

TRENDS IN UTILIZATION OF DECEASED DONOR KIDNEYS BASED ON HEPATITIS C VIRUS STATUS AND IMPACT OF PUBLIC HEALTH SERVICE LABELLING ON DISCARD

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Abbreviations:

Antibody: Ab

Deceased Donor Renal Transplant: DDRT

Hepatitis C Virus: HCV

Kidney Donor Profile Index: KDPI

Nucleic Acid Test: NAT

Public Health Service Increased Risk: PHS-IR

United Network for Organ Sharing: UNOS

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ABSTRACT

Background: Kidneys from deceased donors infected with hepatitis C virus (HCV) are underutilized. Most HCV virus-infected donors are designated as Public Health Service increased donors (PHS-IR). Impact of PHS and HCV designations on discard is not well studied.

Methods: We queried the UNOS dataset for all deceased donor kidneys between January 2015 and December 2018. The final study cohort donors (n= 38,702) were stratified into three groups based on HCV antibody (Ab) and NAT status: 1) Ab-/NAT- (n= 35,861); 2) Ab+/NAT- (n= 973), and 3) Ab±/NAT+ (n=1,868). We analyzed utilization/discard rates of these organs, the impact of PHS-IR and HCV designations on discard using multivariable two-level hierarchical logistic regression models, forecasted number of HCV viremic donors/kidneys by 2023.

Results: During the study period, 1) the number of viremic donor kidneys increased 2 folds; 2) the multilevel mixed-effects logistic regression models showed that, overall, the PHS labeling (OR 1.20, CI 95% CI 1.15-1.29) and HCV designation (OR 2.29; 95% CI 2.15-2.43) were independently associated with increased risk of discard; 3) contrary to the general perception, PHS-non-IR kidneys across all HCV groups, compared to PHS-IR kidneys were more likely to be discarded; 4) we forecasted that the number of kidneys from HCV viremic donor kidneys might increase from 1,376 in 2019 to 2,092 in 2023.

Conclusion: HCV viremic kidneys might represent 10-15% of deceased donor organ pool soon with the current rate of the opioid epidemic. PHS labeling effect on discard requires further discussion of the utility of this classification.

INTRODUCTION:

Renal transplantation (RT) is the treatment of choice for end-stage renal disease (ESRD).^{1,2} Despite recent increases in the number of deceased donor (DD) RT³, there still exists a wide gap between supply and demand for RT. While there has been a concerted effort to maximize the utilization of kidneys from existing donors⁴ and to increase the donor pool as well⁵, the proportion of kidneys discarded remains high. The last decade in the United States (US) has witnessed a significant change in the demographics of opioid users.⁶ Opioid use is increasing amongst Caucasians with even higher rates in the Midwestern US. Heroin use went up fivefold from 2002 to 2013⁷, coinciding with a surge in intravenous drug use (IVDU), hepatitis C virus (HCV) transmission, and opioid-related overdose deaths.⁸⁻¹¹ Donors dying due to overdose are more likely to be infected with HCV ¹¹, and organs from HCV positive donors are underutilized.¹²⁻¹⁵ Single-center studies have utilized HCV antibody positive^{16,17} and viremic donors¹⁸⁻²⁰ for RT with good short-term outcomes. A recent national registry analysis by our group confirms excellent short term outcomes for such transplants.²¹

Kidneys recovered from opioid overdose death donors have predominantly been classified as the Public Health Service increased risk (PHS-IR) donors, implying higher transmission risk of viral infection (mainly HCV, hepatitis B [HBV], and human immunodeficiency virus [HIV]) through organ donation (previously defined as Center for

Disease Control and Prevention high risk donors). $^{22-24}$ New guidelines obligated use of nucleic acid testing (NAT) supplementing serologic ones (mainly for HCV, HBV, and HIV) in 2013 for all PHS-IR donors, and were officially implemented in 2015 in the U.S. 24

The purpose of our study was to analyze trends in center specific, organ procurement organization (OPO) level, regional utilization of adult kidney donors based on donor HCV Ab and NAT status, study the impact of PHS labeling and HCV designation on discard of those kidneys, and forecast the number of HCV viremic donors by 2023.

