Applicability and reproducibility of the CPAT-grading system for pancreas allograft thrombosis

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**ABSTRACT**

Purpose: Although pancreas allograft thrombosis (PAT) incidence has progressively decreased, it remains the most common cause of early graft failure. Currently, there is no consensus on documentation of PAT, which has resulted in a great variability in reporting. The Cambridge Pancreas Allograft Thrombosis (CPAT) grading system has recently been developed for classification of PAT. In this study we aimed to assess the applicability and validate the reproducibility of the CPAT grading system.

Methods: This study is a retrospective cohort study. Selected for this study were all 177 pancreas transplantsations performed at our center between January 1st, 2008 and September 1st, 2018 were included.

Results: A total of 318 Computed Tomography (CT) images was reevaluated according the CPAT system by two local radiologists. Inter-rater agreement, expressed in Cohen’s kappa was 0.403 for arterial and 0.537 for venous thrombosis. Inter-rater agreement, expressed in the Fleiss’ kappa, within clinically relevant thrombosis categories was 0.626 for Grade 2 and 0.781 for Grade 3 venous thrombosis.

Conclusions: Although not perfect, we believe that implementation of the CPAT system would improve current documentation on PAT. However, it is questionable whether identification of a small Grade 1 thrombosis would be relevant in clinical practice. Furthermore, a good quality CT scan, including adequate phasing, is essential to accurately identify potential thrombus and extend after pancreas transplantation.

**Keywords:**
- Pancreas transplantation
- Pancreas allograft thrombosis
- CPAT grading system
- Applicability
- Reproducibility

1. Introduction

Pancreas transplantation has been proven to be a successful treatment for patients with insulin-dependent diabetes mellitus. Survival rates of pancreas transplantation have further improved over the past decades. A recent study showed 1- and 2-years patient survival rates of almost 100 % and 1- and 2-year graft survival rates over 80 % in deceased-after-brain-death (DBD) organs and over 90 % in deceased-after-circulatory-death (DCD) organs [1]. Technical failure is the leading cause for early graft loss in pancreas transplantation [2–6]. Pancreas allograft thrombosis (PAT), which is the most frequent surgical complication and responsible for 29 % of all early graft loss, remains an unsolved problem and usually results in graft loss [7,8]. Although incidence has progressively decreased, pancreas allograft thrombosis is still reported to develop in 3–34 % of all transplanted patients [9–12]. Complete thrombosis is likely to result in graft loss, although cases of successful salvage of the graft by performing a thrombectomy have been reported [8,13]. In contrast to complete thrombosis, little is known about partial thrombosis. Partial thrombosis is presumably underreported and develops and occurs more frequently than complete thrombosis [13]. Different interventions and strategies for detection and treatment of partial thrombosis have been described in literature [8,13,}

**Abbreviations:** CIT, cold ischemia time; CPAT, Cambridge Pancreas Allograft Thrombosis; DBD, donor after brain death; DCD, donor after cardiac death; LMWH, low molecular weight heparin; PACS, picture archive and communication system; PAK, pancreas after kidney; PAT, pancreas allograft thrombosis; PTA, pancreas transplant alone; SA, splenic artery; SD, standard deviation; SMA, superior mesenteric artery; SMV, superior mesenteric vein; SV, splenic vein; SPK, simultaneous pancreas kidney.

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2. Materials and methods

2.1. Study population and design

This study is a retrospective cohort study of all pancreas transplantations (SPK, pancreas transplant alone (PTA), pancreas after kidney (PAK) and retransplantations) performed at the Leiden University Medical Center (LUMC) between January 1st of 2008 and September 1st of 2018. Data regarding recipient and transplant characteristics and the original radiology report were retrieved from the patient charts. Inclusion criterion for the study was the availability of an abdominal Computed Tomography (CT)-scan, performed within the first 3 months after transplantation. Follow-up time was 3 months after transplantation.

2.2. Pancreas transplantation

In our center, arterial revascularization of the pancreas allograft is usually performed using the donor common, external and internal iliac arteries. The donor iliac arteries are converted into a Y-graft conduit and are anastomosed to the splenic artery (SA) and superior mesenteric artery (SMA) of the allograft. Less frequently, when the donor SMA and celiac trunk remained with the pancreas graft including a common, external and internal iliac arteries. The donor iliac arteries are converted into a Y-graft conduit and central occlusive thrombosis (Grade 3).

Currently, there are no solid guidelines for documentation on pancreas allograft thrombosis, especially on partial allograft thrombosis. This has resulted in great variability in reporting of thrombosis. Verification of these reports led to the conclusion that the reports are not suitable for analysis or comparison. Thus, implementation of a grading system, such as the CPAT grading system would be eligible. However, the applicability of the CPAT grading system has not been described by others so far. This study aims to assess the applicability of the CPAT grading system and validate its reproducibility in a different cohort.

