UNIVERSITY^{OF} BIRMINGHAM

University of Birmingham Research at Birmingham

Deceased Donor Characteristics and Kidney Transplant Outcomes

Sharif, Adnan

DOI:

10.3389/ti.2022.10482

License:

Creative Commons: Attribution (CC BY)

Document Version

Publisher's PDF, also known as Version of record

Citation for published version (Harvard):

Sharif, A 2022, 'Deceased Donor Characteristics and Kidney Transplant Outcomes', *Transplant international*, vol. 35, 10482. https://doi.org/10.3389/ti.2022.10482

Link to publication on Research at Birmingham portal

General rights

Unless a licence is specified above, all rights (including copyright and moral rights) in this document are retained by the authors and/or the copyright holders. The express permission of the copyright holder must be obtained for any use of this material other than for purposes permitted by law.

- •Users may freely distribute the URL that is used to identify this publication.
- •Users may download and/or print one copy of the publication from the University of Birmingham research portal for the purpose of private study or non-commercial research.
- •User may use extracts from the document in line with the concept of 'fair dealing' under the Copyright, Designs and Patents Act 1988 (?)
- •Users may not further distribute the material nor use it for the purposes of commercial gain.

Where a licence is displayed above, please note the terms and conditions of the licence govern your use of this document.

When citing, please reference the published version.

Take down policy

While the University of Birmingham exercises care and attention in making items available there are rare occasions when an item has been uploaded in error or has been deemed to be commercially or otherwise sensitive.

If you believe that this is the case for this document, please contact UBIRA@lists.bham.ac.uk providing details and we will remove access to the work immediately and investigate.

Download date: 05. May. 2024





Deceased Donor Characteristics and Kidney Transplant Outcomes

Adnan Sharif 1,2*

¹Department of Nephrology and Transplantation, University Hospitals Birmingham, Queen Elizabeth Hospital, Birmingham, United Kingdom, ²Institute of Immunology and Immunotherapy, University of Birmingham, Birmingham, United Kingdom

Kidney transplantation is the therapy of choice for people living with kidney failure who are suitable for surgery. However, the disparity between supply versus demand for organs means many either die or are removed from the waiting-list before receiving a kidney allograft. Reducing unnecessary discard of deceased donor kidneys is important to maximize utilization of a scarce and valuable resource but requires nuanced decisionmaking. Accepting kidneys from deceased donors with heterogenous characteristics for waitlisted kidney transplant candidates, often in the context of time-pressured decisionmaking, requires an understanding of the association between donor characteristics and kidney transplant outcomes. Deceased donor clinical factors can impact patient and/or kidney allograft survival but risk-versus-benefit deliberation must be balanced against the morbidity and mortality associated with remaining on the waiting-list. In this article, the association between deceased kidney donor characteristics and post kidney transplant outcomes for the recipient are reviewed. While translating this evidence to individual kidney transplant candidates is a challenge, emerging strategies to improve this process will be discussed. Fundamentally, tools and guidelines to inform decision-making when considering deceased donor kidney offers will be valuable to both professionals and patients.

Keywords: mortality, kidney transplant, graft loss, deceased donor, discard, kidney failure

INTRODUCTION

OPEN ACCESS

*Correspondence:

Adnan Sharif adnan.sharif@uhb.nhs.uk

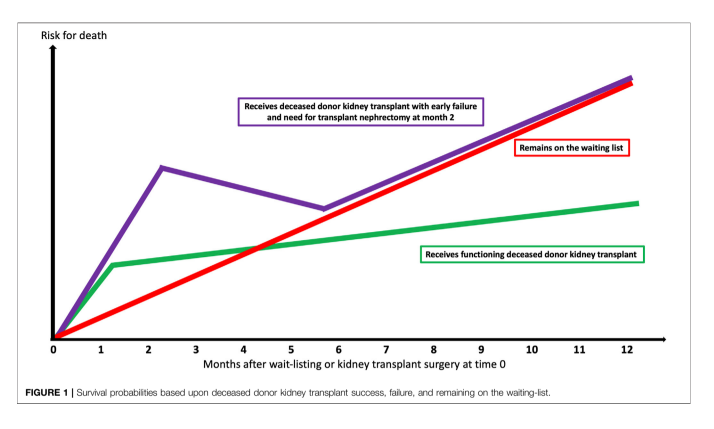
Received: 01 March 2022 Accepted: 25 July 2022 Published: 25 August 2022

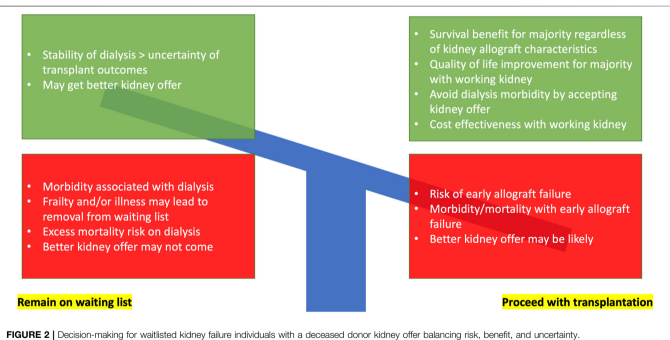
Citation

Sharif A (2022) Deceased Donor Characteristics and Kidney Transplant Outcomes. Transpl Int 35:10482. doi: 10.3389/ti.2022.10482 Kidney transplantation is the treatment modality of choice for kidney failure patients deemed fit enough for surgery. While successful kidney transplantation lowers both cardiovascular (1) and all-cause mortality (2,3), and provides better quality of life and cost-effectiveness in most scenarios (4), kidney transplant failure and return to dialysis is associated with heightened risk for mortality over-and-above transplant-naïve waitlisted dialysis patients (see **Figure 1**) (5,6). Therefore, personalizing use of deceased donors for individual waitlisted kidney transplant candidates at the most appropriate time is challenging (see **Figure 2**) (7).

These factors partly explain unnecessary kidney discards. Mohan et al. observed 17.3% of procured kidneys in the United States between 2000–2015 were discarded, despite partner kidneys of unilaterally discarded kidneys experiencing 1-year death-censored graft survival rates >90% (8). Over 80% of kidney discard rates can be explained by the broadening donor pool and unexplained residual factors (9). Organ discard rates in European countries are lower than the United States (10), although donor characteristics differ (e.g., more opioid-related deaths in the United States) (11). If deceased donor kidney acceptance in the United States mirrored the French model (discard rate 17.9% versus 9.1% respectively, p < 0.001), then utilization of discarded kidneys

1





(n=17,435) could generate 132,445 allograft life-years (12). This is important as declined kidney offers are not benign events. Husain et al., in a cohort study analyzing 280,041 wait-listed kidney transplant candidates in the United States, observed approximately 30% of candidates receiving at least one deceased donor offer declined on their behalf eventually died

or were removed from the waiting-list before receiving a kidney allograft (13). Apart from clinical benefits, transplantation using kidneys of any quality is cost-effective versus remaining on the waiting-list (14).

In view of increasing marginality of kidneys procured from deceased donors (15,16), which contributes to sub-optimal organ

utilization, informed decision-making to accept kidney offers for individual kidney transplant candidates must be the objective. While organ allocation systems continue to evolve (17), which impact upon utilization, the aim of this review is to summarize published evidence regarding kidney transplant outcomes associated with deceased kidney donor characteristics. Translating such data into decision-making pathways is a clinical challenge and emerging ways to foster better organ utilization are discussed.