PATIENTS AND METHODS/MATERIALS AND METHODS:

Study Population:

This study used data from the OPTN STAR files administered by the United Network of Organ Sharing (UNOS), which includes data submitted by members on all donors, waitlisted candidates, and transplant recipients in the US. The Health Resources and Services Administration (HRSA) of the US Department of Health and Human Services (DHHS) oversees the activities of the OPTN and the contractor. The University of Texas Southwestern Institutional Review Board approved the study.

This retrospective cohort study included all deceased donors registered in the OPTN STAR files from January 1st, 2015 through December 31st, 2018. Donors with incomplete HCV Ab and NAT information were excluded. Thus, we identified 38,702 deceased donors as a final cohort during the study period. For our analyses, HCV uninfected donor is defined as a donor with negative HCV Ab and negative NAT (HCV Ab- /NAT-); an HCV seropositive, non-viremic donor is defined as a donor with positive HCV Ab and negative NAT (HCV Ab+ /NAT-); and an HCV viremic donor is identified as a subject with positive HCV NAT, regardless of the HCV Ab status (HCV Ab ± /NAT+). The term "HCV positive donor" refers to donors with a positive HCV Ab and/or positive HCV NAT.

The study cohort donors (n= 38,702) were stratified into three groups based on HCV Ab and NAT status: 1) Ab-/NAT- (n= 35,861); 2) Ab+/NAT- (n= 973), and 3) Ab±

/NAT+ (n=1,868). Under each HCV categories, the kidneys (N=70,450) from above donors were further classified as "PHS-IR" or "PHS-non-IR" for the logistic regression analysis to predict discard: 1) HCV Ab- /NAT- (n=66,224) category was composed of PHS-IR (n=13,411, 20.3%) and PHS-non-IR (n=52,787, 79.7%); 2) HCV Ab+ /NAT- (n=1,459) category was composed of PHS-IR (n=1,030, 70.7%) and PHS-non-IR (n=427, 29.3%); 3) HCV Ab± /NAT+ (n=2,767) category was composed of PHS-IR (n=2,298, 83.0%) and PHS-non-IR (n=469, 17.0%).

Primary Outcomes:

Primary outcome measures were transplantation and discard rates of deceased donors, utilization of NAT+ donors by transplant centers, Organ Procurement Organizations (OPO), UNOS Region, the impact of PHS-IR and HCV designation on discard, and forecasted number of HCV viremic kidneys by 2023.

Statistical Methods:

Donor characteristics were summarized by mean and standard deviation for continuous variables, and count and percent of the total for categorical variables. Comparisons between groups were made using t-test or Wilcoxon rank-sum test (non-parametric), one-way ANOVA or Kruskal-Wallis test by ranks (non-parametric) for continuous variables, and Chi-squared test for categorical variables as appropriate. The Holm multiple comparison adjustments was used as a follow up to one-way-ANOVA to calculate multiplicity adjusted p-values. The magnitude of missing data was minimal (<2%); thus, imputation was not used. A p-value < 0.05 was considered statistically significant. Statistical analyses were performed with Stata/MP14 (StataCorp LP, College Station, TX) and R Free Software Foundation (version 3.5.1 version).

To account for variations in discard rates among the UNOS Regions (there are total of 11 regions in the U.S.) and OPOs (there are total of 58 OPOs under eleven UNOS Regions), we utilized multilevel (two-level and three-level models) mixed-effect logistic regression models. For this analysis, we used the Stata command of "melogit" which fits mixed-effects models for binary responses

(https://www.stata.com/manuals14/melogit.pdf). Mixed-effects logistic regression

contains both fixed and random effects. It is useful for modeling intracluster correlation because donors in the same cluster (the UNOS Region or OPO) are correlated and share common cluster-level random effects. We run three separate mixed-effects logistic regression analysis defining random effects for 1) the UNOS Regions (two-level models); 2) OPOs (two-level models); and 3) OPOs nested within the UNOS Regions (three-level models). For simplicity, we only reported results of the mixed-effects logistic regression models for the UNOS Regions (two-level models) because the results of other two models (for OPOs and OPOs nested within the UNOS Regions) did not show any major differences.