Table 1

| Grade | Description |
|-------|-------------|
| 0     | No thrombosis |
| 1     | Peripheral thrombus – Thrombus lies in the very distal vessel the transected margin of the SMV/SV or SMA/SA and lies in a single branch only, without encroachment into the main trunk of the vessel |
| 2     | Venous: thrombus extending into parenchymal vessels/main trunk of the SMV or SV to the SMV/SV confluence but not into the portal vein |
| 3     | Central occlusive thrombosis |

2.3. CT imaging

Each pancreas recipient received a routine abdominal CT scan within the first week postoperatively as part of the local pancreas-transplant protocol. In case of impaired kidney function, imaging was performed later. All routine abdominal CT scans were selected for this study. In addition, CT scans performed within 3 months after transplantation with specific request for evaluation of the vascular status of the graft, were included. All CT scans were performed on a Toshiba Aquilion or Toshiba Aquilion One 320-slice CT scanner with the following parameters: routine section thickness 1.0 mm; section thickness after reconstruction, 1.0–5.0 mm; filtered back projection reconstruction method; 120 kV; Automatic Exposure Control. Standard post-transplant protocol in our institute constituted of a two-phase contrast-enhanced CT scan of the abdomen, including an early-arterial phase of the lower abdomen (including the pancreas transplant) and a parenchymal phase of the entire abdomen. The two phases were respectively derived with a sure start in the abdominal aorta and a 35 s delay. Non-ionic iobitrol 350 mg I/mL was infused with a dose of 1.4 mL/kg and a flow of 0.05 mL/kg/s for 30 s. All scans received a unique study number and were reconstructed anonymously in the picture archive and communication system (PACS).

2.4. CT analysis

Reevaluation of the 318 CT scans was performed independently by one radiologist and one senior resident specialized in abdominal imaging (JK and BK) with respectively 7 and 4 years of experience. Reviewers were blinded to each other’s results as well as to the original report and the clinical patient characteristics. The CT findings were classified according to the CPAT grading system developed by Hakeem et al. [15]. Additional clarification on the classification was received from the research team which conducted the original study. Based on the additional clarification, our reviewers classified non-occlusive thrombosis in the portal vein and the Y-graft as a Grade 2 thrombosis. Original drawings demonstrating the limits of each specific grade of the CPAT system were provided (Supplement 1). Our reviewers practiced the use and implementation of the CPAT system together on a training set of approximately 40 CT scans from pancreas transplants performed in 2006 and 2007, directly prior to onset of the study. Arterial and venous blood supply of the pancreas graft were evaluated separately, the highest grade of thrombosis was reported, whether this concerned only one of both arteries of veins. The reviewers also reported their opinion on the quality of both the arterial and venous phase of the CT scan (good, moderate, poor). Quality of the scans was primarily defined by vascular enhancement. Reviewers also looked at the presence of artifacts.

2.5. Statistics

Descriptive analyses were performed on patient demographics and frequencies of CPAT scores. The applicability of the CPAT grading system was assessed by the inter-rater agreement. The inter-rater agreement was evaluated using the Cohen’s kappa and the Fleiss’ kappa. The strength of agreement was interpreted using guidelines established by Landis and Koch [16]. According to Landis and Koch, κ values < 0 have a poor strength of agreement, κ values of 0–0.2 a slight strength of...
agreement, κ values of 0.21–0.4 fair agreement, κ values of 0.41–0.6 moderate agreement, κ values of 0.61–0.8 substantial agreement and a κ value of 0.81–1 is considered as perfect agreement. The inter-rater agreement was assessed separately for arterial and venous thrombosis. A Stuart Maxwell test was performed to determine the difference in distribution. All confidence intervals were presented as 95% confidence intervals. Results were considered to be significant if p < 0.05. All data analyses were performed using SPSS Statistics version 25.

3. Results

Between January 1st 2008 and September 1st 2018, 182 patients underwent a pancreas transplantation in our center. Eligible for inclusion were 173 transplant patients corresponding with 177 pancreas transplantations (four cases of re-transplantation) and a total of 318 CT scans. Five patients were excluded from the study because no CT scan was performed within the first three months after transplantation. The study population included 76 females and 101 males (42.9% vs 57.1%). A Y-graft, had been performed in 76.3% (N = 248) of all transplantations. In 22.6% (N = 71) of all patients had developed (partial) thrombosis. This number of new cases of thrombosis. According to Reviewer 1 (R1), 87.7% (279/318) of all scans, 187 scans were scored a Grade 1 arterial thrombosis (56.6%), 58 scans a Grade 2 arterial thrombosis (18.2%) and one case of complete arterial thrombosis (0.3%).

