Has Faetal Whole Faetal Pancreas Grafting Some Perspectives among the Surgical Methods of Diabetes Mellitus Treatment?

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Abstract

Background: Presently the best results in diabetes mellitus treatment are obtained after pancreas transplantation and bariatric surgery which have not negligible consequences. Implantation of pancreatic adult beta cells, islets, stem cells up to now has only temporary effect.

Material and methods: In experiments on 148 rats (including control series) Streptozotocin® diabetes mellitus was induced and followed by subcutaneous syngeneic foetal whole pancreas syngeneic implantation. In clinics, between 1982 and 1990, 59 patients with type 1(53) and insulin requiring type 2(6) diabetes mellitus have undergone allogeneic implantation of foetal pancreas without immunosuppressive drug use after obtaining their and the foetus' parents' informed consent, as well as the agreement of the USSR Ministry of Health Care.

Results: Long lasting correction of Streptozotocin® induced diabetes was obtained in 30% of the rats with syngeneic implantation of foetal pancreas. The best results - in 7 out of 10 animal series were observed when 4 and more foetal pancreas was implanted. In clinics, complete healing of the patients could not be observed, but significant and long lasting (up to 2-10 years, 70% of the cases) improvement of general condition including remission of retinopathy and neuropathy, decrease of glycaemia levels and of the daily delivered insulin amount accompanied by an increase of blood C-peptide were noted in 70% of the humans concerned.

Discussion and conclusion: The positive aspects and difficulties of foetal pancreas implantation are analyzed comparing this experience with literature data. Perspectives of adaptation to contemporary techniques and ethic conditions are suggested.

Keywords: Diabetes mellitus surgical treatment; Pancreas transplantation; Foetal pancreas; Foetal pancreas implantation; Beta cells implantation; Pancreas stem cells endocrine pancreas regeneration

Abbreviations: APD: Animal Protein Deprivation; APDD: Animal Protein Deprivation Diabetes; DM: Diabetes Mellitus; DM1: Type 1 Diabetes Mellitus; DM2: Type 2 Diabetes Mellitus; EEG: Electroencephalogram; FPI: Foetal Pancreas Implantation; Ptx: Adult Pancreas Transplantation

Case Report

In spite of the remarkable development of drugs (insulin’s with various intensity and duration of action; oral anti-diabetic preparations of different classes) and devices for their rationale administration up to artificial pancreas [1-18], conservative treatment of diabetes mellitus type 1 as well as type 2, only can significantly improve the condition of the patients and prolong their life, diminishing the complications of the disease, but does not completely cure it. Only surgical approach allows a total and definite healing in most cases pancreas Transplantation for DM1 and bariatric surgery, especially by-pass for DM2 patients [19-26]. However these operations are complex not easily tolerated and followed by severe consequences: immunosuppressive treatment with its infectious and oncological complications on one side, metabolic disorders of nutriments and vitamin absorption/assimilation on the other one [20,25,26].

It is the reason why since decades another solution to the problem has been considered: a kind of reduced and targeted transplantation of the endocrine part - cells or islet - implantation: acceptable for treatment of the not too severe DM [27-38]. The first results were promising; but the effect duration was limited; collecting, storage and immunologic (if allo- or xenogeneic) remain. In the 80ties foetal pancreas implantation was studied in animal experiments and even in clinics [39-42]. The results of these studies have shown that foetal implants are able to survive
and develop in adult recipient and even to function. Because of the
great 30-year moratorium of the researches on embryonic
and foetal material [43-44], this way was practically abandoned.

New trials were started with pancreatic stem cells, first
cultivated in vitro and later implanted into the patient organism
[45-46]. But cultivated cells have great difficulties to adapt, develop and work in the receptor organism. The aim of the
present work is to come back to our own investigations [40,42,47-
55], to analyse their results taking into account more recently
obtained experimental data, to confront our conclusions with
literature and answer to the question: has the foetal pancreas
grafting a place, a perspective in the DM surgical treatment, at
what conditions?