The mixed-effects logistic regression models were adjusted for previously identified donor factors in the literature^{25,26}, including donor age >50 or not, either kidney biopsied, glomerulosclerosis >20% or not if biopsied, cytomegalovirus (CMV) status, KDPI, cause death due to cerebrovascular accident (CVA), donation after cardiac death (DCD) status, height, weight, history of tattoo, either kidney pumped, cold ischemica time, hepatitis B core antibody status, hepatitis B surface antigen status, history of diabetes, history of hypertension, history of cocaine use, history of IV drug use (IVDU), terminal creatinine >1.5 mg/dl or not, ABO blood type, transplant year, and race. We did not find multicollinearity between individual elements of KDPI (10 donor variables) and KDPI score, therefore we decided to keep KDPI in the multivariable mixed-effects logistic regression models.

The potential number of deceased donors with HCV NAT positivity is forecasted into the year 2023 using time series analysis with trend adjusted exponential smoothing method; Excel's built-in FORECAST.ETS function was utilized for this purpose.

RESULTS:

The Study Cohort Selection:

During the study period, 38,702 potential kidney donors became available (Figure 1). Consent was obtained only for 76,888 kidneys, of which 5,804 kidneys were not procured. Notably, HCV as the reported reason for a kidney not being procured was only 1%. Among the 71,084 kidneys that were procured, 56,833 (73.9%) kidneys were

transplanted, 13,617 (17.7%) kidneys were discarded, and 634 (0.8%) were used for research.

Characteristics of All Deceased Donors by HCV status and Disposition:

Characteristics of the deceased donors by HCV status are shown in Table 1. Ab-/NAT- donors comprised the majority of the study cohort (n=35,861) and served as the reference group. There were 1,868 donors in the Ab±/NAT+ (viremic) group and 973 donors in the Ab+/NAT- group. The highest number of HCV viremic donor kidneys was recovered in the UNOS region 2 (U.S. States DE, DC, MD, NJ, PA, and WV; a total of 373 donors).

Among all deceased donors (Table 1) and recovered kidneys for transplantation (mainly transplanted and discarded ones, shown in Table 2-3), compared to the reference group (HCV Ab- /NAT-), HCV viremic donors were younger more likely to be White, and male, less likely to be diagnosed with diabetes and hypertension, and had less donation after circulatory death (DCD) donors. As expected, HCV viremic donors also had higher KDPI and were also more likely to be labeled as PHS - IR donors.

Disposition of Deceased Donor Kidneys by HCV Status:

Trends in deceased donor kidney disposition by HCV status over time is shown in Figure 2. Number of Ab+ /NAT- kidneys that were transplanted increased from 103 (35.9% of such kidneys) in 2015 to 444 (66%) in 2018. The discard rate in the same group decreased from 32.4% to 22.4%. The percentage of viremic donor kidneys transplanted (from 41% to 50%) and discarded (from 32% to 33%) slightly increased. Disposition categories for the reference group remained stable during the study period.

Comparison of KDPI Categories in Transplanted and Discarded Deceased Donors by HCV status:

The KDPI distributions of transplanted and discarded kidneys for the reference group were widely separated (left-skewed in the discarded group) and stayed stable for four years period (see Figure 3). The similar distribution pattern was observed in HCV Ab+ /NAT- group in 2018. On the other hand, the KDPI distributions of transplanted and

discarded kidneys for the viremic group mostly overlapped, and the median KDPI percentage was persistently higher in the discarded group during the study period.

Reasons for kidney discard by HCV status:

Table 4 shows the reasons for kidney discard by HCV status. 'No recipient located/list exhausted' and biopsy findings uniformly appear to be two most common reasons for discard across all groups.

