/kg) was injected in the cannula of animals in the fed state and then flushed with heparinized saline for the IVGTT. 4 d later the same animals again in the fed state received arginine hydrochloride solution (180 mg/ml, pH 7.4, 300 mg/kg dose) also via the cannula.

† 2P < 0.05, as compared with sham pancreatectomy.
§ 2P < 0.005, as compared with sham pancreatectomy.

secretory patterns were seen in both the neonatal streptozotocin model (4, 5) and these partial pancreatectomy findings, the possibility that streptozotocin caused this acquired defect seems less likely. In the neonatal streptozotocin model the reduction in B cell insulin content could lead to the suggestion that a defect in insulin production or storage led to the secretory defect. However, in the partial pancreatectomies, the B cell insulin content of the remnant was normal. The reason for these differences in insulin content of the two models is unknown, but perhaps the higher plasma glucose levels of the streptozotocin model led to degranulation. This is consistent with the demonstration that islets cultured at high glucose concentrations have depleted insulin stores and reduced insulin responses to acute glucose challenges (20, 21).

Even though the pancreatectomized rats had a clear defect in glucose-induced insulin secretion they were only moderately hyperglycemic. In the fed state plasma insulin concentrations were indistinguishable from control values, but taking into account the hyperglycemia, it can be argued that there is relative hypoinsulinemia. Nonetheless, the fact that the fed insulin levels are as well maintained as they are can probably be attributed to such nonglucose B cell stimuli as amino acids and gut hormones. Indeed, the in vitro and in vivo insulin responses of the remnant to arginine are essentially normal when expressed as a function of B cell mass.

The partial pancreatectomy model (6-10) was a popular one several decades ago and the data from this work led to the often quoted concept that only 10% of the B cell mass is required to maintain a non-diabetic state. This is misleading for several reasons. First, there is regeneration of B cell tissue such that the remnant in the present study contained 42% of the normal mass of B cells. Second, even though the fed plasma glucose levels of ~160 mg/dl may seem unimpressive they are clearly higher than normal and unequivocal glucose intolerance is seen after either a glucose or meal challenge. Third, the partial pancreatectomy removes a substantial mass of glucagon-containing A cells, and this lack would be expected to further ameliorate the hyperglycemia. In a study by Martin and Lacy (10) 90% partial pancreatectomies on rats were also performed and 50% of the animals were clearly hyperglycemic 40 d following surgery. We cannot explain the increased severity of diabetes in their study, but differences in surgical technique or in the strain of the rats could be responsible for the discrepancy.

The regeneration of the remnant is of interest in that there were almost parallel increases in the mass of exocrine tissue, B cells and non-B cells. This was reflected by the concentrations of insulin, glucagon, and somatostatin in the remnant, equivalent to those found in whole control pancreas. The glucagon concentration of the remnant equivalent was lower than
that of either the remnant or the whole pancreas, and the reason for this difference is unclear. Certainly glucose stimulation to B cell growth (22) remains an attractive explanation for the increase of B cell mass, but the manner in which this might be related to the growth of the other elements remains unknown. The

![Graph showing insulin secretion in response to glucose and arginine from the perfused pancreatic remnant 8–11 wk after partial pancreatectomy.]

**Figure 4** Insulin secretion in response to glucose and arginine from the perfused pancreatic remnant 8–11 wk after partial pancreatectomy. Control rats (sham) were sham pancreatectomized 8–11 wk earlier. All animals were studied in the fed state. The data are expressed as percent change from base line, rather than as the absolute quantity of insulin released, to make the two groups easier to compare.

**Table V**

<table>
<thead>
<tr>
<th>Perfusate condition</th>
<th>Insulin output†</th>
<th>Insulin output/B cell mass</th>
<th>Insulin output/insulin content‡</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Sham Px</td>
<td>Px</td>
<td>Sham Px</td>
</tr>
<tr>
<td>Glucose, 120 mg/dl</td>
<td>6.64±2.28</td>
<td>2.27±0.76</td>
<td>1.21±0.41</td>
</tr>
<tr>
<td>Glucose, 300 mg/dl</td>
<td>32.2±8.4</td>
<td>3.78±1.36‖</td>
<td>5.86±1.53</td>
</tr>
<tr>
<td>Arginine, 10 mM</td>
<td>52.1±8.8</td>
<td>16.5±2.87‖</td>
<td>9.48±1.61</td>
</tr>
</tbody>
</table>

* The number of perfusions for Px was nine and for Sham Px was seven.
† This refers to the mean output calculated from the initial 5-min period for 120 mg/dl glucose and the 10-min periods for 300 mg/dl glucose and 10 mM arginine.
‡ The output for 120 mg/dl although measured for only 5 min was calculated as if for 10 min to allow comparisons with the two other conditions.
‖ Differs from Sham Px, 2P < 0.02.
heterogeneity of islet configurations suggests that there may be replication of existing B cells in islets, but also possible neogenesis of complete islets. The pathogenesis of the fibrosis seen in some islets remains a puzzle, but we find no evidence indicating that it can be correlated with the degree of glucose intolerance as has been suggested by Clark et al. (23). Earlier studies have suggested that partial pancreatectomy accompanied by injections of anterior pituitary extract can lead to islet damage and “exhaustion” (7). Even though anterior pituitary extract was not used in this study, we found no evidence of islet destruction, and it does not yet seem reasonable to view the fibrosis in those terms. If the term exhaustion is used in a carefully defined restricted sense, one could, however, argue that there is a selective exhaustion of glucose-induced insulin secretion.

The results of the present study could have implications for our understanding of some of the abnormalities which are seen in NIDDM in man. The available morphological evidence suggests that B cell mass in NIDDM is reduced to ~60% of normal (24, 25). One can therefore speculate that this B cell mass is stressed by hyperglycemia in the same manner as occurs in the partial pancreatectomy model, and that this leads to the selective defect in glucose-induced insulin release seen in NIDDM (26). Further support for this hypothesis comes from several studies which show that control of plasma glucose levels in NIDDM results in improved insulin secretion (27–29). There is emerging evidence for an abnormal structure of the 5’-flanking region of the insulin gene in NIDDM (30, 31) and the relationship of this to the reduced B cell mass and the abnormalities of insulin secretion in this disease remains to be unraveled.

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