3.1. Pancreas allograft thrombosis

Original CT report described arterial thrombosis in 17.3% (55/318), venous thrombosis in 30.2% (96/318) and a combination of arterial and venous thrombosis in 9.7% (31/318), either complete or partial.

Re-evaluation of the 318 scans by both reviewers yielded a great number of new cases of thrombosis. According to Reviewer 1 (R1), 87.7% (279/318) of all patients had developed (partial) thrombosis. This was 82.7% (263/318) according to Reviewer 2 (R2). Of all newly detected arterial thrombosis, Grade 1 was by far the most common (R1:122/150 (81.3%), R2:119/143 (83.2%)). Of all newly detected venous thrombosis, Grade 2 was most common (R1:52/103 (50.5%), R2:54/92 (58.7%)).

3.1.1. Arterial thrombosis

The different grades of arterial thrombosis are demonstrated in Fig. 1. Both reviewers reported higher thrombosis incidences compared to original report. R1 detected arterial thrombosis in 80.5% of all scans (256/318). Of the 318 scans, 187 scans were scored a Grade 1 arterial thrombosis (58.8%), 69 scans a Grade 2 arterial thrombosis (21.7%) and no case of complete arterial thrombosis was reported. R2 detected arterial thrombosis in 75.2% of all scans (239/318). Of the 318 scans, 180 scans were scored a Grade 1 arterial thrombosis (56.6%), 58 scans a Grade 2 arterial thrombosis (18.2%) and one case of complete arterial thrombosis (0.3%).

3.1.2. Venous thrombosis

The different grades of venous thrombosis are demonstrated in Fig. 2. Both reviewers reported higher thrombosis incidences compared to the original report (Fig. 1). R1 detected venous thrombosis in 61.6% of all scans (196/318). Fifty-seven scans were scored a Grade 1 venous thrombosis (17.9%), 107 scans a Grade 2 venous thrombosis (33.6%) and 32 scans a Grade 3 venous thrombosis (10.1%). R2 detected venous thrombosis in 75.2% (239/318) of all scans. Of the 318 scans, 180 scans were scored a Grade 1 venous thrombosis (56.6%), 58 scans a Grade 2 venous thrombosis (18.2%) and one case of complete venous thrombosis (0.3%).

Table 2

| Transplant Factor | 2008 | 2009 | 2010 | 2011 | 2012 | 2013 | 2014 | 2015 | 2016 | 2017 | 2018 |
|------------------|------|------|------|------|------|------|------|------|------|------|------|
| Donor Type* | DBD | 80.8% (N = 143) | DCD | 16.9% (N = 30) | Female | 56.5% (N = 100) | Male | 41.2% (N = 73) | Donor Age | 37.3 (SD = 13.06; range 10–65) | Donor BMI | 23.36 (SD = 2.81; range 15.4–29.4) |
| Transplant Type | SPK | 84.2% (N = 149) | PAK | 14.1% (N = 25) | PTA | 1.7% (N = 3) | Transplantectomy | 14.7% (N = 26) | Retransplantation | 2.3% (N = 4) | CIT, hrs | 9.83 (SD = 2.25; range 4.0–15.3) | WIT, min | 27.5 (SD = 8.01; range 10.0–64.0) | Reconstruction Type | Y-graft | 76.3% (N = 135) | Aortic Patch | 22.6% (N = 40) | Other | 1.1% (N = 2) |
| Recipient Factor | Recipient Sex | Female | 42.9% (N = 76) | Male | 57.1% (N = 101) | Recipient Age, y | 43.0 (SD = 8.23; range 23–64) | Recipient BMI | 24.9 (SD = 3.57; range 16.5–34.7) |
| Primary Disease | 98.3% (N = 174) | Diabetes Mellitus Type 1 | 0.0% (N = 1) | Other | 1.1% (N = 2) | Medical History | 13.6% (N = 24) | Coronary Vascular Disease | 7.9% (N = 14) | Thromboembolic Events | 5.1% (N = 9) | Induction Therapy | Postoperative Factor | Alemtuzumab | 96.0% (N = 170) | IL-2-Receptor Blocker | 2.8% (N = 5) | Other | 1.1% (N = 2) |
| Thrombosis (original report) | No Thrombosis | 57.1% (N = 101) | Thrombosis | 42.9% (N = 76) | Arterial | 6.8% (N = 12) | Venous | 26.0% (N = 46) | Both | 10.7% (N = 19) | Therapeutic anticoagulation | 58.2% (N = 103) | No anticoagulation | 41.8 (N = 74) |