DONOR CLINICAL FACTORS

Expanded Criteria Donor

An ECD is one who, at the time of death, is aged ≥60 years or aged 50–59 years with any two the following three criteria: 1) cause of death is cerebrovascular accident, 2) pre-existing history of systemic hypertension, and 3) terminal serum creatinine >1.5 mg/dl. The criteria for defining ECD was based on the presence of variables that historically increased the risk for graft failure by 70% compared with a standard criteria donor (SCD) kidney (18).

Previous systematic reviews suggested ECD kidneys should not be offered to younger (aged <40 years) kidney transplant candidates or those undergoing re-transplantation (19). ECD kidneys may be better prioritized for older recipients by ignoring immunology-based allocation. Using this strategy, the Eurotransplant Senior program have shown favorable 5-year outcomes using ECD kidneys in older recipients (20).

However, recent analyses support broadening access with careful risk stratification. Querard et al. conducted a systematic review and meta-analysis of 32 studies comparing survival outcomes between SCD and ECD kidneys (21). Pooled 5-year patient survival probabilities were 78.4% versus 86.4% in ECD versus SCD recipients respectively. A significant difference in mortality was observed comparing North American and European studies, with 5-year pooled patient survival closer in European studies (ECD versus SCD; 85.3% versus 90.3% respectively) than in North American studies (ECD versus SCD, 73.4% versus 83.6% respectively). The corresponding pooled RR was estimated at 1.50 (95% CI 0.50–3.43) for the European studies versus 1.62 (95% CI 1.18–2.22) for the North American studies. Similar effect sizes were seen with regards to death-censored graft survival.

ECD kidney allograft survival may be improved in the absence of circulating donor-specific antibody (p < 0.001) and CIT <12 h (p = 0.030) according to a French study (22). Optimal utilization of ECD kidneys may also be stratified by recipient age, with studies suggesting recipients aged ≥ 60 years (23) or ≥ 65 years (24) be prioritized. However, while a 1.75-fold (95% CI 1.53–2.00, p < 0.0001) increased risk for graft failure using ECD versus SCD kidneys was observed in one study, population-average effect using propensity scores with 10-year follow-up highlighted a minimal absolute effect of only 8 months (95% CI 2–14 months) quicker time to graft failure attributed to ECD kidneys (24). Therefore, the absolute risk difference between SCD and ECD

kidneys in the long-term may be marginal when compared to remaining on the waiting-list.

Donation After Cardiac Death

DCD refers to a donor who does not meet the criteria for donation after brain death (DBD) but in whom cardiac standstill or cessation of cardiac function occurred before organs were procured, with cessation of cardiac function initiated deliberately (controlled) or occurring spontaneously (uncontrolled) (18).

Data from the United Kingdom, examining outcomes in adult recipients receiving a deceased donor kidney transplant between 2000-2007, compared survival outcomes between 8,289 DBD kidneys and 845 DCD kidneys (25). Despite increased rates of delayed graft function (DGF) after DCD kidney transplantation, first-time recipients of DCD kidneys (n = 739) or DBD kidneys (n = 6.759) showed no difference in 5-year graft survival (HR 1.01, 95% CI 0.83–1.19, p = 0.97). Increasing donor or recipient age, repeat transplantation, and CIT >12 h were associated with worse graft survival for recipients of DCD kidneys. Subsequent analyzes demonstrate equivalent 5-year patient survival or 10year death-censored graft survival comparing DCD versus DBD kidneys (26). Prolonged CIT (>24 h versus <12 h) was associated with poorer graft survival for DCD versus DBD kidneys in cohorts from the United Kingdom and United States (27). The rate of primary nonfunction for both DCD and DBD kidneys was low (3.1 and 2.5% respectively) and not significantly different (risk-adjusted OR 1.18, 95% CI 0.9-1.5, p = 0.21) (28). These reassuring long-term outcomes suggest DCD kidneys of any age should be actively considered for all kidney transplant candidates, if logistics and resources can facilitate timely surgery to avoid prolonged CIT.

Studies report heterogeneous outcomes for ECD-DCD kidneys. Locke et al., exploring data from the United States between 1993-2005, observed donor age was associated with increased graft failure risk, although graft survival was similar between ECD-DBD and >50-year old DCD kidneys (29). Singh et al., analyzing data from the United States including 562 ECD-DCD kidneys, showed slightly increased risk for graft loss in recipients receiving DCD versus non-DCD kidneys, which was not significantly modified by ECD status (30). Across a number of studies, ECD-DCD kidneys report acceptable 3-year deathcensored graft survival rates between 70%-90%, which are inferior to SCD-DCD kidneys but not ECD-DBD kidneys (26-32). However, Montero et al. demonstrate how important selecting the most appropriate donor-recipient combination is in a recently published risk modelling study (33). In their multicenter cohort study, mortality risk for the highest riskstratification group receiving ECD-DCD kidneys was significant. Although survival was better post-transplantation compared to remaining waitlisted, it raises a level of caution in decision making when dealing with donor-recipient extremes. Therefore, use of ECD-DCD kidneys is acceptable for select waitlisted kidney transplant candidates when carefully balanced against their mortality risk without transplantation and quality of life considerations.

Kidney Donor Risk Index

The KDRI is a risk quantification score developed in 2009 by Rao et al. using data from the United States between 1995–2005 containing 14 donor- or transplant-specific variables (34). A recent re-evaluation using more contemporary United States data reported the original KDRI remains robust for discrimination and predictive accuracy for graft failure (35). KDRI has been implemented into allocation policy within the United States, with low KDRI (i.e., better quality) kidneys preferentially allocated to kidney transplant candidates with the greatest expected longevity (36).

A pan European study including 24,177 adult kidney transplant recipients demonstrated an increase in KDRI by 1.3% annually, from 1.31 (IQR 1.08–1.63) in 2005 to 1.47 (IQR 1.16–1.90) in 2015, driven by increased donor age, hypertension, and use of DCD kidneys (16). No difference was observed in 5-year patient or allograft survival outcomes, with survival probabilities improving over time for the highest KDRI kidneys. Within any given KDRI interval, although ECD kidneys have higher rates of discard and graft failure risk, the ECD categorization does not confer additional risk of discard or graft failure compared with SCD kidneys within the same KDRI interval (37).

However, caution should be exercised with the KDRI. It contains components which can increase the risk quantification score but now demonstrate comparable outcomes (e.g., DCD). Translatability of the KDRI to population cohorts outside the United States may not be valid (38,39). Due to disparate survival outcomes observed for kidney failure patients treated with dialysis (40,41) versus kidney transplantation (42) in the United States versus elsewhere, and different utilization of deceased donors (e.g., greater use of older and DCD kidneys in the United Kingdom versus the United States for example) (43), generalizability may be invalid.

Donor Age

Donor age has the strongest independent association with longterm kidney transplant outcomes (44). These accepted deleterious effects justify donor age being a component of the KDRI risk score but also the fundamental stratification for ECD classification. Donor age is one of the most frequent explanations for organ discard (8), despite an increasing proportion of deceased organs over time being procured from older donors (16). While many studies dichotomize at an arbitrary cut-off donor age of 60 years, deleterious effects for kidney transplant recipients may start earlier. Keith et al. analyzed data from the United States between 1990 and 1997 and observed adjusted 10-year patient survival starts to drop with deceased donor ages 36-40 years (45). There is a strong interaction between donor and recipient age, with additive detrimental effect on allograft survival with a combination of older kidneys into older recipients (46), although many allocation systems prioritize on this like-for-like basis.