Material and Methods

In this work we have put together for a new analysis animal
experiences (part 1) and clinical trial (part 2). Both parts have
been the subject of separate and partial publications, to which
we refer when necessary.

Part 1

Experiments were carried on 148 rats (Fischer, Lewin and
Wistar, males and females, aged 2-4 months at the beginning of
the observation).

They were dispatched into the following series:

a. Control 1 -5 sham operations - creating a site for
implantation without performing the FPI (NB. data obtained
during several day observation before the STZ injection were
also used as control).

b. Control 2 - FPI in healthy animals -10 animals.

c. Streptozotocin diabetes - total 59 - without treatment
25 with implantation of foetal pancreas at days 1, 4, 7, 12, 42
after the STZ injection: 34 Animal protein deprivation and foetal
pancreas implantation.

d. -75 Streptozotocin (75 or 100mg/kg) was injected intra
peritoneum. In experimental series the injection was provided 4
days before syngeneic implantation of foetal pancreas (FPI).

Animal protein deprivation (APD) consisted in exclusion
from the food any animal protein (milk, cheese, meat) during the
female gravidity and the first 4-6 weeks after birth. Drink water
and other food such as vegetables and corns were delivered ad
libido. At this moment animal protein deprivation was stopped,
usual diet was restored and implantation of allogeneic foetal
pancreas was provided, in order to imitate the possible human
situation in cases of chronic hunger [40] (see also “Diabetes
Mellitus” Report of the WHO research group- 1975). NB. For
STZ diabetes treatment Syngeneic grafting was adopted in
order to test the graft potential capacities without the bias of
biological incompatibility. In all the series, donors were rat
foetuses collected after 14-18 days of intrauterine development.
Recipients were adult rats with a body weight of 200±10g
(female rats) or 250±30g (male rats). All the operations were
performed under general anesthesia (0.1ml of 0.015% Nembutal
solution/100g BW injected intra peritoneum).

For implant collect, after extracting a foetus from the
mother’s womb the foetal organ was removed and placed in a
saline solution. The implantation site was: ear (104 rats), spleen
hilum (14 rats). In the first experimental group, the foetal tissue
(1 or 2 foetal pancreas were introduced into a subcutaneous
pouch of both ear pavilions of the recipient. The skin wound
was closed with “No becutane®” spray. In the second group,
laparotomy was performed and 3-8 pancreases were introduced
under the peritoneum and the spleen capsule. Abdominal wound
was closed by 2-layer running suture with Ethilon 4 °C (Figure
1). After the operation, the recipient was placed in a separate
standard cage and fed with a standard maintenance diet in an
agreed animal husbandry. Some of them received 1 or more IU
of Actrapid® subcutaneously to maintain glycaemia within 250
and 350mg/dl.

The following investigations were provided: BW, amount
of daily drunk water, measure of capillary glycaemia and of
glycosuria, proteinuria, ketonuria, haematuria and pH by
strips (“Acutrends”, USA). In some cases IGF 1 was determined
in blood serum. Observation delays have run up to 9 months.
Post-operative animals were euthanized by anesthetic overdose
at varying time points (days 2, 4, 7, 14, 58, 80, 120, 397) for
morphological investigations of the implant and main organs of
the recipient. The material was fixed in 12% formaldehyde,
and paraffin inclusion was performed. Subsequently, embedded
tissue slices 3-4 microns in thickness were stained with
hematoxylin eosin, in some cases determination of insulin,
glucagon and somatostatin by immune histochemistry methods,
and electron microscopy was performed after glutaraldehyde
fixation and araldite embedding. All experiments were conducted
in conformity the Helsinki Bioethics Convention and recent
experiments were approved by the Brussels Free University
Ethics Committee. (Protocol 50)