* missing: N = 4. Continuous variables are summarized by means and standard deviations. Categorical variables are summarized by percentages and frequencies. Recipient age was measured at time of transplantation. DBD, donor after brain death; DCD, donor after cardiac death; CIT, cold ischemia time; IL-2-Receptor, interleukin-2-receptor; LMWH, low molecular weight heparin; MMF, mycophenolate mofetil; PAK, pancreas after kidney; PTA, pancreas transplant alone; SPK, simultaneous pancreas kidney; VKA, vitamin-K-antagonist; WIT, warm ischemia time.
thrombosis in 59.4% of all scans (189/318), which is a lower incidence compared to R1. Fifty-one scans were scored a Grade 1 venous thrombosis (16.0%), 115 scans were scored a Grade 2 venous thrombosis (36.2%) and 23 scans were scored a Grade 3 venous thrombosis (7.2%).

3.2. Inter-rater agreement

Both reviewers classified all 318 CT scans into 1 of 4 categories. All scans received an arterial thrombosis score and a venous thrombosis score. For arterial thrombosis, overall Cohen’s kappa was 0.403 \((SE = 0.046, 95\% CI 0.313;0.493)\). For venous thrombosis, overall Cohen’s kappa was 0.537 \((SE = 0.038, 95\% CI 0.463;0.611)\). The CT scans were also classified according the quality of arterial and venous phase scan and Cohen’s kappa values were measured for different categories of quality (Table 3). For arterial thrombosis, Cohen’s kappa was 0.206 \((SE = 0.165, 95\% CI 0.028;0.618)\) in scans of medium quality (22/318) and 0.248 \((SE = 0.049, 95\% CI 0.334;0.522)\) in scans of good quality (284/318). For venous thrombosis, Cohen’s kappa was 0.250 \((SE = 0.141, 95\% CI 0.074;0.626)\) in scans of poor quality (24/318), 0.348 \((SE = 0.112, 95\% CI 0.128;0.568)\) in scans of medium quality (43/318) and 0.580 \((SE = 0.041, 95\% CI 0.506;0.660)\) in scans of good quality (251/318).

For evaluation of the inter-rater agreement in individual grades of thrombosis, Fleiss’ kappa values were calculated (Table 4). For arterial thrombosis, Fleiss’ kappa for Grade 0 was 0.533 \((95\% CI 0.243;0.463)\), for Grade 1 was 0.349 \((95\% CI 0.239;0.459)\), for Grade 2 was 0.538 \((95\% CI 0.428;0.647)\). For venous thrombosis, Fleiss’ kappa within Grade 0 was 0.520 \((95\% CI 0.410;0.629)\), within Grade 1 was 0.286 \((95\% CI 0.176;0.396)\), within Grade 2 was 0.626 \((95\% CI 0.516;0.736)\) and within Grade 3 was 0.781 \((95\% CI 0.671;0.891)\).

The difference in distribution between both reviewers was calculated using the Stuart Maxwell Test for marginal homogeneity. Difference in distribution was considered to be significant for arterial thrombosis \((p = 0.025)\), whereas for venous thrombosis it was not \((p = 0.233)\).

3.3. Clinical outcome

In this study population, 26 transplantectomies were performed in 177 transplantations \((14.7\%)\). One patient lost both his first and second transplant and underwent two transplantectomies during the duration of this study. Nineteen grafts \((10.7\%)\) were removed for other reasons including bleeding, leakage and one case of proven rejection. Sensitivity and specificity were calculated for evaluation of the accuracy of assessment of Grade 3 thrombosis. Sensitivity was 80.0% and specificity 96.4%.

4. Discussion

This study was primarily designed to assess the applicability and the reproducibility of the CPAT grading system on our pancreas transplant database. The CPAT grading system is the first classification scheme developed for pancreas allograft thrombosis. Although appearing promising, the applicability and reproducibility of the CPAT grading system has not been reported by others to date. To evaluate whether we can implement this grading system into our clinical practice, we assessed the inter-rater agreement of two local radiologists on a set of 318 CT scans from 177 pancreas transplant patients. Our results showed a moderate inter-rater agreement for both arterial and venous thrombosis. Cohen’s kappa for overall arterial thrombosis was 0.403 and for overall venous thrombosis 0.537. According to the guidelines of Landis and Koch [16] these kappa values correspond with fair strength of agreement in arterial thrombosis and moderate strength of agreement in venous thrombosis.

For arterial thrombosis, CPAT scores from both reviewers differed significantly \((p = 0.025)\). Analysis of our results suggest that there is a correlation between the quality of the scans and the extent of agreement. Not surprisingly, the agreement in scans, classified as poor quality, was lower in both arterial and venous thrombosis. Results in this category showed different CPAT scores between both reviewers (Supplement 2).

