The impact of vascular anastomosis time on early kidney transplant outcomes

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Abstract

Background: Most studies have found cold ischemic time to be an important predictor of delayed graft function in kidney transplantation. Relatively less is known about the warm time associated with vascular anastomosis and early outcomes.

Methods: A retrospective cohort of 298 consecutive solitary deceased donor kidney recipients from January 2006 to August 2012 was analyzed to examine the association between anastomosis time and delayed graft function (need for dialysis) and length of hospital stay.

Results: Delayed graft function (DGF) was observed in 56 patients (18.8%). The median anastomosis time was 30 minutes (interquartile range 24, 45 minutes). Anastomosis time was independently associated with DGF in a multivariable, binary logistic regression analysis (odds Ratio (OR) 1.037 per minute, 95% CI 1.016, 1.057, \(P = 0.001\)). An anastomosis time >29 minutes was also associated with a 3.5 fold higher (OR 3.5, 95% CI 1.6, 7.3, \(P = 0.001\)) risk of DGF. Median days in hospital was 9 (interquartile range 7, 14 days). Every 5 minutes of longer anastomosis time (0.20 days per minute, 95% CI 0.13, 0.27, \(P <0.001\)) was associated with 1 extra day in hospital in a multivariable linear regression model. An anastomosis time >29 minutes was associated with 3.8 (95% CI 1.6, 6.0, \(P <0.001\)) more days in hospital.

Conclusion: Anastomosis time may be an underappreciated but modifiable variable in dictating use of hospital resources. The impact of anastomosis time on longer term outcomes deserves further study.

Keywords: Delayed graft function, Warm ischemic time, Cold ischemic time, Hospitalization

Background

Over the last two decades cold ischemic time (CIT) has been found to be an important independent risk factor for delayed graft function (DGF) in deceased donor kidney transplantation [1-4]. DGF is also associated with inferior graft survival and function [1-3]. All supporting evidence for the adverse events associated with DGF is derived from cohort studies. DFG likely results in longer stays in hospital and additional resource use (dialysis). Surprisingly there are few studies examining the effects of warm ischemic time (WIT) on DGF [5,6]. The importance of WIT has become a hot area of research in Urology especially with greater use of partial nephrectomy for cancer surgery [7,8]. There is also recent information that long WITs may reduce graft survival in live donor kidney transplantation [9].

By definition there are two WITs. The first relates to organ procurement and the second relates to the vascular anastomosis time (AT) in the recipient. The purpose of this study is to examine the second warm time or the AT during the recipient operation and examine whether this contributes to delayed graft function.

Methods

Consecutive deceased donor organ transplant recipients of solitary kidneys were identified in the Program database from January 2006 until August 2012. Combined organ and living donor transplant procedures were excluded. Data were extracted retrospectively from our
electronic health records, which came into existence in January 2006. Approval for this retrospective study was obtained from our institution’s ethics board.

All patients received induction with either basiliximab or antithymocyte globulin (ATG) and all received preoperative methylprednisolone (500 mg) intravenously. ATG was reserved for those who were repeat transplant recipients, those who received organs from deceased cardiac donors, and those with a calculated panel reactive antibody level >20%. Post-operative patients received oral tacrolimus, mycophenolate, and prednisone. All patients were treated with sulfamethoxazole-trimethoprim for pneumocystis prophylaxis and oral ranitidine for gastrointestinal ulcer prophylaxis. For patients on a proton pump inhibitor pre transplant, this agent was continued in place of the ranitidine. Cytomegalovirus (CMV) prophylaxis with valganciclovir was used in CMV-negative recipients receiving CMV-positive organs and CMV-positive recipients receiving ATG. All patients received subcutaneous heparin for venous thromboembolism prophylaxis.

Data extracted included recipient age, gender, body mass index (BMI), weight, calculated panel reactive antibodies (cPRA), human leukocyte antigen mismatch (HLA MM), cause of end-stage renal disease (ESRD), diabetes mellitus status at transplantation, duration of dialysis pretransplantation, blood transfusion pre transplantation, CIT, donor cause of death, donor age, gender, BMI, weight, terminal creatinine, and donation after circulatory death (DCD). Variables that were significant at $P < 0.1$ in a univariable analysis were included into the multivariable models. Overall significance was assumed at the $P = 0.05$ level (two-sided). Statistical analysis was performed using IBM SPSS Statistics software version 20.0 (New York, NY, USA).