Kidney discards by the HCV groups and PHS designations

Table 5 shows relevant characteristics discards by the HCV categories and PHS designations in recovered kidneys (excluding the ones used for research) for transplantation and demonstrates the effect of HCV and PHS designation on discard using multivariable mixed-effect logistic regression models. In all cohort (N=70,450), 23.8% of the kidneys were designated as PHS-IR, had a mean (SD) KDPI of 51.0 ± 29.0%, and experienced a discard rate of 19.3%. PHS IR (odds ratio [OR] 1.20, 95% confidence interval [CI] 1.15-1,29) and HCV designations (OR 2.29, 95% CI 2.15-2.43) were independently associated with increased risk of discard.

In HCV Ab-/NAT- group (n=66,224), 20.3% of the kidneys were designated as PHS IR, and the donor age, mean KDPIs and discard rates were lower in PHS-IR group compared to PHS-non-IR group. The PHS designation was associated with a 17% increased risk of discard (OR 1.24, 95% CI 1.15-1.34) in PHS-IR group compared to the reference group (PHS non-IR) in this category.

In HCV Ab+ /NAT- group (n=1,459), 70.7% of the kidneys were designated as PHS-IR, and the donor age, mean KDPIs and discard rates were lower in PHS-IR group compared to PHS non-IR group. While the PHS designation was not associated with increased risk of discard (OR 1.24, 95% CI 0.84-1.83) in PHS-IR group compared to the reference group (PHS-non-IR in this category), HCV Ab+ /NAT- status increased the odds of discard by approximately 2 folds (OR 2.07, 95% CI 1.78-2.40) compared to the reference group (HCV Ab- /NAT- group).

In HCV Ab± /NAT+ group (n=2,767), 83.0% of the kidneys were designated as PHS IR, and similarly, the donor age, mean KDPIs and discard rates were lower in PHS-IR group compared to PHS-non-IR group. While the PHS designation was not associated with increased risk of discard (OR 1.04, 95% CI 0.79-1.38) in PHS-IR group compared to the reference group (PHS-non-IR in this category), HCV Ab± /NAT+ status increased the odds of discard by approximately 5 folds (OR 5.21, 95% CI 4.62-5.89) compared to the reference group (HCV Ab- /NAT- group).

In the PHS-non-IR kidneys across all HCV groups, compared to the PHS-IR, more recovery biopsies were performed (slightly higher than 50%) that showed a higher percentage of glomerulosclerosis>20% and moderate-to-severe interstitial fibrosis.

Kidney Transplant Center/OPO/UNOS Region Utilization of HCV Viremic Kidneys:

Figures 4 shows the heat map geographic data (the number of kidneys) from viremic donors recovered (Fig 4a) and transplanted (Fig 4b) based on the UNOS Regions. The UNOS Regions 2 and 3 were more likely to procure, and transplant kidneys from viremic donors. Figure 5 and 6 shows the geographic distribution of transplantation with viremic kidneys according to the OPOs and individual transplant centers, respectively. The number of OPOs that transplanted at least 25 kidneys from viremic donors increased from only one in 2015 to six in 2018. There were at least two centers that transplanted more than 60 viremic donor kidneys in 2018.

Forecasting Number of Potential viremic Kidneys by 2023:

We forecasted a potential number of HCV NAT+ DD kidneys that may become available in 2023, based on actual numbers of such kidneys from 2015-2018, using time series trend adjusted exponential smoothing method. We predict about 2,092 HCV positive kidneys from deceased donors would be available by 2023 (Figure 7), the model assumes that the opioid epidemic and related overdose deaths continue to rise exponentially with the same trend.