Cases of disagreement between no thrombosis and Grade 2 \((6/318, both arterial and venous thrombosis)\) and between no thrombosis and Grade 3 \((3/318, venous thrombosis)\) thrombosis were documented. In clinical practice, these cases of disagreement would correspond with disagreement on administration of anticoagulants (Grade 2/Grade 3) or not (Grade 0). This depicts the importance of the quality of a CT scan for adequate evaluation of the vascularization. Acceptance of poor CT-scans could increase the risk of incorrect diagnosis in real clinical practice in which evaluation is only performed by a single reviewer. As was described in the introduction, results from the study by Hakeem et al. [15] showed that Grade 1 and 2 of arterial thrombosis and Grade 1 of venous thrombosis can be managed without anticoagulation. However, patients with Grade 2 venous thrombosis or complete arterial/venous thrombosis have shown to benefit from anticoagulative therapy. Hence, identification and consensus in these cases is paramount. Also, in clinical practice the radiologists are not blinded to the patient’s clinical characteristics and it is likely that a more weighted decision can be made concerning the thrombosis grade, presumably after consultation of the surgeon.

Inter-rater agreement for grade subgroups was measured and expressed in the Fleiss’ kappa. In the arterial thrombosis set, reviewers disagreed on one CT scan: whether it was proximal partial (Grade 2) or complete (Grade 3) thrombosis. Because of the small number of cases in this category \((N = 1)\), a valid Fleiss’ kappa value could not be measured for this category. Although outcomes on the overall inter-rater agreement are relatively low, agreement on thrombosis grades, in which treatment is considered to be effective, was found to be substantial.

In our experience, description of the CPAT grading system provided in the article by Hakeem et al. was not specific enough for accurate implementation. Even with the additional information provided by the research team which developed the CPAT grading system, our radiologists experienced difficulty with some parts of the classification. This difficulty was mostly experienced in differentiation between a Grade 0 and Grade 1 thrombosis, stated by the Cambridge research team as respectively the absence or presence of a visible thrombus at the transected margin, not extending into the main vessel. In our opinion, development of a thrombus at the transected margin is considered to be a physiological process in a blind ending vessel. For that reason, presence of a thrombus at the distal end of the vessel may be expected or even assumed. However, due to the small caliber of the vessels it is not always possible to follow the vessel until it’s transected margin or to distinguish the contours of a distal thrombus on a CT scan. In this situation, the decision between presence or absence of a distal thrombus (Grade 0 vs Grade 1) will most likely be based on the reviewer’s personal intuition.

Difficulty was also experienced in the differentiation between Grade 1 and Grade 2 thrombosis in the small branches of SA, as a result of the definition in the CPAT grading system. The CPAT system defines Grade 1 thrombosis as the presence of a thrombus in the very distal vessel at the transected margin but not reaching into the main vessel. In our opinion, thrombosis have shown to benefit from anticoagulative therapy. Hence, identification and consensus in these cases is paramount. Also, in clinical practice the radiologists are not blinded to the patient’s clinical characteristics and it is likely that a more weighted decision can be made concerning the thrombosis grade, presumably after consultation of the surgeon.
**Fig. 1.** CT images demonstrating different grades of arterial allograft thrombosis.

A. Grade 0 no arterial thrombosis: axial arterial phase image on the level of the left common iliac artery (white arrowhead) and superior mesenteric artery (SMA) (white arrow) up to the transection margin of the graft (black dotted arrow) without any thrombus. (Note: the transection margin is visible because the mesenteric root is stapled off)

B. Grade 1 arterial thrombus: arterial phase coronal reformatted image demonstrating thrombus (black arrow) in the distal SMA (white arrow), distally to the branch of the inferior pancreaticoduodenal artery (black arrowhead). Y-graft (white dotted arrow), transection margin of the graft (black dotted arrow). Pancreatic head (*).

C. Grade 2 arterial thrombus: arterial phase coronal reformatted image demonstrating thrombus (black arrows) extending into the mid splenic artery (white arrow). Y-graft (white dotted arrow), pancreas allograft (*) The SMA is not shown in this picture.
allograft thrombosis. Since the CPAT grading system is, to our best
knowledge, the only available grading system, implementation of this
system would be a good first step in systematically reviewing pancreas
allograft CT scans. Also, classification according to the CPAT system
would considerably enhance both communication between radiologist
and surgeon and the quality of documentation, which will benefit both
clinical and future scientific purposes.

It was a remarkable finding that on re-evaluation 90 % of all CT-
scans showed some form of thrombosis. We cannot conclude whether all
the detected thromboses should be considered relevant in clinical
practice, since the focus of this study was to evaluate the CPAT grading
system.

