**Name of Journal:** *World Journal of Transplantation*

**ESPS Manuscript NO: 24748**

**Manuscript Type:** Case Report

**Islet Autotransplantation in a Patient with Hypercoagulable Disorder**

**Running Title**: Islet auto-transplants in challenging condition

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**Conflict of interest:** The authors have no potential conflict of interest.

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**ABSTRACT**

Total pancreatectomy and islet auto transplantation is a good option for chronic pancreatitis patients who suffer from significant pain, poor quality of life, and the potential of type 3c diabetes and pancreatic cancer. Portal vein thrombosis is the most feared complication of the surgery and chances are increased if the patient has a hypercoagulable disorder. We present a challenging case of islet auto transplantation from our institution. A 29-year-old woman with *PAI-4G/4G* variant and a clinical history of venous thrombosis was successfully managed with a precise peri- and post-operative anticoagulation protocol. In this paper we discuss the anti-coagulation protocol for safely and successfully caring out islet transplantation and associated risks and benefits.

**Key words**: islet transplantation, autoislet transplant, pancreatectomy, chronic pancreatitis, hypercoagulable disorder, and heparin

**Core tip:** Total pancreatectomy and islet auto-transplantation is an option for select patients with chronic pancreatitis. Portal vein thrombosis is the most feared surgical complication and chances are increased if the patient has a hypercoagulable disorder. The paper describes important topics like the management of the anticoagulation in the peri-operative period.

**Citation:**

**Desai CS**, Khan KM, Cui W. Islet Autotransplantation in Patients with Hypercoagulable Disorder

**Audio Core Tip**

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**INTRODUCTION**

Patients with chronic pancreatitis suffer from significant pain and associated decrease in the quality of life and also a potential of forming type 3C diabetes and the pancreatic cancers[[1-4](#_ENREF_1" \o "Chen, 2006 #225)]. It is an inflammatory disease, which is characterized by irreversible, morphological changes that cause permanent loss of function, and fibrosis and development of severe pain and complications. Over time, fibrosis in the pancreas, results in destruction of the islet cells, and patients are at risk of diabetes[[1](#_ENREF_1" \o "Chen, 2006 #225), [3](#_ENREF_3" \o "Malka, 2000 #227), [4](#_ENREF_4" \o "Schneider, 2002 #226)]. The risk of pancreatic cancer is 10 to 15 fold higher in patients with chronic pancreatitis and if it is associated with hereditary pancreatitis with genetic mutations, then the lifetime risk is 75%[[2](#_ENREF_2" \o "Freelove, 2006 #224), [5](#_ENREF_5" \o "Lowenfels, 1997 #417)]. Many surgical, medical, endoscopic and intervention radiological treatments are applied to these patients, despite which many still suffer from continuous dependence on narcotics and bad quality of life.

Total pancreatectomy and autologous islet cell transplantation is a great option for selected patients with chronic pancreatitis[[6-14](#_ENREF_6" \o "Kesseli, 2015 #352)]. Islet auto transplantation helps to take care of 3 Ps that are necessary for this disorder: 1) Pain relief, 2) Prevention of the brittle diabetes mellitus, 3) Prevention of pancreatic cancer[[15](#_ENREF_15" \o "Desai, 2015 #255)]. At times, the results of the autologous islet cell transplantation are criticized because the variable insulin independence rate reported[[16](#_ENREF_16" \o "Sutherland, 2008 #143), [17](#_ENREF_17" \o "Wilson, 2014 #72)]. We have previously argued that the insulin independence is not the only marker of the success, the wide marker of the success would be euglycemia, preventing cancer and having better quality of life[[15](#_ENREF_15" \o "Desai, 2015 #255)].