Some centers consider dual versus single kidney transplants using older kidneys. However, when using donors aged ≥60 years, no graft survival advantage at 5-year was observed comparing dual versus single kidney transplantation in an analysis from the

United Kingdom between 2005–2017 (adjusted HR 0.81, 95% CI 0.59–1.12). However, dual kidney transplantation did result in slightly higher 1-year estimated glomerular filtration rate [eGFR] (40 versus 36 ml/min/1.73 m² respectively, p < 0.001) (47).

Donor Ethnicity

Non-white ethnicity demonstrates conflicting associations with kidney transplantation outcomes. Pisavadia et al., exploring data from the United Kingdom between 2003–2015, observed higher risk for graft loss with south Asian (HR 1.38, 95% CI 1.12–1.70, p=0.003) and Black (HR 1.66, 95% CI 1.30–2.11, p<0.001) donated kidneys independent of recipient ethnicity, with no survival advantage from donor-recipient ethnicity matching (48). Locke et al., exploring data from the United States between 1993–2006, suggested DCD kidneys from Black donors, but not DBD kidneys, were associated with better patient and graft survival for Black recipients (49). This contrasts with evidence from registry data that kidneys donated by ethnic minorities (especially Black individuals) are associated with poorer graft survival for any kidney transplant recipient (50,51).

However, using ethnicity for risk stratification of deceased donors is questionable. Ethnicity is not a reliable proxy for genetic difference between individuals (52). While incorporating ethnicity into clinical decision-making can be considered a form of personalized medicine, it may not add additional value. For example, Chong et al., in an analysis of data from the United States between 2000–2017, demonstrated removal of donor ethnicity from KDRI calculations makes negligible difference to patient and kidney allograft survival, strongly advocating for removal of donor ethnicity as a risk factor (53).

Donor Body Mass Index

In a population cohort study from the United Kingdom, Arshad et al. observed an independent association between donor BMI and delayed graft function (54), with risk increased in recipients of kidneys from overweight (Odds Ratio [OR] 1.12, 95% CI 1.00-1.23, p = 0.022), obese (OR 1.23, 95% CI 1.08–1.39, p <0.001), and morbidly obese (OR 1.38, 95% CI 1.16-1.63, p < 0.001) donors when compared to normal donor BMI group. However, donor BMI did not influence long-term patient or graft survival. This is corroborated with data from the United States. In a study of 6,932 recipients of DCD kidneys in the United States, Ortiz et al. reported donors with a BMI between 30.0–34.9 kg/m² incurred 1.77-fold increased odds of developing DGF, with similar odds for DGF in donors with a BMI between $35.0-39.9 \text{ kg/m}^2$ (OR 1.78, p < 0.001) (55). However, only DCD kidneys from donors with a BMI >45.0 kg/m² were associated with an increased risk of death-censored graft failure (adjusted HR 1.84, 95% CI 1.23–2.74, p < 0.001) relative to a normal donor BMI category.

Donor Size

The influence of donor-to-recipient size matching has shown conflicting results. Arshad et al., exploring data from the United Kingdom between 2003–2015, showed no association between donor-to-recipient size match difference and risk for

DGF or death-censored graft survival (56). Donor-to-recipient difference in body weight was associated with higher 12-month creatinine in large recipients receiving small donor kidneys. Increased mortality was observed in recipients receiving larger kidneys (HR 1·21, 95% CI 1.05–1.40 p=0.009), which conflicts with other population-cohort studies that show inferior long-term patient and graft survival associated with larger recipients receiving smaller donor kidneys (57-59). Some show negative effects of size mismatch (large kidney into small recipient) only in the context of ECD kidneys (58) or male recipients of female kidneys (59).

Donor Acute Kidney Injury

The relationship between donor AKI and kidney transplant outcomes has been reviewed by Koyawala and Parikh (60). In total, 37 studies were identified comparing transplant outcomes between kidneys with versus without donor AKI. Donor AKI was associated with DGF, with prolonged nights in hospitals and additional attributed costs. In a separate meta-analysis of 14 cohort studies exploring 15,345 donors, Zheng et al. estimate the relative risk of DGF to be 1.76 (95% CI 1.52–2.04) for recipients of kidneys with versus without donor AKI (61).

No association is seen between donor AKI and risk for rejection after 6 months or 1 year, either in a review of published studies (60) or meta-analysis of empirical data (RR 0.87, 95% CI 0.66–1.15) (61). No association was seen between donor AKI and graft function (60).

From a graft survival perspective, donor AKI was not associated with graft failure in 25/29 studies (60). However, some studies provide more granular insight. Botha et al. analyzed 11,219 transplanted kidnevs United Kingdom, comprising 1,869 (17%) with AKI (62). While 1-year graft survival difference was statistically significant comparing AKI versus non-AKI donor kidneys, the numerical difference was clinically insignificant (89% versus 91% respectively, p = 0.02). DGF rates increased with severity of AKI (no AKI = 28%, AKI stage 1 = 35%, AKI stage 2 = 43%, AKI stage 3 = 55%, p < 0.005). Primary nonfunction rates were higher with donor AKI stage 3 versus no AKI kidneys (9% versus 4%, p = 0.04) and graft function was lower among donor AKI kidneys (OR 1.25, 95% CI 1.08-1.31, p < 0.005). This study differed from other cohorts due to its higher sample size, with a larger proportion of donor kidneys with severe AKI and donation after circulatory death, meaning this study may be better powered to observe differences in outcomes among donor kidneys with higher levels of injury. Other studies observed higher rates of graft failure only among a sub-select of studies using ECD donor kidneys with AKI (63,64).

Donor AKI is more acceptable with high versus low quality kidneys. Single center outcomes using donors with both AKI (comparing advanced stages 2–3 versus 0–1) and high KDPI (\geq 85%) demonstrated more DGF (71% versus 37% respectively, p < 0.001), more primary nonfunction (5.3% versus 0.6% respectively, p = 0.02), no difference in eGFR in ml/min/1.73² (44 versus 46 respectively, p = 0.42) and lower 1-year death-

censored graft failure 14.5% versus 3.5% for AKI 2-3 versus AKI 0-1 high KDPI kidneys respectively (HR 2.40, 95% CI 1.24–4.63, p=0.01) (65).

Donor Diabetes

Cohen et al. studied survival outcomes for kidney transplant patients receiving diabetic versus non-diabetic kidney allografts in the United States between 1994-2013 (66). Recipients of diabetic donor kidneys had higher rates of all-cause allograft failure (HR 1.21, 95% CI 1.16-1.26) and death (HR 1.19, 95% CI 1.13-1.24) compared to receiving non-diabetic donor kidneys. Allograft survival was worse for younger (≤45 years of age) versus older recipients of diabetic donor kidneys, but no difference was observed in patient survival. Due to a significant interaction between donor and recipient diabetes status (with diabetic recipients receiving diabetic donor kidneys having the worst patient and allograft survival), paired analyzes of mate-kidneys from the same donor were undertaken where one recipient was diabetic and the other non-diabetic. In this analysis, diabetic recipients had significantly higher risk of allograft failure (HR 1.27, 95% CI 1.05-1.53) and death (HR 1.53, 95% CI 1.22-1.93) compared to non-diabetic recipients. Diabetic recipients of nondiabetic donor kidneys and non-diabetic recipients of diabetic donor kidneys had similar rates of all-cause allograft survival.