Part 2

The clinical study was performed in Moscow Hospital N
64 at the Gynecology Unit under the supervision of the head
and medical staff of this department, with the agreement of
the USRR Public Health Ministry from 1980 till1990 years.
Pregnant women, exempt of local infection, tumor, and infectious diseases, with spontaneous or induced for medical reasons early delivery of not viable foetus (aged 14-22 weeks, BW<500g), both sexes), were informed and gave their consent to the foetal pancreas collecting. In sterile conditions of operation room the donor’s laparotomy was performed and the pancreas isolated and placed in a small amount (1-2ml) of conserving solution used for tissue culture, minced with scissors (whole pancreas dimensions: 2x2x20 to 4x5x25mm, fragment volumes: 1-2mm³). After operation the donor’s wound was closed and the foetus returned to the family or to the hospital service for engraving. Recipients were patients with compensated (at the moment of the operation) but severe or middle complicated type 1 diabetes (53) and type 2 insulin requiring DM(6) aged 16-49 years, having given their informed consent. During to the implant collection, the recipient was prepared, abdominal skin disinfected. In the Para umbilical region a small incision of 0.5cm was performed and the foetal material was injected through a special trocar (diameter 2.5mm) into the subcutaneous fat at a depth of 2-3cm. This site was supposed to ensure a venous drainage partly in direction of the portal vein through the Para umbilical venous web. One silk ditch closed the wound.

After operation usual investigations were provided: blood glucose, urine glucose and acetone, blood creatinine, urea and other routine indices determination [42]. In some cases blood c-peptide and insulin were determined, as well as some immunologic investigations (CD4, CD8, global amount of Theophylline sensible lymphocytes and active lymphocytes). The follow up was realized by family polyclinics physicians and under the control by endocrinologists and by our team. Only 3 patients have systematically received higher doses of insulin during the first post operation 3 days. For the others insulin was given according to the usual schema of glycaemia correction needs. Ultrasound investigation of the site of FPI and abdomen was performed to follow the morphological evolution of the implant in 16 patients. EEG was also performed in patients with severe neuropathy and encephalopathy.

The observation delay varied from 6 months to 10 years and more. Repeated implantation were performed 6 times according to the patients’ requests. Biopsies of the implant were not performed.*NB. Since 1990 the study has been interrupted: the initial one (5x3x2mm versus 2x1x1.5mm) the typical structure of adult pancreas with well-structured exocrine and endocrine parts were never observed. Nevertheless after 4 months a pancreas-like formation was observed with tubules, endocrine cells positive to insulin, glucagon and somatostatin (Figure 2B). These endocrine cells might be functional, that was verified in series with STZ diabetes.

In control healthy rats glucose blood levels varied between 80 and 130mg/dl and daily drunk water volume was between 20 and 40 ml; urine was free of blood, protein, glucose, urea and pH was 5-6. We considered as normalized treated diabetic animals whose glycaemia had come back to 100±20mg/dl, daily drunk water amount - to 30±6ml polyuria, urine ketone have disappeared, and BW has reached normal level according to the strain evolution curve depending on age and sex. After STZ injection without any treatment the mean survival delay was 12±7 days, glycaemia reached 500 and more mg/dl with polyuria, high glycosuria, presence of ketone in urine, severe decrease of BW and general condition worsening.

FPI has given positive results in 8 of the 24 animals concerned (only 30%), that was complete and stable during several months, normalization of glycaemia, disappearance of glycosuria and improvement of general condition including BW increase and normalization (compared to standard BW curves by strain, sex and age). A whole series of 16 STZ rats has failed

Results

Part 1

Syngeneic FPI at the ear site has allowed the following observations: during the first week, complete destruction of the organ structure: only weakly differentiated cells, fibroblasts and necrotic or apoptotic cells may be observed (Figure 2A). At the end of the first post implantation week some lacunas and capillaries appeared and the pancreas development has started (Figure 2A). But though the volume of the new-formed organoids was much more the initial one (5x3x2mm versus 2x1x1.5mm) the typical structure of adult pancreas with well-structured exocrine and endocrine parts were never observed. Nevertheless after 4 months a pancreas-like formation was observed with tubules, endocrine cells positive to insulin, glucagon and somatostatin (Figure 2B). These endocrine cells might be functional, that was verified in series with STZ diabetes.