Good outcomes of islet auto transplantation are based on various factors from selection of the case to performing safe surgery, good isolation and safe injection of the cells followed by good engraftment of the islet cells. Once the islets are isolated and brought back to the patient, a small angiocatheter is introduced in one of the vessels either the splenic vein stump or any vessels draining into the superior mesenteric vein to infuse these cells into the portal vein so that they can flow to the liver. Safety is important in terms of decreasing the risk of thrombogenesis in these vessels by paying attention to the details of the procedure, the physiology of the patient, and the liver pathology[[18](#_ENREF_18" \o "Desai, 2013 #337)]. Surgical complications are most dreaded compared to the long- term outcome and insulin dependency because they can add to significant morbidity and therefore poor quality of life to the patient. Porto-venous thrombosis would arguably be the most important complication. It can vary in magnitude from a segmental vein to thrombosis of the main portal vein and potentially complete thrombosis of the superior mesenteric access requiring a bowel resection and consequent problems[[19](#_ENREF_19" \o "Thomas, 2010 #342), [20](#_ENREF_20" \o "Memsic, 1984 #348)]. The risk of portal vein thrombosis will be increased if the patient has a hypercoagulable disorder such as factor V Leiden mutation or plasminogen activator inhibitor *(PAI)-1* gene mutation.

We report a case from our new program with physiological challenge in the context of issues described. These include a case of islet autotransplantation performed in a patient with a hypercoagulable disorder. To our knowledge, it is the first such case in the literature.

**Case Report**

The patient was a 29-year-old lady (body weight 83 kg, body mass index 29.3 kg/m2) with a history of chronic abdominal pain related to chronic pancreatitis. At the time of her initial visit she was in the emergency room or hospitalized on a weekly basis. Her history dated back 13 years and she had been on narcotics for 6 years. She had undergone 7 ERCPs over the years and MRI had shown pancreas divisum. Our own MRI scoring system[[21](#_ENREF_21" \o "Khan, 2013 #176)] indicated minimal pancreatic damage (atrophy, 1/6). The pre-operative C-peptide was 1.75 ng/mL and hemoglobin A1c was 5.5%. We also considered gall stone disease, alcohol and completed a genetic analysis for common hereditary gene mutations that are causally associated with chronic pancreatitis. She had also reported having developed thrombosis related to PICC line placement on multiple occasions at an outside institution. During her evaluation we obtained hypercoagulability studies, which included factor V Leiden mutation, prothrombin gene mutation, plasminogen activator inhibitor *(PAI)-1* gene mutation and level, clotting factor VII, VII, protein C, protein S levels, methylenetetrahydrofolate reductase *(MTHFR)* gene mutations and an autoimmune thrombophilia screen. She was found to be homozygous for the 4G variant of the *PAI-1* gene and heterozygote for the *MTHFR A1298C*.

Her surgery was performed using the technique described earlier[[22](#_ENREF_22" \o "Desai, 2011 #178)] and islet infusion was also done through splenic vein stump. Islet preparation was performed at the cGMP facility in the Islet Cell Laboratory at the Georgetown University Hospital.

The pancreas was explanted post 1 and half min of warm ischemia time and placed immediately into an ice-cold Viaspan solution in a sterile container and delivered to the lab on ice. On arrival of the lab, the pancreatic duct was cannulated after trimming. The pancreas was then divided into two portions at the neck. On the cut surface both openings of the pancreatic duct were cannulated with a 14-gauge cannula and connected with a 60cc syringe through an extension tube. A warm mixed enzyme solution of collagenase, buffers, and proteases were infused directly into the cannulated pancreatic duct using a 60 cc syringe. The parenchyma was then repeatedly injected with mixed enzyme solution under manual pressure generated by a 60 cc syringe to monitor the gland for optimal distension and distribution of the enzyme solution throughout the parenchyma. The distended pancreas was then digested using the semi-automated method of Ricordi[[23](#_ENREF_23" \o "Ricordi, 1988 #422)]. The pancreas weighed 65.9 g. The total cold ischemia time from removal of the pancreas to completion of trimming was 51 min. The digestion rate was 92.2% post 18 min of digestion. After purification using a modified continuous density gradient method with cell processor COBE2991[[24](#_ENREF_24" \o "Anazawa, 2011 #137)], the final pellet was reduced from 36 mL to 12 mL[[25](#_ENREF_25" \o "Wilhelm, 2013 #431)]. The total islet yield was 459164 IEQ which was quantified as islet equivalents (IEQ) by normalizing the islet mass to an islet size of 150micrometer diameter. The islet recovery was 7552 IEQ/gram of pancreas tissue. The final pellet was suspended with 5% human serum albumin and 35 units of Heparin per kilogram of patient body weight. The islet infusion dose was 5532 IEQ per kilogram recipient body weight (IEQ/kg).