The critical question is whether waitlisted patients should accept diabetic donor kidneys versus waiting for better kidneys. Cohen et al. compared survival benefits of kidney transplantation using diabetic donor kidneys versus remaining on the waiting-list in the United States between 1994-2015 (67). They observed recipients of diabetic donor kidneys had lower mortality compared with remaining on the waiting-list and/or transplantation later with a non-diabetic donor kidney (adjusted HR 0.91, 95% CI 0.84-0.98). Although recipients of non-diabetic donor kidneys with high KDPI scores had lower mortality risk (adjusted HR 0.86, 95% CI 0.81-0.91), recipients of diabetic donor kidneys with similar high KDPI scores showed no survival difference (adjusted HR 1.09, 95% CI 0.97-1.22). Younger waitlisted patients (aged <40 years) had no survival benefit from transplantation with diabetic donor kidneys, while diabetic patients with longer waiting-list times attained the greatest survival benefit.

Donor Hypertension

Donor hypertension is increasing in prevalence and observed in nearly a third of deceased donors (16). Altheaby et al., in a systematic review and meta-analysis, identified 15 studies published between 1963–2014 exploring the association between donor hypertension and kidney transplant outcomes (68). Pooled risk ratios (RR) demonstrate donor hypertension is associated with kidney allograft failure (RR 1.31, 95% CI 1.06-1.63, p=0.014) but not mortality (RR 0.996, 95% CI 0.652-1.519, p=0.984).

Donor Smoking

Donor smoking and kidney transplant outcome associations are unclear. Lin et al. explored data from the United States between 1994–1999 and observed smoking history among deceased

kidney donors was associated with increased transplant recipient risk for death and graft loss (69). However, Gillott et al. explored data from the United Kingdom between 2001–2013 and observed no association between donor smoking and allograft survival for kidney transplant recipients, although an association with mortality was observed (adjusted HR 1.16, 95% CI 1.03–1.29, p=0.011) (70).

Donor Cause of Death

Death by cerebrovascular accident remains the commonest cause of death, varying little across Europe between 2005–2015 (16), and contributes to ECD classification for donors aged ≥50 years. Few studies have explored the impact of cause of donor death and recipient outcomes, although cause of death that can result in disease transmission has been of greater concern.

Donor-Derived Disease Transmission

Risk for donor-derived disease transmission (defined as either infection or malignancy) leading to morbidity or mortality occurred in only 0.96% of all solid organ transplantation in the United States (71). Increased risk for disease transmission (IRD) kidneys tend to be better quality (defined as lower KDPI scores) and associated with survival benefits. For example, Bowring et al. analyzed data in the United States between 2010-2014 and demonstrated: 1) recipients who declined IRD kidneys and subsequently received non-IRD kidneys accepted a higher median KDPI (21 versus 52 respectively); and 2) after a short risk period in the first 30 days following IRD acceptance (adjusted HR 2.06, 95% CI 1.22–3.49, p = 0.008) (absolute mortality 0.8% versus 0.4%), those who accepted IRDs had lower risk of death 1-6 months (adjusted HR 0.67, 95% CI 0.50-0.90, p =0.006) and beyond 6 months (adjusted HR 0.52, 95% CI 0.46-0.58, p < 0.001) (72).

However, most cases of donor-disease transmission will occur in clinically covert donors. For example, in a systematic review of published literature, donors with a history of cancer or an ongoing malignancy contributed to disease transmission in only 17.1% (n=32) of cases (73). Using data from the United Kingdom, it is estimated the risk of transmitting cancer from a donor not known to have a malignancy is very low at 1 in 2,000 (0.05%) (74).

Donor Increased Risk Behavior

Increased risk behavior (IRB) among deceased donors can be classed as intravenous drug use (IVDU), imprisonment, or highrisk sexual behavior. Trotter et al, analyzing data from the United Kingdom between 2003–2015, studied the outcomes associated with use of IRB deceased donor kidneys (75). Donors with IRB provided 1,091 organs for transplantation (including 624 kidneys) and transplant outcomes were similar in recipients of organs from donors with versus without IRB. Only three cases of unexpected hepatitis C virus transmission were identified, all from an active IVDU donor who was hepatitis C virus seronegative at time of donation but was found to be viremic on retrospective testing. National registry data and single

center studies from the United States have shown excellent outcomes and minimal risk associated with using deceased donor kidneys from IVDU individuals (76-79). High decline rates observed with IRB kidneys (75,76) suggest a valuable but underutilized resource due to a subjective perception of heightened risk for kidney transplant recipients not supported by objective evidence.

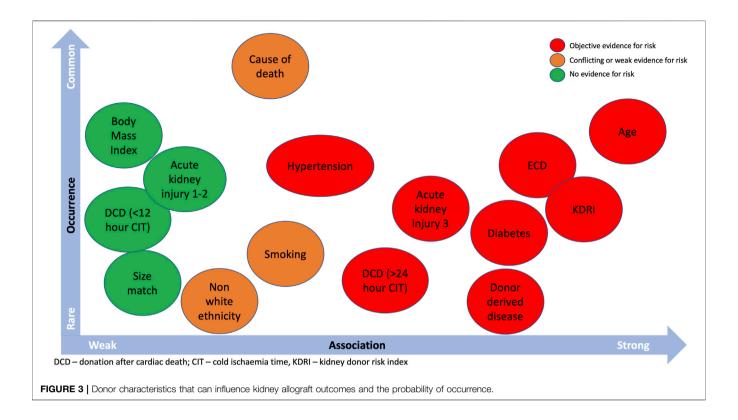
DONOR HISTOPATHOLOGY

The benefit of obtaining donor histopathology to guide kidney utilization is unclear. In a systematic review of published evidence, Wang et al. combined empirical evidence from 47 studies (80). In these retrospective studies exploring heterogenous histopathological criteria, no semi-quantitative scoring system was conclusively associated with posttransplant outcomes including DGF, grant function, and/or graft failure. This may relate to weak inter-observer correlation and variability between pathologists, which could be improved using a dedicated pool of specialist pathologists (81). Preimplantation biopsy analysis may be useful in a subset of deceased donor kidneys where chronic injury is prevalent like ECD kidneys. Based upon this rationale, the PreImplantation Trial of Histopathology In renal Allografts (PITHIA) study is an open, multicenter, stepped-wedge cluster randomized study, that involved all UK adult kidney transplant centres (82). Using a pool of dedicated pathologists, it will explore whether a national, 24-h, digital histopathology service improves organ utilization from deceased donors aged 60 years and over. The results from this national study are awaited but should provide clarity regarding the value of pre-implantation donor histopathology.

DECISION CHALLENGES

Translating this evidence to nuanced decision-making is the big challenge. No guidelines or recommendations exist to support this process, which is difficult considering the nature of available data. For example, most deceased donors will have a combination rather than individual clinical factors (see Figure 3). This requires individualized considerations of population-level data which are not amenable to simple flowcharts. Organ utilization has behavioral components, from both patients and professionals, that will influence decision-making, and it is important every kidney transplant candidate receives the same opportunities (83). Therefore, consensus recommendations to support decision-making may be welcomed by the transplant community.

However, this is a challenge due to the multi-factorial factors that influence post-transplant outcomes. Kerr et al., exploring data from the United States, quantified the magnitude of paired deceased donor effects when transplanted into different recipients (84). In analyses adjusted for KDPI, Kerr et al. demonstrated moderate donor effects for DGF and minimal donor effects for 1- and



3-year graft failure, with 4%–8% excess absolute risk over baseline for a graft if the mate kidney failed. Therefore, it is important to appreciate that post kidney transplant outcomes are influenced by a complex interplay of factors that include, but are not exclusive to, donor characteristics.

