DETECTION OF PANCREATIC ISLET TISSUE FOLLOWING ISLET ALLOTRANSPLANTATION IN MAN

Qualitative and quantitative standards have been recently proposed for islet isolation assessment in humans and large animals (1). The ultimate test of viability is functional activity after transplantation into a diabetic recipient. Morphologic identification of insulin-containing transplanted islets could provide important correlative data and eventually confirm the functional results. Recently, several centers reported prolonged insulin independence after human islet allotransplantation (2-5). The present report describes the first identification of islet tissue in a liver biopsy after intrahepatic islet allotransplantation.

**Figure 1.** Liver biopsy showing a well-preserved human islet (top) in a portal space 17 days following human islet allotransplantation. No significant surrounding inflammatory reaction was observed (H & E: original magnification x80).

**Figure 2.** Human islet in liver biopsy after immunohistochemical stain (immunoperoxidase) for insulin. Beta granulations (dark) can be distinguished in the cytoplasm of the majority of the islet cells. (original magnification: x350).

Human islets were obtained from 2 multiorgan donors by a modification (6) of the automated method for human islet isolation (7). The islets were purified on Eurocol-lins-Ficoll gradients (8) using a cell separator (9, 10) (COBE 2991, Lakewood, CO). On March 16, 1990 258,000 islets of an average diameter of 150 μ (1) were transplanted via portal vein injection following an upper abdominal exenteration and liver replacement (2). The procedure was performed in a 52-year-old woman for adenocarcinoma of the ampulla of Vater infiltrating the duodenal wall, with liver metastasis. Immunosuppression was with FK506 as described previously (2).

An exploratory laparotomy was performed 17 days after
transplantation to rule out intraabdominal abscess. During the procedure a wedge biopsy was taken from the liver. A well-preserved human islet in a portal space (Fig. 1), with no significant surrounding inflammatory reaction was demonstrated. Immunohistochemical stains (immunoperoxidase) for insulin and glucagon confirmed the presence of beta (Fig. 2) and alpha (Fig. 3) cells. The insulin stain unequivocally demonstrated beta granulation in the majority of the islet cells. The alpha cells appeared more intensely stained for glucagon. A possible reason for the difference in stain intensity could be that total parenteral nutrition was administered to the patient. The hyperalimentation may have induced relative beta cell degranulation. An additional explanation may be that she received a marginal number of islets that could have been maximally stimulated. In support of this, she was the only one out of 9 patients who was still requiring daily insulin injections 4 months after transplantation (2). Nevertheless, islet function eventually improved, resulting in insulin independence 6 months after human islet transplantation.

The morphologic documentation of intrahepatic pancreatic islet tissue following islet allotransplantation in man indicated that human islets can survive in the liver of transplanted patients, adding histologic evidence to previously reported functional results (2). A larger series of patients who received combined liver-islet or kidney-islet allografts is in progress (n=20).

This report provides the first evidence of islet tissue in a liver biopsy following human islet allotransplantation. These initial findings have been confirmed in other patients in whom islets have been found both in wedge and needle liver biopsies.

Camillo Ricordi
Andreas Tzakis
Rodolfo Alejandro
Yijun Zeng
Anthony J. Demetris
Patricia Carroll
Daniel H. Mintz
Thomas E. Starzl
The Departments of Surgery and Pathology
University of Pittsburgh School of Medicine
Pittsburgh, Pennsylvania 15213
The Diabetes Research Institute
University of Miami School of Medicine
Miami, Florida 33101

REFERENCES


Received 18 March 1991.
Accepted 12 April 1991.