DISCUSSION:

This study reveals some key insights about the recent trends in kidney transplant utilization in the U.S.: 1) an increasing number and utilization rates of Ab+ /NATkidneys (annual transplant rate increased from 35.9% in 2015 to 66% in 2018) showing a positive change in transplant centers' behavior and patient acceptance of minimal infectious transmission risk organs; 2) the number of viremic donor kidneys increased from 658 in 2015 to 1,144 in 2018, and the number of OPOs transplanting at least 25 viremic donor kidneys increased from one in 2015 to six in 2018; 3) no recipient located/list exhausted' was the most common reason for discard across all groups (40.4%), and even higher in the viremic donor group (65.4%); 4) PHS designation (OR 1.20, CI 95% CI 1.15-1.29) and HCV status (2.29; 95% CI 2.15-2.43) were independently associated with increased risk of discard; 5) PHS-non-IR kidneys across all HCV groups, compared to PHS IR kidneys, were more likely to be discarded (contrary to common perception), had higher KDPI scores, and underwent more biopsies showing slightly higher percentage of glomerulosclerosis (GS) >20% and moderate-to-severe interstitial fibrosis (IF); 6) the reasons for high kidney discards are multifactorial, could partially be explained by KDPI score, the performance of procurement biopsy and its findings for HCV infected kidneys; 7) We forecasted that the number of kidneys from HCV viremic donors would increase from 1,376 in 2019 to 2,092 in 2023 which might represent 10-15% of deceased donor organ supply over the next few years with current rate of opioid epidemic.

Decision to discard a deceased donor kidney is influenced by several factors including variability in regional/OPO/center wait time and wait-list size, center transplant rates/aggressiveness, KDPI score, CIT, decision to biopsy and biopsy findings, pump parameters, regional and national share, living donation access, PHS designation, HCV status, perceived risk/benefit ratio, recipients socioeconomic status. Accepting a PHS-IR organ offers survival benefit to recipients compared to those who declined it and are waiting for a PHS-non-IR donor offer and staying on dialysis. 5,27-30 There exist a notion that a disproportionate number of discarded kidneys originate from PHS-IR donors. 31,32 In our study cohort (the kidneys recovered for transplantation), the PHS-IR kidneys accounted for 23.8% of total organ pool and 21.6% discarded kidneys, and, contrary to common perception, the PHS-IR kidneys experienced lower discard rates across all

HCV groups compared to the PHS-non-IR kidneys under same HCV categories. Lower discard rates in PHS-IR designated groups could be explained by their donor's younger age, lower KDPI scores, and a lower likelihood of undergoing procurement biopsies. However, when adjusting for factors associated with discard (using regression analysis), we found that PHS-IR designation is independently associated with increased discard risk in HCV Ab-/NAT- group, but not in HCV Ab+ and/or HCV NAT+ groups. HCV Ab+ and NAT+ designations seem to negate PHS-IR's relatively small effect on discard.

The American Society of Transplantation Consensus Conference on HCV donors and organ transplantation recently recommended that HCV Ab+ /NAT- donors (without other increased risk factors) not be considered at increased risk of HCV transmission.³³ A single-center study also demonstrated the safety of transplanting HCV Ab+ /NATdonor kidneys into HCV negative recipients. 16 Accordingly, our study documents the increased nationwide utilization of HCV Ab+ /NAT- donor kidneys in the last three years. This represents a pool of donors that is probably still underutilized, and so far has not resulted in a documented viral transmission and hence may not need antiviral Our study brings to light some challenging ethical dilemmas. Allocating HCV Ab or NAT+ organs to HCV Ab and NAT + recipients while bypassing HCV negative recipients is a thought-provoking concern when considering longer wait-list time for HCV negative recipients not willing to accept PHS organs, primarily due to lack of access or education on disease transmission. Whether HCV Ab or NAT+ organ utilization for HCV Ab or NAT+ recipients is more beneficial than for HCV Ab or NATrecipients, in terms of graft or patient survival and cost-effectiveness, remains to be seen. Education of public and private payers is crucial to help provide payment for initial HCV treatment as well as additional therapy should resistance be a challenge posttransplant (<5%); thus, we propose that every transplant institution establish an individual or a group of HCV champion providers tasked with education and consenting of patients, being a facilitator in negotiations with insurance carriers, and in-depthanalyzers of all outcomes of HCV Ab and/or NAT+ organs.