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after FPI of 1 or 2 pancreases and no other intervention. The best results were obtained when glycaemia level during the first days and weeks after grafting was kept between 250 and 350mg/dl by insulin injection of 2-4U1 daily at the beginning, and then slowly decrease in dose and injection frequency.

Insulin supplement was stopped when glycaemia has fallen under the end point 250mg/dl (Figure 3). The result of APD was a slowing of the development: BW gain, sexual differentiation and in 50.1% of the animals the development of diabetes some months after the nutrition normalization. The allogeneic FPI has hastened the correction of the animal development after restoration of a normal diet and significantly decreased the incidence of post APD diabetes (Table 1). Interesting that the allogeneic implant acute rejection was observed only at the end of the restoration process, i.e. about 1 month after FPI, when animals have practically reached their normal BW.

### Table 1: Results of Foetal pancreas implantation in protein deprived rats.

<table>
<thead>
<tr>
<th>Series</th>
<th>APD Rats Number</th>
<th>APDD Cases Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>FPI</td>
<td>38</td>
<td>5(13.1%)</td>
</tr>
<tr>
<td>Control</td>
<td>37</td>
<td>19(50.1%)</td>
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Ultrasound investigation has shown a significant increasing of the implant dimensions, especially in patients with a type 2 effect and a fibrous reduction in late delays. A temporary increase of the patient’s own pancreas dimensions was also noted. Encephalography has shown normalization of the brain electric activity in 15 out of 16 investigated patients. As to hormonal investigation results increase of blood C-peptide (0.4- 4ng/ml) in 10 out of 20 investigated patient. An increase of blood somatomedin (from 52±10 versus 91±20ng/ml) and IGF-1, normalization to control levels of increased or decreased glucagon were also observed in isolated patients investigated during the first year after FPI. Results of blood lymphocytes investigation have shown some changes correlated in time to the FPI effect, but not significant enough.

According to the results of the inquiries, different improvements in the disease development maintained during 4-7 years. Stability of the DM was noted during 4-10 years after FPI in 74±1, 3% of the patients (interval of liability 95%). The second and third FPI has a less expressed positive effect, including a shorter restoration of the patient’s potency. In order to diminish the hypoglycaemia and yo-yo events, Biostator and Apparatus of Diabetes Diagnosis and complex insulin treatment [7,12-15] were first used insulin delivery monitoring with positive stabilizing effect not only during the sessions but during several days after them.

### Discussion

Our results have confirmed that foetal pancreas grafting has proved to be possible, to give growth to adult organ-like formations with specific morphology and endocrine function [39,41] and others. Complete correction of STZ induced Diabetes in rats under certain conditions as well as significant improvement of diabetic patients’ condition and Indices during several years in 70% of the cases. Some remarks are necessary. Considering the syngeneic implant structure evolution in rats, we never have observed the development of an adult structured exocrine and endocrine pancreas. Both types of tissue co-existed but without the connexion they have in the foetal organ before grafting, or in the normal adult organ. This could be due to the implantation site - subcutaneous ear pouch - which did not ensure the place enough for a normal expansion and no possibility for evacuation/excretion of the secreted juice that could inhibited its secretion.

It could explain that other authors have described the restoration of both endo and exocrine parts of the implant placed under liver or spleen capsule [36]. Another explanation is the
absence in the pancreas of a well-organized intramural nervous system, the compact character of the gland. The same observations were obtained with foetal liver transplantation [56-57]. This makes the difference with the hollow organs of the digestive tract which give growth to a normal adult oesophagus, stomach or intestines [51-52] and with the heart [58-59]. However the very presence pancreatic tubules in the developed FPI can explain the long lasting effect of the graft in STZ rats and even in patients. The role of these formations is known in the endocrine cells production and endocrine pancreas natural regeneration. Hence the existence and function of FPI were found to be significantly more important that the one of differentiated endocrine cells or clusters implants. In rats, the success of FPI was conditioned by several factors such as: no less than 4 foetal pancreas implanted, warm ischaemia no more than 1 hour, a well vascularized site (ear pavilion, organ parenchyma - spleen or liver), regulation of glycaemia between 250 and 350 mg/dl by insulin injections, that is high enough to stimulate insulin production by the graft, but not too high, that can inhibit or paralyze it. We have also noted this inhibitory influence of maintaining high insulin doses after FPI on the early post operation evolution of the patients.