One important limitation of this study is the absence of a gold-
standard, different from the CT scan which was re-evaluated for this
study, to compare the results of the re-evaluation to. The only available
clinical outcome is whether a transplantectomy was performed due to

A. Grade 0 no visible venous thrombotic portal phase coronal reformatted image. Completely contrasted superior mesenteric vein (SMV) (white arrow) and portal vein (PV) (white dotted arrow) without any thrombus. Pancreatic head (*), transection margin of the graft (black dotted arrow) B. Grade 1 venous thrombus: portal phase coronal reformatted image demonstrating thrombus (black arrows) only in the very distal vessel at the transected margin but not into the main splenic vein (SV) (white arrow). Pancreas allograft (*) C. Grade 2 venous thrombus: portal phase oblique coronal reformatted image demonstrating a focal thrombus (black arrows) in the distal and mid SV (white arrow) not extending in the PV (white dotted arrow). No thrombus in SMV (white arrowhead). Pancreas allograft (*) D. Grade 3 venous thrombus: portal phase coronal reformatted image demonstrating thrombus in the SMV, SV and PV (black arrows) extending into the inferior vena cava (black arrowheads). Non-enhancing pancreas allograft (*).

Table 3
Inter-rater agreement between both raters using the Cohen’s Kappa (95 % confidence intervals).

| Thrombosis Category | Cohen’s Kappa [95 % confidence intervals] | Interpretation of reliability coefficient | P-value |
|---------------------|------------------------------------------|------------------------------------------|---------|
| Arterial            |                                          |                                          |         |
| Poor quality        | 0.086 [-0.241;0.413]                     | Slight                                   | 0.590   |
| Medium              | 0.295 [-0.028;0.618]                     | Fair                                     | 0.059   |
| Good quality        | 0.428 [0.334;0.522]                     | Moderate                                 | < 0.001 |
| Overall             | 0.403 [0.312;0.493]                     | Fair                                     | < 0.001 |
| Venous              |                                          |                                          |         |
| Poor quality        | 0.350 [0.074;0.626]                     | Fair                                     | 0.003   |
| Medium              | 0.348 [0.128;0.568]                     | Fair                                     | 0.001   |
| Good quality        | 0.580 [0.500;0.660]                     | Moderate                                 | < 0.001 |
| Overall             | 0.537 [0.463;0.611]                     | Moderate                                 | < 0.001 |

* Interpretation for Cohen’s kappa: < 0: poor; 0 – 0.20: slight; 0.21 – 0.40: fair; 0.41 – 0.60: moderate; 0.61 – 0.80 substantial; 0.81 – 1.00: perfect.

Table 4
Inter-rater agreement between both raters using the Fleiss’ kappa (95 % confidence intervals), separately for each thrombosis grade.

| Thrombosis Category | Fleiss’ Kappa [95 % confidence intervals] | Interpretation of reliability coefficient | P-value |
|---------------------|-------------------------------------------|------------------------------------------|---------|
| Arterial            |                                          |                                          |         |
| Grade 0             | 0.353 [0.243;0.463]                     | Fair                                     | < 0.001 |
| Grade 1             | 0.349 [0.239;0.459]                     | Fair                                     | < 0.001 |
| Grade 2             | 0.538 [0.428;0.647]                     | Moderate                                 | < 0.001 |
| Venous              |                                          |                                          |         |
| Grade 0             | 0.520 [0.410;0.629]                     | Moderate                                 | < 0.001 |
| Grade 1             | 0.286 [0.176;0.396]                     | Fair                                     | < 0.001 |
| Grade 2             | 0.626 [0.516;0.736]                     | Substantial                              | < 0.001 |
| Grade 3             | 0.781 [0.671;0.891]                     | Substantial                              | < 0.001 |

* Interpretation for Cohen’s kappa: < 0: poor; 0 – 0.20: slight; 0.21 – 0.40: fair; 0.41 – 0.60: moderate; 0.61 – 0.80 substantial; 0.81 – 1.00: perfect.
complete vascular occlusion. Complete vascular occlusion corresponds with a Grade 3 thrombosis according the CPAT system. Therefore, it was possible to calculate the sensitivity and specificity for the identification of a Grade 3 thrombosis. The Grade 3 thrombosis was considered identified correctly only if both reviewers agreed on the presence of a Grade 3 thrombosis. Unfortunately, there were no clinical features or parameters which could prove the presence of a Grade 1 or Grade 2 thrombosis. Hence, it is not possible to assess the accuracy of re-evaluation of the CT scans classified as a Grade 0, Grade 1 and Grade 2.