A recent analysis showed that transplanting viremic donor kidneys into negative recipients could be cost-effective with an incremental cost-effectiveness ratio of \$56,018

per quality-adjusted life-year (QALY) from the payer's perspective, and \$4,647 per QALY from the societal perspective, compared to remaining on the waitlist for one additional year. ³⁴ Also, Gupta et al. found that kidney transplants using HCV+ donors for HCV- recipients was a less costly approach (\$138,000 versus \$329,000) and resulted in slightly more years of life (YOL) (4.7 versus 4.8) when compared to HCV-donors for HCV- recipients.³⁵

According to a recent OPTN data analysis, overdose-deaths (N=63,632 in 2016) and overdose-death donors (N=1,804 in 2016), although accounting to a meager 3% of such deaths, continued to increase exponentially last several (the study period ended in 2016). Based on our analysis, the number of the HCV NAT+ kidneys doubled in four-years during our study period (from 658 in 2015 to 1,144 in 2018), and we would expect those kidneys to reach around 2,000 in 2023 assuming current trends in opioid use and related death rates remain unchanged. The rising trend for available HCV NAT+ kidneys to transplant is supported by a recent publication (our estimation of 344 vs. actual number of 374 for the first quarter of 2019).

Unfortunately, during the study period, the discards for HCV NAT+ kidneys were unacceptably high around 40%, and those kidneys carried a tenfold higher risk of discard. In era of DAA therapy curing HCV with >95% success and expectation of 10-15% of deceased donor pool originating from HCV NAT+ kidneys (based on our forecasting), an urgent policy changes are needed to tackle opioid epidemic, minimize discard with efficient allocation of those kidneys to the centers routinely using for HCV naive or infected recipients, mitigate PHS labeling effect, and disseminate evidence-based experience on this evolving topic.

Another issue with HCV viremic donors is that they are unlikely to be placed in younger recipients (longevity matched donor-recipient pairs, mainly allocation of KDPI<20% kidneys to young recipients with the longest post-transplant survival expectancy) due to adverse impact of HCV Ab positivity on KDPI calculation, even though they are otherwise good quality kidneys.³⁷ One UNOS study of Ab+/NAT-kidneys concludes that if such kidneys are considered to be HCV-, their survival would be comparable to the matched non-HCV-infected kidneys, less likely to be classified as

KDPI> 85%, and the risk of DGF was significantly lower when compared to non-HCV kidneys.³⁸ A recent companion study also confirms similar or superior short term outcomes from transplantation with HCV+ kidneys. ²¹ Hence some authors even question the need for including HCV Ab result with donor offers and KDPI calculations and recommend uniform utilization of NAT status alone. ³⁹⁻⁴¹

Strengths of our study include a large sample size of a national dataset. Limitations of our study include: 1) it is a retrospective registry data analysis without a control group; 2) the OPTN dataset does not include information regarding potential donors in whom a donor consent was not obtained and not recovered for transplant; 3) missing data can introduce bias; 4) reporting delays and labelling errors might happen.

In conclusion, HCV positive donors are likely to increase in near future years, unless there is a dramatic reduction in the current opioid crisis. Organs from HCV positive donors could potentially expand the organ pool, especially given the effective antiviral therapy available against HCV, and increase access to transplant across all patients, including HCV negative recipients. PHS labeling effect on discard requires rediscussion of purpose and utility of classification. We predict that as center level and patient comfort level spreads in accepting HCV viremic donors, HCV NAT+ organ utilization will increase significantly soon, similar to the increasing trend observed in HCV Ab+/NAT- organ utilization in the last three years.

References:

- Wolfe RA, Ashby VB, Milford EL, et al. Comparison of mortality in all patients on dialysis, patients on dialysis awaiting transplantation, and recipients of a first cadaveric transplant. N Engl J Med. Dec 2 1999;341(23):1725-1730.
- 2. Schnitzler MA, Lentine KL, Burroughs TE. The cost effectiveness of deceased organ donation. *Transplantation*. Dec 15 2005;80(11):1636-1637.
- **3.** Hart A, Smith JM, Skeans MA, et al. OPTN/SRTR 2016 Annual Data Report: Kidney. *Am J Transplant*. Jan 2018;18 Suppl 1:18-113.