Developing and validating novel strategies and/or techniques to improve the process is therefore necessary. Various tools to aid decision-making are currently available or under investigation. These include donor risk scores in the setting of DCD kidneys (85), donor-recipient characteristics (86), donor-specific features (87), monitoring of perfusion parameters and assessment of tissue viability function ex situ (88), molecular diagnostics (89), and machine learning and artificial intelligence (AI) algorithms (90-92). The latter remains in its infancy, with tremendous potential to augment the decision-making regarding transplantation (93), but requires more granular data, generalizability, and validation across different population cohorts to enter mainstream use. Such AI tools must provide survival probabilities for kidney transplant candidates to proceed with an individual organ offer versus remaining on the waiting-list to allow a meaningful decision to be made about transplantation. While some risk communication tools are available (http:// www.transplantmodels.com or https://www.odt.nhs.uk/ transplantation/tools-policies-and-guidance/risk-communicationtools/for the US and UK respectively), they lack the machine learning capability or enhanced AI to provide more personalized risk probabilities.

CONCLUSION

Complex deceased donor kidney offers, with time-pressured decision-making, can lead to unnecessary decline and/or discard of acceptable kidneys. By outlining donor clinical factors associated with post kidney transplant outcomes, the aim of this review is to support clinical decision-making. However, donor characteristics are only one component of a complex interplay that influence post-transplant outcomes. While any kidney allograft may not be better than no kidney allograft in every clinical scenario, the objective evidence would argue most kidney allografts are better than being denied the opportunity of kidney transplantation if deemed suitable for waitlisting.

AUTHOR CONTRIBUTIONS

The author confirms being the sole contributor of this work and has approved it for publication.

CONFLICT OF INTEREST

AS is the local Clinical Lead for Utilisation (CLU) supported by NHS Blood and Transplant. AS has participated in the Stakeholder Forum for the Organ Utilisation Group set up by the Department of Health and Social Care. Both organisations have had no role in the writing of this manuscript.

REFERENCES

- Jardine AG, Gaston RS, Fellstrom BC, Holdaas H. Prevention of Cardiovascular Disease in Adult Recipients of Kidney Transplants. *Lancet* (2011) 378(9800):1419–27. doi:10.1016/S0140-6736(11)61334-2
- 2. Chapman JR. The Consequences of Successful Transplantation. Lancet~(2011)~378(9800):1357-9.~doi:10.1016/S0140-6736(10)61111-7
- Chaudhry D, Chaudhry A, Peracha J, Sharif A. Survival for Waitlisted Kidney Failure Patients Receiving Transplantation versus Remaining on Waiting List: Systematic Review and Meta-Analysis. BMJ (2022) 376:e068769. doi:10.1136/ bmj-2021-068769
- Watson CJ, Dark JH. Organ Transplantation: Historical Perspective and Current Practice. Br J Anaesth (2012) 108(1):i29–42. doi:10.1093/bja/aer384
- Gill JS, Abichandani R, Kausz AT, Pereira BJ. Mortality after Kidney Transplant Failure: the Impact of Non-immunologic Factors. Kidney Int (2002) 62(5):1875–83. doi:10.1046/j.1523-1755.2002.00640.x
- Rao PS, Schaubel DE, Jia X, Li S, Port FK, Saran R. Survival on Dialysis postkidney Transplant Failure: Results from the Scientific Registry of Transplant Recipients. Am J Kidney Dis (2007) 49(2):294–300. doi:10.1053/j.ajkd.2006. 11.022
- Foley DP, Sawinski D. Personalizing Donor Kidney Selection: Choosing the Right Donor for the Right Recipient. Clin J Am Soc Nephrol (2020) 15(3): 418–20. doi:10.2215/CJN.09180819
- Mohan S, Chiles MC, Patzer RE, Pastan SO, Husain SA, Carpenter DJ, et al. Factors Leading to the Discard of Deceased Donor Kidneys in the United States. Kidney Int (2018) 94(1):187–98. doi:10.1016/j.kint.2018.02.016
- Stewart DE, Garcia VC, Rosendale JD, Klassen DK, Carrico BJ. Diagnosing the Decades-Long Rise in the Deceased Donor Kidney Discard Rate in the United States. *Transplantation* (2017) 101(3):575–87. doi:10.1097/TP. 0000000000001539
- Council of Europe. Newsletter Transplant: International Figures on Donation and Transplantation (2018). EQDM 2019. ISSN: 2171-4118.
- Maghen A, Mone TD, Veale J. The Kidney-Transplant Waiting List and the Opioid Crisis. N Engl J Med (2019) 380(23):2273–4. doi:10.1056/ NEJMc1817188
- Aubert O, Reese PP, Audry B, Bouatou Y, Raynaud M, Viglietti D, et al. Disparities in Acceptance of Deceased Donor Kidneys between the United States and France and Estimated Effects of Increased US Acceptance. *JAMA Intern Med* (2019) 179(10):1365–74. doi:10.1001/jamainternmed.2019.2322
- Husain SA, King KL, Pastan S, Patzer RE, Cohen DJ, Radhakrishnan J, et al. Association between Declined Offers of Deceased Donor Kidney Allograft and Outcomes in Kidney Transplant Candidates. *JAMA Netw Open* (2019) 2(8): e1910312. doi:10.1001/jamanetworkopen.2019.10312
- Senanayake S, Graves N, Healy H, Baboolal K, Barnett A, Sypek MP, et al. Donor Kidney Quality and Transplant Outcome: An Economic Evaluation of Contemporary Practice. Value Health (2020) 23(12):1561–9. doi:10.1016/j. jval.2020.07.007
- Chapman JR. Marginal Kidneys for Transplantation. BMJ (2015) 351:h3856. doi:10.1136/bmi.h3856
- Pippias M, Stel VS, Arnol M, Bemelman F, Berger SP, Buturovic Ponikvar JB, et al. Temporal Trends in the Quality of Deceased Donor Kidneys and Kidney Transplant Outcomes in Europe: an Analysis by the ERA-EDTA Registry. Nephrol Dial Transpl (2021) 37(1):175–86. doi:10.1093/ndt/gfab156
- Glorioso JM. Kidney Allocation Policy: Past, Present, and Future. Adv Chronic Kidney Dis (2021) 28(6):511–6. doi:10.1053/j.ackd.2022.01.006
- Rao PS, Ojo A. The Alphabet Soup of Kidney Transplantation: SCD, DCD, ECD-Ffundamentals for the Practicing Nephrologist. Clin J Am Soc Nephrol (2009) 4(11):1827–31. doi:10.2215/CJN.02270409
- Pascual J, Zamora J, Pirsch JD. A Systematic Review of Kidney Transplantation from Expanded Criteria Donors. Am J Kidney Dis (2008) 52(3):553–86. doi:10. 1053/j.ajkd.2008.06.005
- Frei U, Noeldeke J, Machold-Fabrizii V, Margreiter R, Fricke L, Voiculescu A, et al. Prospective Age-Matching in Elderly Kidney Transplant Recipients-Aa 5-year Analysis of the Eurotransplant Senior Program. Am J Transpl (2008) 8(1): 50–7. doi:10.1111/j.1600-6143.2007.02014.x