We acknowledge that there is a possibility that our reviewers did not classify all CT scans correctly during the re-evaluation. This type of error is called perceptual error and it is an acknowledged and unfortunately unconquered problem in radiology. Complete elimination of this type of error cannot be achieved since radiologic interpretation is a human enterprise which cannot be automated [17]. To minimize the risk of perceptual error in this study, all CT scans were anonymized, and reviewers were blinded to each other’s evaluation. Furthermore, results were documented in structured and uniform reports. Previous studies have reported on the accuracy of CT imaging as a diagnostic tool for pancreas allograft thrombosis [18–21]. Studies report different statements on whether US Doppler or CT imaging should be used for the detection of thrombosis. Tolat et al. [18] described that both arterial and venous pancreatic allograft thrombosis are best displayed with volumetric high-spatial-resolution CT scanning. These statements were supported by O’Malley et al. [19] which reported that cross-sectional imaging plays a key role in the accurate assessment of vascular complications after pancreas transplantation. In our center, CT imaging is considered most accurate for assessment of post-operative complications including thrombosis. For that reason, a routine abdominal CT scan is performed in all pancreas transplant in the first week after transplantation.

Another limitation of our study is the assessment of the inter-rater agreement of the CPAT grading system for only two reviewers. Theoretically, involvement of a third reviewer would contribute to a higher reliability of our results. However, it is no guarantee that the third reviewer will score the CT scans more correctly or as correctly as the first two reviewers. Since there is no gold-standard for interpretation of pancreas allograft thrombosis, it is not possible to state that the two out of three reviewers with the highest agreement implemented the CPAT grading system most accurately.

Since this study is a retrospective analysis, results were based on CT scans performed in the past. For that reason, it was not possible to avoid the inclusion of CT scans which were found to be of poor quality (arterial: 12 scans, venous: 24 scans). Reason for the poor quality was predominantly a relatively hypovolemic state of the patient, resulting in a poor visibility of the venous vessels. However, all included scans were originally requested for evaluation of vascular status of the pancreas allograft and therefore the presence of poor scans in our set reflects the reality in clinical practice.

5. Conclusion

In conclusion, currently there are no solid guidelines for documentation on pancreas allograft thrombosis, which has resulted in variability within CT reports. The CPAT grading system is, to our best knowledge, the first and only available grading system for pancreas allograft thrombosis. Although some training and extra information was needed to precisely understand the classification and to correctly score the CT images, the reviewers considered the CPAT grading system reproducible and applicable. Implementation of the grading system will firstly enhance communication between radiologist and surgeon. Also, it will improve both the documentation and the understanding of pancreas allograft thrombosis. Therefore, we would recommend to implement the CPAT grading into clinical practice. It remains questionable whether identification of Grade 1 thrombosis would be relevant in clinical practice since this is considered to be a physiological process in a blind ending vessel. The agreement within this category has shown to be low and treatment is not considered to be effective in patients with arterial or venous Grade 1 thrombosis. Furthermore, this study depicts the importance of a good quality of CT-scan in diagnosis of pancreas allograft thrombosis. Poor CT scans showed diverging CTAP scores. Therefore, a CT scan, which meets the quality standard in regard to correct contrast phasing, should be demanded to ensure accuracy of the reviewer’s judgment.

CRediT authorship contribution statement

SA Simonis: Conceptualization, Methodology, Formal analysis, Investigation, Data curation, Writing - original draft, Visualization, Project administration. BM de Kok: Investigation, Writing - review & editing. JC Korving: Investigation, Writing - review & editing. WH Kopp: Resources, Writing - review & editing. AG Baranski: Writing - review & editing. VAL Huurman: Writing - review & editing. MNJM Wasser: Writing - review & editing. PJM van der Boog: Writing - review & editing. AE Braat: Conceptualization, Methodology, Validation, Writing - review & editing, Supervision, Project administration.

Declaration of Competing Interest

The authors of this manuscript have no conflicts of interest to disclose.

Appendix A. Supplementary data

Supplementary material related to this article can be found, in the online version, at doi:https://doi.org/10.1016/j.ejrad.2020.109462.