- **4.** Schold JD, Segev DL. Increasing the pool of deceased donor organs for kidney transplantation. *Nat Rev Nephrol.* Mar 27 2012;8(6):325-331.
- **5.** Bowring MG, Holscher CM, Zhou S, et al. Turn down for what? Patient outcomes associated with declining increased infectious risk kidneys. *Am J Transplant*. Mar 2018;18(3):617-624.
- Martins SS, Sarvet A, Santaella-Tenorio J, Saha T, Grant BF, Hasin DS. Changes in US Lifetime Heroin Use and Heroin Use Disorder: Prevalence From the 2001-2002 to 2012-2013 National Epidemiologic Survey on Alcohol and Related Conditions. *JAMA Psychiatry*. May 1 2017;74(5):445-455.
- Jones CM, Logan J, Gladden RM, Bohm MK. Vital Signs: Demographic and Substance Use Trends Among Heroin Users United States, 2002-2013. MMWR Morb Mortal Wkly Rep. Jul 10 2015;64(26):719-725.
- **8.** Durand CM, Bowring MG, Thomas AG, et al. The Drug Overdose Epidemic and Deceased-Donor Transplantation in the United States: A National Registry Study. *Ann Intern Med.* May 15 2018;168(10):702-711.
- **9.** Gonzalez SA, Trotter JF. The rise of the opioid epidemic and hepatitis C-positive organs: A new era in liver transplantation. *Hepatology*. Apr 2018;67(4):1600-1608.
- **10.** Goldberg DS, Blumberg E, McCauley M, Abt P, Levine M. Improving Organ Utilization to Help Overcome the Tragedies of the Opioid Epidemic. *Am J Transplant*. Oct 2016;16(10):2836-2841.
- Abara WE, Collier MG, Moorman A, et al. Characteristics of Deceased Solid Organ Donors and Screening Results for Hepatitis B, C, and Human Immunodeficiency Viruses United States, 2010-2017. MMWR Morb Mortal Wkly Rep. Jan 25 2019;68(3):61-66.
- 12. Kling CE, Perkins JD, Landis CS, Limaye AP, Sibulesky L. Utilization of Organs From Donors According to Hepatitis C Antibody and Nucleic Acid Testing Status: Time for Change. *Am J Transplant*. Nov 2017;17(11):2863-2868.
- **13.** Kucirka LM, Singer AL, Ros RL, Montgomery RA, Dagher NN, Segev DL. Underutilization of hepatitis C-positive kidneys for hepatitis C-positive recipients. *Am J Transplant*. May 2010;10(5):1238-1246.
- **14.** Reese PP, Abt PL, Blumberg EA, Goldberg DS. Transplanting Hepatitis C-Positive Kidneys. *N Engl J Med.* Jul 23 2015;373(4):303-305.
- **15.** Bowring MG, Kucirka LM, Massie AB, et al. Changes in Utilization and Discard of Hepatitis C-Infected Donor Livers in the Recent Era. *Am J Transplant*. Feb 2017;17(2):519-527.