### Results

In this retrospective cohort of 307 consecutive deceased donor solitary kidney transplant recipients in the database, ATs were available in 298 subjects. The nine (2.9%) missing subjects were excluded from the analysis. Of the remaining 298 patients, 56 (18.8%) had DGF. Characteristics of the patients are shown in Table 1. Figure 1

| Table 1 Population descriptors (n = 298) |
|-----------------------------------------|
| Descriptor                              | Value                  |
| Age, years, mean ± SD                   | 51 ± 13                |
| Gender, male, number (%)                | 181 (60.7%)            |
| Regraft, number (%)                     | 29 (9.0%)              |
| Diabetes mellitus, number (%)           | 69 (23.0%)             |
| End-stage renal disease                 |                        |
| Polycystic kidney disease, number (%)   | 55 (19%)               |
| Glomerulonephritis, number (%)          | 85 (29%)               |
| Diabetes mellitus, number (%)           | 50 (17%)               |
| Recipient weight, kg, mean ± SD         | 78 ± 17                |
| Recipient body mass index, kg/m², mean ± SD | 28 ± 5               |
| Blood transfusion, never, number (%)    | 205 (66%)              |
| Human leukocyte antigen mismatch, mean ± SD | 4.1 ± 1.4             |
| Dialysis duration, days, median (IQR)   | 819 (419, 1462)        |
| Calculated panel reactive antibody, %, median (IQR) | 0 (0, 8)             |
| Cold ischemic time, hours, median (IQR) | 12 (8, 16)            |
| Anastomosis time, minutes, median (IQR) | 30 (24, 45)            |
| Donation after circulatory death, number (%) | 16 (5.4%)            |
| Donor cause of death                    |                        |
| Anoxia, number (%)                      | 24 (8%)                |
| Cerebrovascular event, number (%)       | 189 (62%)              |
| Donor age years, mean ± SD              | 47 ± 17                |
| Donor male, number (%)                  | 135 (45%)              |
| Donor weight kg, mean ± SD              | 76 ± 19                |
| Donor body mass index kg/m², mean ± SD  | 27 ± 6                 |
| Donor creatinine μmol/L, mean ± SD      | 67 ± 31                |
| Delayed graft function, number (%)      | 56 (18.8%)             |
| Days in hospital, days, median (IQR)    | 9 (7, 14)              |
| Creatinine day 7, μmol/L, median (IQR)  | 168 (125, 314)         |
shows the distribution of ATs in the study. Since the distribution is significantly skewed the median is the more appropriate descriptor (median 30 minutes, IQR 24, 45 minutes). Figure 2 (box and whisker plot) shows that patients experiencing DGF had significantly longer ATs. AT was independently associated with DGF in the binary logistic regression analysis (odds ratio (OR) 1.037 per minute, 95% CI 1.016, 1.057, \( P = 0.001 \)). Other variables that were significant in the model were DCD (OR 11, 95% CI, 3.3, 34, \( P < 0.001 \)), donor BMI (OR 1.06 per kg/m\(^2\), 95% CI, 1.003, 1.12, \( P = 0.041 \)), and dialysis duration pre transplantation (OR 1.001 per day, 95% CI, 1.000, 1.002, \( P < 0.001 \)). Diabetes mellitus, donor sex, and repeat transplants were included in the multivariable model (\( P < 0.10 \) in the univariable model) but were not independently significant. In a separate multivariable model, an AT >29 minutes was associated with an increased risk of DGF (OR 3.5, 95% CI 1.6, 7.3, \( P = 0.001 \)).

The median days in hospital was 9 (IQR 7, 14 days). Figure 3 shows the relationship between AT and days in hospital (\( r = 0.32, P < 0.001 \)). AT (B coefficient 0.20 days per minute AT, 95% CI, 0.13, 0.27, \( P < 0.001 \)) was associated with longer stays in hospital in the multivariable linear regression analysis model. Therefore every 5 minutes of longer AT was associated with one extra day in hospital. Other significant covariables were DCD (B coefficient 11 days, 95% CI, 7, 16, \( P < 0.001 \)), dialysis duration pre transplantation (B coefficient 0.002 days per day, 95% CI, 0.001, 0.003, \( P = 0.001 \)) and donor sex (B coefficient 3.5 days for male, 95% CI, 1.4, 5.6, \( P = 0.001 \)). Recipient age, diabetes mellitus, donor age, and repeat transplants were included in the multivariable model (\( P < 0.10 \) in the univariable model) but were not independently significant. In a separate multivariable model an AT >29 minutes was associated with 3.8 days (95% CI 1.6, 6.0, \( P < 0.001 \)) longer in hospital.

The median serum creatinine on day 7 was 168 μmol/L (IQR 125, 314). AT (B coefficient 4.0 μmol/L per minute AT, 95% CI, 2.1, 5.0, \( P < 0.001 \)) was associated with higher day 7 creatinine levels. Other significant factors in the multivariable model were donor terminal serum creatinine (B coefficient 1.8 μmol/L per μmol/L, 95% CI, 1.04, 2.5, \( P = 0.001 \)) and DCD (B coefficient 312 μmol/L, 95% CI, 213, 411, \( P < 0.001 \)). Recipient sex, recipient weight, diabetes mellitus, and donor BMI were included in the multivariable model (\( P < 0.10 \) in the univariable model) but were not independently significant. An AT >29 minutes was associated with a higher creatinine at day 7 (B coefficient 75 μmol/L, 95% CI, 30, 121, \( P < 0.001 \)).