References

[1] P.J.M. van der Boog, E.J.F. de Koning, J.W. de Fijter, A.G. Baranski, A.E. Braat, Pancreas transplantation with grafts from donors deceased after circulatory death: 5 years single-center experience, Transplantation 102 (2018) 353–339.
[2] C. Troppmann, A.C. Graessner, D.L. Dunn, et al., Surgical complications requiring early relaparotomy after pancreas transplantation: a multivariate risk factor and economic impact analysis of the cyclosporine era, Ann. Surg. 227 (1998) 256–268.
[3] W. Steurer, J. Malaise, W. Murt, et al., Spectrum of surgical complications after simultaneous pancreas-kidney transplantation in a prospectively randomized study of two immunosuppressive protocols, Nephrol. Dial. Transplant. (2005) 20, i54 – i61.
[4] W.O. Bechstein, J. Malaise, F. Saudek, et al., Efficacy and safety of tacrolimus compared with cyclosporine microemulsion in primary simultaneous pancreas-kidney transplantation: 1-year results of a large multicenter trial, Transplantation 77 (2004) 1221–1228.
[5] J. Malaise, W. Steurer, A. Koenigsrainer, et al., Simultaneous pancreas-kidney transplantation in a large multicenter study: surgical complications, Transplant. Proc. 37 (2005) 2859–2860.
[6] A. Manrique, C. Jimenez, R.M. Lopez, et al., Relaparotomy after pancreas transplantation: causes and outcomes, Transplant. Proc. 41 (2009) 2472–2474.
[7] G.W. Burke III, G. Gancio, J. Figuero, R. Buiga, L. Olsen, D. Roth, W. Kupin, J. Miller, Hypercoagulable State associated with Kidney–Pancreas transplantation. Thromboelastogram-directed anti-coagulation and implications for future therapy, Clin. Transplant. 18 (2004) 423–428.
[8] A.H. Stockland, D.L. Willingham, R. Paz-Fumagalli, H.P. Greival, J.M. McKinney, C.B. Hughes, E.M. Wals, Pancreas transplant venous thrombosis: role of endovascular interventions for graft salvage, Cardiovasc. Intervent. Radiol. 32 (2009) 279–283.
[9] C. Troppmann, Complications after pancreas transplantation, Curr. Opin. Organ Transplant. 15 (2010) 112–118.
[10] W.H. Kopp, C.A.T. van Leeuwen, H.D. Lam, V.A.L. Huurman, J.W. de Fijter, A. F. Schaapberder, A.G. Baranski, A.E. Braat, Retrospective study on detection, treatment, and clinical outcome of graft thrombosis following pancreas transplantation, Transplant. Int. (2018).
[11] A. Vaidya, A.S. Muthusamy, V.G. Hadjianastasiou, D. Roy, D.E. Elker, P. Moutafellos, A. Muktarid, S. Sihna, P.J. Friend, Simultaneous pancreas-kidney transplantation: to anticoagulate or not? Is that a question? Clin. Transplant. 21 (2007) 554–557.
[12] P. Schenker, O. Vonend, N. Ertas, A. Wunsch, M. Schaeffer, L.-C. Rump, P. Moorthy, Incidence of pancreas graft thrombosis using low-molecular-weight heparin, Clin. Transplant. 23 (2009) 407–414.
[13] G. Gancio, M. Cespedes, L. Olsen, J. Miller, G.W. Burke, Partial venous thrombosis of the pancreatic allograft after simultaneous pancreas-kidney transplantation, Clin. Transplant. 14 (2000) 464–471.
[14] J.A. Fridell, R.S. Mangus, A.B. Mull, T.E. Taber, C.E. Sanders, R.C. Slisher, M.L. Goble, J.A. Powelson, Early reexploration for suspected thrombosis after pancreas transplantation, Transplantation 91 (2011) 902–907.

[15] A. Hakeem, J. Chen, S. Iype, et al., Pancreatic allograft thrombosis: suggestion for a CT grading system and management algorithm, Am. J. Transplant. 18 (2018) 163–179.

[16] J.R. Landis, G.G. Koch, The measurement of observer agreement for categorical data, Biometrics 33 (1977) 159–174.

[17] M.A. Bruno, E.A. Walker, H.H. Abujudeh, Understanding and confronting our mistakes: the epidemiology of error in radiology and strategies for error reduction, RadioGraphics 35 (2015) 1668–1676.

[18] P.P. Tolat, W.D. Foley, C. Johnson, M.D. Hohenwalter, F.A. Quiroz, Pancreas transplant imaging: how I Do It, Radiology (2015) 275.

[19] R.B. O’Malley, M. Moshiri, S. Osman, C.O. Menias, D.S. Katz, Imaging of pancreas transplantations and its complications, Radiol. Clin. N. Am. 54 (2016) 251–266.

[20] F.Q. Vandermeer, M.A. Manning, A.A. Frazier, J.J. Wong-You-Cheong, Imaging of whole-organ pancreas transplants, RadioGraph. 32 (2012) 411–435.

[21] S.Y. Liong, R.E. Dixon, N. Chalmers, A. Tavakoli, T. Augustine, S. O’Shea, Complication following pancreatic transplantations: imaging features, Abom Imaging 36 (2011) 206–214.