The islet cells were infused via catheter into the portal vein or mesenteric venous tributaries for engraftment into the liver. To reduce complication rates of acute portal hypertension and thrombosis in this case, an endotoxin free, low-volume (12 mL pellet), homogenous cell suspension, which was prepared through purification procedure, was infused while the patient is given intravenous heparin. We gave the patient 35 U/kg intravenously in addition to the 35 U/kg of Heparin with the islets; the patient therefore received a therapeutic dose of 70 U/kg of heparin. Portal pressures were closely monitored during infusion, as it has been recently demonstrated that the risk of thrombosis increases tenfold (1.52–15.2%) in those with portal pressure changes greater than 25 cm H2O[[25](#_ENREF_25" \o "Wilhelm, 2013 #431)]. The pre infusion portal pressure was 4.5 cm/saline and the post infusion pressure was 15 cm/saline.

Heparin was started intra-operatively. Fifty IU/Kg of body weight bolus before the infusion of islet cells followed by 25000 IU mixed with 500 mL of D5 ½ NS at the rate of 10 IU/Kg/hour. Postoperatively, the patient was continued on a heparin drip according to our protocol and activated thromboplastin time was maintained in the range of 50 to 60 seconds. At the end of three days when she started on clear liquid diet, we continued the patient on low molecular weight heparin and monitored with anti-Xa activity factors maintained between 0.6 to 1 international units/mL. Postoperative Doppler ultrasound of the liver was performed on day 1, 2, and 5 and once weekly for one month and biweekly for another two months. Specifically, the doppler studies during the first week demonstrated patency and normal flow in the portal veins, hepatic arteries and veins; the main portal vein peak velocities ranged between 25-38 cm/sec, left and right portal vein velocities ranged from 11-27 cm/sec. The patient was discharge home after 14 days. At three months the patient was off insulin with a C-peptide of 1.95 ng/mL. At the end of three months, the dose of low molecular weight heparin was reduced to maintain anti-Xa level between 0.3 to 0.6 international units/mL. Six months after the surgery, the low molecular weight heparin was discontinued after consultation with hematology. The patient did not develop venous thrombosis of any form during follow-up and was able to resume a normal life.

 **DISCUSSION**

Total pancreatectomy and islet auto transplantation is being described by some as a radical procedure for patients with chronic pancreatitis though it has a clear role in the treatment of patients with chronic pancreatitis. Patients undergo multiple endoscopic procedures and fail to get a satisfactory outcome and all the time their narcotic requirement keeps escalating. This definitive procedure is feared because of surgical complications like portal vein thrombosis and also the failure of the islets to prevent diabetes.

Hypercoagulability is a significant risk factor for portal vein thrombosis. In one study 28% of patients with portal vein thrombosis had an inherited thrombophilic disorder[[26](#_ENREF_26" \o "Dutta, 2008 #349)]. Of this factor V Leiden mutation was the most common (11%) followed by anti-thrombin III deficiency (11%) and protein c deficiency (8%). Prothrombin gene mutations are also commonly implicated in venous thrombosis[[27](#_ENREF_27" \o "Rosendaal, 1998 #350)]. The PAI 4G variant and MTHFR mutations are considered less severe though do have an increased risk for venous thrombosis after major surgery including transplantation. Such situations are challenging because of the post-operative risk of thrombosis leading to graft failure or bleeding from anti-coagulation. However, many such transplants are carried out in a safe manner. Our patient had a *PAI-1* gene mutation, which was only diagnosed after diligent history taking helped us to obtain the risk in this case. The authors have previously worked at different auto islet cell transplantation centers and as with other surgeries it was not routine to do a hypercoagulable workup since obtaining this panel in every patient is very expensive and may not be cost effective[[15](#_ENREF_15" \o "Desai, 2015 #255), [18](#_ENREF_18" \o "Desai, 2013 #337), [22](#_ENREF_22" \o "Desai, 2011 #178)].