- Querard AH, Foucher Y, Combescure C, Dantan E, Larmet D, Lorent M, et al. Comparison of Survival Outcomes between Expanded Criteria Donor and Standard Criteria Donor Kidney Transplant Recipients: a Systematic Review and Meta-Analysis. *Transpl Int* (2016) 29(4):403–15. doi:10.1111/tri.12736
- Aubert O, Kamar N, Vernerey D, Viglietti D, Martinez F, Duong-Van-Huyen JP, et al. Long Term Outcomes of Transplantation Using Kidneys from Expanded Criteria Donors: Prospective, Population Based Cohort Study. BMJ (2015) 351:h3557. doi:10.1136/bmj.h3557
- Ma MK, Lim WH, Craig JC, Russ GR, Chapman JR, Wong G. Mortality Among Younger and Older Recipients of Kidney Transplants from Expanded Criteria Donors Compared with Standard Criteria Donors. Clin J Am Soc Nephrol (2016) 11(1):128–36. doi:10.2215/CJN.03760415
- Hellemans R, Kramer A, De Meester J, Collart F, Kuypers D, Jadoul M, et al. Does Kidney Transplantation with a Standard or Expanded Criteria Donor Improve Patient Survival? Results from a Belgian Cohort. Nephrol Dial Transpl (2021) 36(5):918–26. doi:10.1093/ndt/gfab024
- Summers DM, Johnson RJ, Allen J, Fuggle SV, Collett D, Watson CJ, et al. Analysis of Factors that Affect Outcome after Transplantation of Kidneys Donated after Cardiac Death in the UK: a Cohort Study. *Lancet* (2010) 376(9749):1303–11. doi:10.1016/S0140-6736(10)60827-6
- Summers DM, Johnson RJ, Hudson A, Collett D, Watson CJ, Bradley JA. Effect
 of Donor Age and Cold Storage Time on Outcome in Recipients of Kidneys
 Donated after Circulatory Death in the UK: a Cohort Study. *Lancet* (2013)
 381(9868):727–34. doi:10.1016/S0140-6736(12)61685-7
- Gill J, Rose C, Lesage J, Joffres Y, Gill J, O'Connor K. Use and Outcomes of Kidneys from Donation after Circulatory Death Donors in the United States. J Am Soc Nephrol (2017) 28(12):3647–57. doi:10.1681/ASN.2017030238
- Summers DM, Watson CJ, Pettigrew GJ, Johnson RJ, Collett D, Neuberger JM, et al. Kidney Donation after Circulatory Death (DCD): State of the Art. Kidney Int (2015) 88(2):241–9. doi:10.1038/ki.2015.88
- Locke JE, Segev DL, Warren DS, Dominici F, Simpkins CE, Montgomery RA.
 Outcomes of Kidneys from Donors after Cardiac Death: Implications for
 Allocation and Preservation. Am J Transpl (2007) 7(7):1797–807. doi:10.1111/
 j.1600-6143.2007.01852.x
- Singh SK, Kim SJ. Does Expanded Criteria Donor Status Modify the Outcomes of Kidney Transplantation from Donors after Cardiac Death? Am J Transpl (2013) 13(2):329–36. doi:10.1111/j.1600-6143.2012.04311.x
- 31. Nagaraja P, Roberts GW, Stephens M, Horvath S, Kaposztas Z, Chavez R, et al. Impact of Expanded Criteria Variables on Outcomes of Kidney Transplantation from Donors after Cardiac Death. *Transplantation* (2015) 99(1):226–31. doi:10.1097/TP.0000000000000304
- 32. Doshi MD, Hunsicker LG. Short- and Long-Term Outcomes with the Use of Kidneys and Livers Donated after Cardiac Death. *Am J Transpl* (2007) 7(1): 122–9. doi:10.1111/j.1600-6143.2006.01587.x
- Montero N, Toapanta N, Pallares N, Crespo M, Diekmann F, Guirado L, et al. Deciphering Transplant Outcomes of Expanded Kidney Allografts Donated after Controlled Circulatory Death in the Current Transplant Era. A Call for Caution. *Transpl Int* (2021) 34(12):2494–506. doi:10.1111/tri.14131
- Rao PS, Schaubel DE, Guidinger MK, Andreoni KA, Wolfe RA, Merion RM, et al. A Comprehensive Risk Quantification Score for Deceased Donor Kidneys: the Kidney Donor Risk index. *Transplantation* (2009) 88(2): 231–6. doi:10.1097/TP.0b013e3181ac620b
- Zhong Y, Schaubel DE, Kalbfleisch JD, Ashby VB, Rao PS, Sung RS. Reevaluation of the Kidney Donor Risk Index. *Transplantation* (2019) 103(8):1714–21. doi:10.1097/TP.000000000002498
- Israni AK, Salkowski N, Gustafson S, Snyder JJ, Friedewald JJ, Formica RN, et al. New National Allocation Policy for Deceased Donor Kidneys in the United States and Possible Effect on Patient Outcomes. J Am Soc Nephrol (2014) 25(8):1842–8. doi:10.1681/ASN.2013070784
- 37. Woodside KJ, Merion RM, Leichtman AB, de los Santos R, Arrington CJ, Rao PS, et al. Utilization of Kidneys with Similar Kidney Donor Risk index Values from Standard versus Expanded Criteria Donors. *Am J Transpl* (2012) 12(8): 2106–14. doi:10.1111/j.1600-6143.2012.04146.x
- Dahmen M, Becker F, Pavenstadt H, Suwelack B, Schutte-Nutgen K, Reuter S. Validation of the Kidney Donor Profile Index (KDPI) to Assess a Deceased Donor's Kidneys' Outcome in a European Cohort. Sci Rep (2019) 9(1):11234. doi:10.1038/s41598-019-47772-7