**Discussion**

AT may be an underappreciated but modifiable variable in dictating use of hospital resources. There were significant increases in both need for dialysis and length of
stay at this center with increased AT. An AT >29 minutes was associated with particularly worse outcomes. The study also shows that the serum creatinine was higher 7 days after transplantation with longer ATs, consistent with slower recovery of kidney function.

Surprisingly few studies have examined WIT as a covariate in studies of delayed graft function or survival. In a normogram to predict DGF, Irish et al. identified WIT as a significant variable but the overall effect was small [6]. The reasons why this study demonstrated a greater effect with AT is not clear since most of the variables included in the DGF normogram were also considered in this study. The DGF normogram was examined in one center and found to have limited accuracy [10]. In Grossberg’s study, transplant recipients with and without DGF had DGF normogram scores that were not significantly different. DGF as defined in this paper and by Irish et al. may not be the best predictor of later graft function [6]. Moore et al. found the DGF normogram to be less predictive of later kidney function at 1 year than other measures. Creatinine at day 5, and to a similar extent creatinine at day 7 post kidney transplantation were predictive of later function at 1 year [11].

In a single center study from the Netherlands of over 1,000 solitary deceased and live donor kidney recipients transplanted between 1981 and July 2000, neither the first nor second warm time was significantly associated with outcomes [5]. However, a more recent study from the same country in live donor kidney recipients found that prolonged warm times (>45 minutes) were associated with a 3-fold higher risk of graft loss [9]. Most research has focused on CIT, preservation solutions or the use of machine perfusion pumps to avoid reperfusion injury and DGF and these do not directly address the AT [12]. Even the large recent randomized trials such as the international trial on the use of machine pumps to prevent DGF or recent cohort trials examining risk factors for DGF did not include WIT in their analysis as a potential confounder [4,13].

There is a growing body of evidence in the urologic literature that WIT in partial nephrectomy is detrimental and there should be no reason why these adverse effects associated with WIT would be irrelevant in kidney transplantation [7,8]. Consistent with a debate in the urologic literature, there may not be a threshold effect and every minute may count [7]. In a recent publication examining the effect of WIT in a cohort undergoing partial nephrectomy of a solitary kidney, it was found that each minute of warm ischemia was associated with a 6% increased risk of acute kidney injury and 4% increased risk of new onset ESRD [14]. Clearly some prolonged times are unavoidable, especially in those with multiple vessels that have significant atherosclerosis or that require re-exploration. Minimizing warm time by back table surgery would also be recommended. Recognizing that pursuing speed should not impact the quality of anastomosis, increasing the awareness of this issue amongst surgeons may be sufficient to improve ATs. It might be necessary to include more simulation and skills laboratory time for trainees to gain adequate expertise prior to performing anastomoses in clinical practice.

There are significant limitations to this study. This was retrospective and single center, and long-term effects on graft or patient survival were not examined because of the low event rates (98% patient and 96% graft survival at one year). Given the growing body of clinical and experimental evidence that acute ischemic kidney injury is associated with later progressive chronic kidney disease in the non-transplant setting, a more detailed analysis of larger registries should be undertaken [15-17]. If an association between greater AT and worse long-term graft survival is observed, then there would be even greater interest in testing early pharmacologic interventions or devising better means to minimize the warm time. A number of strategies have been tested experimentally and in ongoing clinical trials (clinicaltrials.gov) to prevent ischemia-reperfusion injury. In addition to previously studied interventions such as spironolactone in animals, therapies to prevent leukocyte recruitment and inflammation (NCT00248040), cytokine damage (NCT00298168) and the use of small interfering RNAs (NCT00802347) to prevent renal injury may hold promise [17,18].

In summary, this study highlights the importance of the anastomosis time in kidney transplantation on short-term outcomes that may impact long-term outcomes. More significantly this event may be modifiable, or potentially an area and time for other types of intervention.

Abbreviations
AT: Anastomosis time; ATG: Antithymocyte globulin; BMI: Body mass index; CIT: Cold ischemic time; CMV: Cytomegalovirus; CPRA: Calculated panel-reactive antibody; DCD: Donation after circulatory death; DGF: Delayed graft function; DIH: Days in hospital; ESRD: End-stage renal disease; GN: Glomerulonephritis; HLA MM: Human leukocyte antigen mismatch; PCKD: Polycystic kidney disease; OR: Odds ratio; WIT: Warm ischemic time; IQR: Interquartile range.

Competing interests
None of the authors declare any funding for this study or competing interests. However JL, IA and BK are participating in a contract research study with Quark Pharmaceuticals.

Authors’ contributions
KM collected data and helped draft the manuscript. JL co-designed the study, and critically appraised the analysis and significance. IA critically appraised the analysis and significance. BK co-designed the study, collected and analyzed data, and assisted in manuscript preparation. All authors read and approved the final manuscript.

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