The incidence of portal vein thrombosis after islet auto-transplant is low but can be risky and life threatening. There are few previous individual reports of portal vein thrombosis after islet auto-transplantation[[20](#_ENREF_20" \o "Memsic, 1984 #348)] and one series that indicated a prevalence of 3.7% after clinical islet transplantation[[28](#_ENREF_28" \o "Kawahara, 2011 #430)]. There is however no systematic study of the cause of thrombosis in such cases. In a previous publication we have noted that there may be unrecognized mild fibrosis and or steatosis[[18](#_ENREF_18" \o "Desai, 2013 #337)]. We were however unable to show that any specific histologic pattern is was more susceptible to venous thrombus formation. To prevent portal venous thrombosis in patients such as ours above with pre-existing risk factors it is imperative to identify at risk patients and manage these patients with therapeutic anticoagulation with heparin. Heparin also has advantage in the islet engraftment process and hence it has dual advantage, but has a significant risk of post-operative bleeding and hence it is very important that the surgery is performed with good hemostasis. Heparin is given by almost all the centers performing auto-islet cell transplant to their patients. However, there are no consensus guidelines on the amount and duration it needs to given. We adapted an approach in which we start a heparin drip in operating room at the time of starting islet infusion after giving bolus. It is continued for the next three days maintaining the activated thromboplastin time in the range of 50 to 60 seconds. At the end of three days when the patient starts taking clears, we continue with low molecular weight heparin two times a day dose based on patient’s weight with anti-Xa activity factors maintained between 0.6 to 1 international units/mL. Patient's postoperative Doppler ultrasound on the liver is done on postoperative day 1, 2, and 5 and subsequently was done once weekly for one month and then twice weekly for another two months if they are at high risk. High risk is defined by three main factor; 1) hypercoagulable disorder 2) previous history of deep venous thrombosis other than segmental splenic vein thrombosis related to chronic pancreatitis (even if the hypercoagulable panel is normal) 3) high portal pressure after infusion (more than 25 cm of saline). If the patient is high risk then at the end of three months, low molecular weight heparin dose is reduced to maintain anti-Xa level to be between 0.3 to 0.6 international units/mL. Six months after the surgery, the low molecular weight heparin is discontinued after consultation with hematology. If the patient is not at high risk then after two weeks dose is reduced and then stopped after another two weeks.

In summary, islet auto transplantation in itself is a challenging procedure and even more challenges can arise medically if there are physiological challenges like a hypercoagulable disorder. Despite all these challenges with careful teamwork and experience, these patients can be safely managed.

**CONCLUSION**

Islet auto transplantation is a challenging procedure and even more challenges can arise medically; if there are physiological challenges like a hypercoagulable disorder. Despite all these challenges with careful teamwork and experience, these patients can be safely managed.

**COMMENTS**

(1) Case characteristics: total pancreatectomy and islet autotransplantation complicated by primary hypercoagulability that presented as repeated thrombosis of indwelling venous lines.

(3) Clinical diagnosis: the presentation was characterized by symptoms of chronic pancreatitis but a history of deep venous thrombosis.

(3) Differential diagnosis: An alternative explanation to a primary hypercoagulability to account for thrombosis if IV lines would be that the presence of intravenous lines themselves was the cause of catheter thrombosis.

(4) Laboratory findings: screening for hypercoagulability included plasma proteins, genetic defects and autoimmunity as potential causes of thrombosis with the patient having a plasminogen activator inhibitor *(PAI)-1* variant.

(5) Imaging diagnosis: serial ultrasounds were used to monitor for portal vein thrombosis after islet infusion in to the portal vein after total pancreatectomy

(6) Pathological diagnosis: confirmation of chronic pancreatitis as the casuse for abdominal pain

(7) Treatment: heparin infusion followed by low molecular weight heparin and aspirin as prophylaxis for a prothrombotic state.

(8) Related reports: there are previous cases of a hypercoagulability giving rise to deep venous thrombosis, most notably with Factor V Leiden mutation.

(9) Term explanation: hypercoagulability refers to a pathological increase in the tendency to form intravascular clots

(10) Patients undergoing major intraabdominal operations should be screened for a hypercoagulable state if there is any history of abnormal venous clot formation.

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**Abbreviation**: TPAIT-total pancreatectomy and autologous islet transplantation