- Sexton DJ, O'Kelly P, Kennedy C, Denton M, de Freitas DG, Magee C, et al. Assessing the Discrimination of the Kidney Donor Risk Index/Kidney Donor Profile Index Scores for Allograft Failure and Estimated Glomerular Filtration Rate in Ireland's National Kidney Transplant Programme. Clin Kidney J (2019) 12(4):569–73. doi:10.1093/ckj/sfy130
- Foley RN, Hakim RM. Why Is the Mortality of Dialysis Patients in the United States Much Higher Than the Rest of the World? J Am Soc Nephrol (2009) 20(7):1432–5. doi:10.1681/ASN.2009030282
- Foster BJ, Mitsnefes MM, Dahhou M, Zhang X, Laskin BL. Changes in Excess Mortality from End Stage Renal Disease in the United States from 1995 to 2013. Clin J Am Soc Nephrol (2018) 13(1):91–9. doi:10.2215/CJN.04330417
- Merion RM, Goodrich NP, Johnson RJ, McDonald SP, Russ GR, Gillespie BW, et al. Kidney Transplant Graft Outcomes in 379 257 Recipients on 3 Continents. Am J Transpl (2018) 18(8):1914–23. doi:10.1111/ajt.14694
- 43. Ibrahim M, Vece G, Mehew J, Johnson R, Forsythe J, Klassen D, et al. An International Comparison of Deceased Donor Kidney Utilization: What Can the United States and the United Kingdom Learn from Each Other? Am J Transpl (2020) 20(5):1309–22. doi:10.1111/ajt.15719
- Dayoub JC, Cortese F, Anzic A, Grum T, de Magalhaes JP. The Effects of Donor Age on Organ Transplants: A Review and Implications for Aging Research. Exp Gerontol (2018) 110:230–40. doi:10.1016/j.exger.2018.06.019
- Keith DS, Demattos A, Golconda M, Prather J, Norman D. Effect of Donor Recipient Age Match on Survival after First Deceased Donor Renal Transplantation. J Am Soc Nephrol (2004) 15(4):1086–91. doi:10.1097/01. asn.0000119572.02053.f2
- Meier-Kriesche HU, Cibrik DM, Ojo AO, Hanson JA, Magee JC, Rudich SM, et al. Interaction between Donor and Recipient Age in Determining the Risk of Chronic Renal Allograft Failure. *J Am Geriatr Soc* (2002) 50(1):14–7. doi:10. 1046/j.1532-5415.2002.50002.x
- 47. Ibrahim M, Greenhall GHB, Summers DM, Mumford L, Johnson R, Baker RJ, et al. Utilization and Outcomes of Single and Dual Kidney Transplants from Older Deceased Donors in the United Kingdom. *Clin J Am Soc Nephrol* (2020) 15(9):1320–9. doi:10.2215/CJN.02060220
- Pisavadia B, Arshad A, Chappelow I, Nightingale P, Anderson B, Nath J, et al. Ethnicity Matching and Outcomes after Kidney Transplantation in the United Kingdom. PLoS One (2018) 13(4):e0195038. doi:10.1371/journal. pone.0195038
- Locke JE, Warren DS, Dominici F, Cameron AM, Leffell MS, McRann DA, et al. Donor Ethnicity Influences Outcomes Following Deceased-Donor Kidney Transplantation in Black Recipients. J Am Soc Nephrol (2008) 19(10):2011–9. doi:10.1681/ASN.2008010078
- Swanson SJ, Hypolite IO, Agodoa LY, Batty DS, Hshieh PB, Cruess D, et al. Effect of Donor Factors on Early Graft Survival in Adult Cadaveric Renal Transplantation. Am J Transpl (2002) 2(1):68–75. doi:10.1034/j.1600-6143. 2002.020112.x
- Chakkera HA, O'Hare AM, Johansen KL, Hynes D, Stroupe K, Colin PM, et al. Influence of Race on Kidney Transplant Outcomes within and outside the Department of Veterans Affairs. J Am Soc Nephrol (2005) 16(1):269–77. doi:10. 1681/ASN.2004040333
- Vyas DA, Eisenstein LG, Jones DS. Hidden in Plain Sight Reconsidering the Use of Race Correction in Clinical Algorithms. N Engl J Med (2020) 383(9): 874–82. doi:10.1056/NEJMms2004740
- Chong K, Litvinovich I, Chen S, Zhu Y, Argyropoulos C. Reconsidering Donor Race in Predicting Allograft and Patient Survival Among Kidney Transplant Recipients. Kidney360 (2021) 2:1831–5. doi:10.34067/KID.0002932021
- Arshad A, Hodson J, Chappelow I, Inston NG, Ready AR, Nath J, et al. The Impact of Donor Body Mass index on Outcomes after Deceased Kidney Transplantation - a National Population-Cohort Study. *Transpl Int* (2018) 31(10):1099–109. doi:10.1111/tri.13263
- Ortiz J, Gregg A, Wen X, Karipineni F, Kayler LK. Impact of Donor Obesity and Donation after Cardiac Death on Outcomes after Kidney Transplantation. Clin Transpl (2012) 26(3):E284–92. doi:10.1111/j.1399-0012.2012.01649.x
- Arshad A, Hodson J, Chappelow I, Nath J, Sharif A. The Influence of Donor to Recipient Size Matching on Kidney Transplant Outcomes. *Transpl Direct* (2018) 4(10):e391. doi:10.1097/TXD.0000000000000826
- Kasiske BL, Snyder JJ, Gilbertson D. Inadequate Donor Size in Cadaver Kidney Transplantation. J Am Soc Nephrol (2002) 13(8):2152–9. doi:10.1097/01.asn. 0000024564.22119.3d

- Goldberg RJ, Smits G, Wiseman AC. Long-term Impact of Donor-Recipient Size Mismatching in Deceased Donor Kidney Transplantation and in Expanded Criteria Donor Recipients. *Transplantation* (2010) 90(8):867–74. doi:10.1097/TP.0b013e3181f24e75
- Miller AJ, Kiberd BA, Alwayn IP, Odutayo A, Tennankore KK. Donor-Recipient Weight and Sex Mismatch and the Risk of Graft Loss in Renal Transplantation. Clin J Am Soc Nephrol (2017) 12(4):669–76. doi:10.2215/CJN. 07660716
- Koyawala N, Parikh CR. A Review of Donor Acute Kidney Injury and Posttransplant Outcomes. Transplantation (2020) 104(8):1553–9. doi:10. 1097/TP.000000000003144
- Zheng YT, Chen CB, Yuan XP, Wang CX. Impact of Acute Kidney Injury in Donors on Renal Graft Survival: a Systematic Review and Meta-Analysis. *Ren Fail* (2018) 40(1):649–56. doi:10.1080/0886022X.2018.1535982
- Boffa C, van de Leemkolk F, Curnow E, Homan van der Heide J, Gilbert J, SharplEs E, et al. Transplantation of Kidneys from Donors with Acute Kidney Injury: Friend or Foe? Am J Transpl (2017) 17(2):411-9. doi:10. 1111/ajt.13966
- 63. Ko KJ, Kim YH, Kim MH, Jun KW, Kwon KH, Kim HS, et al. Kidney Transplantation Using Expanded Criteria Deceased Donors with Terminal Acute Kidney Injury: a Single center Experience in Korea. Ann Surg Treat Res (2018) 95(5):278–85. doi:10.4174/astr.2018.95.5.278
- 64. Park WY, Choi MS, Kim YS, Choi BS, Park CW, Yang CW, et al. Impact of Acute Kidney Injury in Expanded Criteria Deceased Donors on posttransplant Clinical Outcomes: Multicenter Cohort Study. BMC Nephrol (2019) 20(1):39. doi:10.1186/s12882-019-1225-1
- Jadlowiec CC, Hanna WA, Ninan J, Ryan MS, Das DM, Smith M, et al. Transplant Outcomes Using Kidneys from High KDPI Acute Kidney Injury Donors. Clin Transpl (2021) 35(5):e14279. doi:10.1111/ctr.14279
- Cohen JB, Bloom RD, Reese PP, Porrett PM, Forde KA, Sawinski DL. National Outcomes of Kidney Transplantation from Deceased Diabetic Donors. Kidney Int (2015) 89:636–47. doi:10.1038/ki.2015.325
- Cohen JB, Eddinger KC, Locke JE, Forde KA, Reese PP, Sawinski DL. Survival Benefit of Transplantation with a Deceased Diabetic Donor Kidney Compared with Remaining on the Waitlist. Clin J Am Soc Nephrol (2017) 12(6):974–82. doi:10.2215/CJN.10280916
- Altheaby A, Al Dalbhi S, Alghamdi Y, Almigbal TH, Alotaibi KN, Batais MA, et al. Effect of Donor Hypertension on Renal Transplant Recipients' Blood Pressure, Allograft Outcomes and Survival: a Systematic Review and Meta-Analysis. Am J Cardiovasc Dis (2019) 9(4):49–58.
- Lin SJ, Koford JK, Baird BC, Hurdle JF, Krikov S, Habib AN, et al. Effect of Donors' Intravenous Drug Use, Cigarette Smoking, and Alcohol Dependence on Kidney Transplant Outcome. *Transplantation* (2005) 80(4):482–6. doi:10. 1097/01.tp.0000168154.14458.28
- Gillott H, Jackson Spence F, Tahir S, Hodson J, Nath J, Sharif A. Deceased-Donor Smoking History Is Associated with Increased Recipient Mortality after Kidney Transplant: A Population-Cohort Study. Exp Clin Transpl (2019) 17(2):183–9. doi:10.6002/ect.2017.0198
- Ison MG, Hager J, Blumberg E, Burdick J, Carney K, Cutler J, et al. Donorderived Disease Transmission Events in the United States: Data Reviewed by the OPTN/UNOS Disease Transmission Advisory Committee. *Am J Transpl* (2009) 9(8):1929–35. doi:10.1111/j.1600-6143.2009.02700.x
- Bowring MG, Holscher CM, Zhou S, Massie AB, Garonzik-Wang J, Kucirka LM, et al. Turn Down for what? Patient Outcomes Associated with Declining Increased Infectious Risk Kidneys. Am J Transpl (2018) 18(3):617–24. doi:10. 1111/ajt.14577
- Eccher A, Girolami I, Motter JD, Marletta S, Gambaro G, Momo REN, et al. Donor-transmitted Cancer in Kidney Transplant Recipients: a Systematic Review. J Nephrol (2020) 33(6):1321–32. doi:10.1007/s40620-020-00775-4
- Desai R, Collett D, Watson CJ, Johnson P, Evans T, Neuberger J. Cancer Transmission from Organ Donors-Unavoidable but Low Risk. Transplantation (2012) 94(12):1200-7. doi:10.1097/TP.0b013e318272df41
- Trotter PB, Summers DM, Robb M, Hulme W, Ushiro-Lumb I, Watson CJE, et al. Deceased Organ Donors with a History of Increased Risk Behavior for the Transmission of Blood-Borne Viral Infection: The UK Experience. Transplantation (2017) 101(7):1679–89. doi:10.1097/TP.00000000000001727
- 76. Durand CM, Bowring MG, Thomas AG, Kucirka LM, Massie AB, Cameron A, et al. The Drug Overdose Epidemic and Deceased-Donor Transplantation in