An important question is the FPI immunogenicity. In our experiments, the foetal material used was syngeneic that allowed to avoid graft rejection problems and be sure of the FPI potentiality. We have proved that allogeneic foetal organ graft is rejected as well as adult organ one [1994]. Maybe embryonic material is less antigenic [46], but we have not used it. It is also possible that DM1 patients are immunologically slightly modified as shown by altered ratio CD4/CD8 lymphocytes (not published own observations) and other literature data [60-63]. If true, this can explain why our FPI underwent no acute, but more or less quick chronic rejection.

Interesting that in APPD series a similar phenomenon could be observed: the implanted was tolerated by weak dystrophic rats, but the implant was acutely rejected as soon as the animal condition was improved. This may be important when considering the complex treatment of serious malnutrition. The possible origin of the so called ‘before 1975 “DM type 3” probably lays in the lack of protein for the adequate insulin synthesis corresponding to the needs of an adult organism. So, for DM treatment FPI has the following advantages:

a. No need to pass by an expensive culture in vitro, long lasting effect on the different aspects of the disease including its vascular, neurologic and other complications.

b. Organization: (the potential recipient has to be ready to answer the calling as soon as possible when a potential donor is considered).

c. Storage (in our cases the maximal warm storage between the donor’s death and the implantation at 19-20 °C was 60-90 min, more often less than 30 min, that was considered as a factor of success), tolerance problem between host and implant.

d. Ethic problems which can be overcome if considering the foetus as a child and, as for the children) donors, obtain the informed consent of the parents or the mother if no father declared.

Coming back to the development of FPI within the adult organism, our experimental observations has shown that the first post operation week presents a certain interest. During this period the implant loses its morphologic structure: part of already differentiated cells die (apoptosis, ischemic necrosis), part seems to dedifferentiate, part of weakly differentiated cells - precursor stem cells - survive.

The cause might be the diffusion conditions of “feeding” which end only after 5-7 days when host capillaries penetrate the graft and blood elements can be observed within it. It marks the beginning of re-differentiation of the implant elements which corresponds to the start of IGF1 elevation in the host blood. The process is the same in all the foetal organ implants we have observed [47, 55]. The phenomenon has not be systematically investigated as for the nature of these not differentiated cells, the presence of growth factors in this medium, the role of the host own stem cells and growth factors (base level or induced by FIU) in the implant development. Among the observed cells of the implant there are probably precursor cells specific for the grafted organ, as far as from a FPI only pancreas cells and tissue will grow, and the same is observed with other foetal organ implants. Teratoma growth is an exception [64]. So, may be, FPI and other foetal organ implantation constitutes a natural short cut for in vivo production of these organoids which are going to revolutionize the regenerative medicine? [65-68]

Conclusion

I. FPI for DM treatment remains promising but under certain conditions of collect, storage, material quantity implanted and host adequate management and follow up.

II. The presence of pancreatic tubules in the grown FPI is probably the warrant of beta-cell further production which allows long lasting effect of the procedure.

III. Implantation of fetal organs could also be applied to clinical purposes, the main problems to resolve being: a) development of acquired immunological tolerance rather than immune suppression and b) constitution of fetal tissue/organ banks with in vitro as well as in vivo storage.

IV. Syngeneic implantation of foetal organs with its features of regression followed by ontogenetic development seems to be an interesting model for the IN VIVO study of such phenomena as possible de-differentiation of tissues and managing re-differentiation growth factors.

V. The organ-like development of FPI (and other foetal organ grafting) could be an effective short cut for obtaining IN VIVO well-formed and vascularized organoids able to repair or replace altered adult organ.
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