- the United States: A National Registry Study. Ann Intern Med (2018) 168(10): 702-11. doi:10.7326/M17-2451
- Limkemann AJ, Wolfe L, Sharma A, King AL, Behnke M, Shah MJ, et al. Outcomes of Kidney Transplants and Risk of Infection Transmission from Increased Infectious Risk Donors. Clin Transpl (2016) 30(8):886–93. doi:10. 1111/ctr.12761
- Kucirka LM, Sarathy H, Govindan P, Wolf JH, Ellison TA, Hart LJ, et al. Risk of Window Period HIV Infection in High Infectious Risk Donors: Systematic Review and Meta-Analysis. Am J Transpl (2011) 11(6):1176–87. doi:10.1111/j. 1600-6143.2010.03329.x
- Kucirka LM, Sarathy H, Govindan P, Wolf JH, Ellison TA, Hart LJ, et al. Risk of Window Period Hepatitis-C Infection in High Infectious Risk Donors: Systematic Review and Meta-Analysis. Am J Transpl (2011) 11(6): 1188–200. doi:10.1111/j.1600-6143.2011.03460.x
- Wang CJ, Wetmore JB, Crary GS, Kasiske BL. The Donor Kidney Biopsy and its Implications in Predicting Graft Outcomes: A Systematic Review. Am J Transpl (2015) 15(7):1903–14. doi:10.1111/ajt.13213
- Girolami I, Gambaro G, Ghimenton C, Beccari S, Calio A, Brunelli M, et al. Pre-implantation Kidney Biopsy: Value of the Expertise in Determining Histological Score and Comparison with the Whole Organ on a Series of Discarded Kidneys. J Nephrol (2020) 33(1):167–76. doi:10.1007/s40620-019-00638-7
- 82. Ayorinde JO, Summers DM, Pankhurst L, Laing E, Deary AJ, Hemming K, et al. PreImplantation Trial of Histopathology in Renal Allografts (PITHIA): a Stepped-Wedge Cluster Randomised Controlled Trial Protocol. *BMJ Open* (2019) 9(1):e026166. doi:10.1136/bmjopen-2018-026166
- Sharif A. Risk Aversion, Organ Utilization and Changing Behavior. Transpl Int (2022) 35:10339. doi:10.3389/ti.2022.10339
- Kerr KF, Morenz ER, Thiessen-Philbrook H, Coca SG, Wilson FP, Reese PP, et al. Quantifying Donor Effects on Transplant Outcomes Using Kidney Pairs from Deceased Donors. Clin J Am Soc Nephrol (2019) 14(12):1781–7. doi:10. 2215/CJN.03810319
- Plata-Munoz JJ, Vazquez-Montes M, Friend PJ, Fuggle SV. The Deceased Donor Score System in Kidney Transplants from Deceased Donors after Cardiac Death. *Transpl Int* (2010) 23(2):131–9. doi:10.1111/j.1432-2277. 2009.00951.x

- Senanayake S, Kularatna S, Healy H, Graves N, Baboolal K, Sypek MP, et al. Development and Validation of a Risk index to Predict Kidney Graft Survival: the Kidney Transplant Risk index. BMC Med Res Methodol (2021) 21(1):127. doi:10.1186/s12874-021-01319-5
- 87. Huang E, Danovitch G, Bunnapradist S. Transplantation: Donor Risk Scores: Can They Predict Renal Transplant Outcomes? *Nat Rev Nephrol* (2011) 7(5): 247–8. doi:10.1038/nrneph.2011.39
- Resch T, Cardini B, Oberhuber R, Weissenbacher A, Dumfarth J, Krapf C, et al. Transplanting Marginal Organs in the Era of Modern Machine Perfusion and Advanced Organ Monitoring. Front Immunol (2020) 11:631. doi:10.3389/ fimmu.2020.00631
- von Moos S, Akalin E, Mas V, Mueller TF. Assessment of Organ Quality in Kidney Transplantation by Molecular Analysis and Why it May Not Have Been Achieved, yet. Front Immunol (2020) 11:833. doi:10.3389/fimmu.2020. 00833
- Yoo KD, Noh J, Lee H, Kim DK, Lim CS, Kim YH, et al. A Machine Learning Approach Using Survival Statistics to Predict Graft Survival in Kidney Transplant Recipients: A Multicenter Cohort Study. Sci Rep (2017) 7(1): 8904. doi:10.1038/s41598-017-08008-8
- Threlkeld R, Ashiku L, Canfield C, Shank DB, Schnitzler MA, Lentine KL, et al. Reducing Kidney Discard with Artificial Intelligence Decision Support: the Need for a Transdisciplinary Systems Approach. Curr Transpl Rep (2021) 8(4): 263–71. doi:10.1007/s40472-021-00351-0
- Bae S, Massie AB, Caffo BS, Jackson KR, Segev DL. Machine Learning to Predict Transplant Outcomes: Helpful or Hype? A National Cohort Study. Transpl Int (2020) 33(11):1472–80. doi:10.1111/tri.13695
- Clement J, Maldonado AQ. Augmenting the Transplant Team with Artificial Intelligence: Toward Meaningful AI Use in Solid Organ Transplant. Front Immunol (2021) 12:694222. doi:10.3389/fimmu.2021.694222

Copyright © 2022 Sharif. This is an open-access article distributed under the terms of the Creative Commons Attribution License (CC BY). The use, distribution or reproduction in other forums is permitted, provided the original author(s) and the copyright owner(s) are credited and that the original publication in this journal is cited, in accordance with accepted academic practice. No use, distribution or reproduction is permitted which does not comply with these terms.