- de Vera ME, Volk ML, Ncube Z, et al. Transplantation of hepatitis C virus (HCV) antibody positive, nucleic acid test negative donor kidneys to HCV negative patients frequently results in seroconversion but not HCV viremia. *Am J Transplant*. Jul 24 2018.
- Sageshima J, Troppmann C, McVicar JP, Santhanakrishnan C, de Mattos AM, Perez RV. Impact of Willingness to Accept Hepatitis C Seropositive Kidneys Among Hepatitis C RNA-Positive Waitlisted Patients. *Transplantation*. Jul 2018;102(7):1179-1187.
- **18.** Durand CM, Bowring MG, Brown DM, et al. Direct-Acting Antiviral Prophylaxis in Kidney Transplantation From Hepatitis C Virus-Infected Donors to Noninfected Recipients: An Open-Label Nonrandomized Trial. *Ann Intern Med.* Mar 6 2018.
- 19. Reese PP, Abt PL, Blumberg EA, et al. Twelve-Month Outcomes After Transplant of Hepatitis C-Infected Kidneys Into Uninfected Recipients: A Single-Group Trial. *Ann Intern Med.* Sep 4 2018;169(5):273-281.
- 20. Morales JM, Campistol JM, Dominguez-Gil B, et al. Long-term experience with kidney transplantation from hepatitis C-positive donors into hepatitis C-positive recipients. *Am J Transplant*. Nov 2010;10(11):2453-2462.
- 21. La Hoz RM, Sandikci B, Ariyamuthu VK, Tanriover B. Short-Term Outcomes of Deceased Donor Renal Transplants of HCV Uninfected Recipients from HCV Seropositive Non-viremic donors and Viremic Donors in the Era of Direct-Acting Antivirals. *Am J Transplant*. Jun 17 2019.
- **22.** Chow EK, Massie AB, Muzaale AD, et al. Identifying appropriate recipients for CDC infectious risk donor kidneys. *Am J Transplant*. May 2013;13(5):1227-1234.
- 23. Seem DL, Lee I, Umscheid CA, Kuehnert MJ, United States Public Health S. PHS guideline for reducing human immunodeficiency virus, hepatitis B virus, and hepatitis C virus transmission through organ transplantation. *Public Health Rep.* Jul 2013;128(4):247-343.
- **24.** Theodoropoulos N, Nowicki MJ, Chinchilla-Reyes C, et al. Deceased organ donor screening for human immunodeficiency virus, hepatitis B virus and hepatitis C virus: Discordant serology and nucleic acid testing results. *Transpl Infect Dis.* Feb 2018;20(1).
- 25. Marrero WJ, Naik AS, Friedewald JJ, et al. Predictors of Deceased Donor Kidney Discard in the United States. *Transplantation*. Jul 2017;101(7):1690-1697.
- 26. 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*. Mar 2017;101(3):575-587.

- **27.** Fernandez HE, Chiles MC, Pereira M, et al. Outcomes for potential kidney transplant recipients offered public health service increased risk kidneys: A single-center experience. *Clin Transplant*. Dec 2018;32(12):e13427.
- 28. Lonze BE, Dagher NN, Liu M, et al. Outcomes of renal transplants from Centers for Disease Control and Prevention high-risk donors with prospective recipient viral testing: a single-center experience. *Arch Surg.* Nov 2011;146(11):1261-1266.
- **29.** Reese PP, Feldman HI, Asch DA, et al. Transplantation of kidneys from donors at increased risk for blood-borne viral infection: recipient outcomes and patterns of organ use. *Am J Transplant*. Oct 2009;9(10):2338-2345.
- **30.** Freeman RB, Cohen JT. Transplantation risks and the real world: what does 'high risk' really mean? *Am J Transplant*. Jan 2009;9(1):23-30.
- **31.** Duan KI, Englesbe MJ, Volk ML. Centers for Disease Control 'high-risk' donors and kidney utilization. *Am J Transplant*. Feb 2010;10(2):416-420.
- **32.** Reese PP, Harhay MN, Abt PL, Levine MH, Halpern SD. New Solutions to Reduce Discard of Kidneys Donated for Transplantation. *J Am Soc Nephrol*. Apr 2016;27(4):973-980.
- **33.** Levitsky J, Formica RN, Bloom RD, et al. The American Society of Transplantation Consensus Conference on the Use of Hepatitis C Viremic Donors in Solid Organ Transplantation. *Am J Transplant*. Nov 2017;17(11):2790-2802.
- **34.** Kadatz M, Klarenbach S, Gill J, Gill JS. Cost-effectiveness of using kidneys from hepatitis C nucleic acid test-positive donors for transplantation in hepatitis C-negative recipients. *Am J Transplant*. Oct 2018;18(10):2457-2464.
- **35.** Gupta G, Zhang Y, Carroll NV, Sterling RK. Cost-effectiveness of hepatitis C-positive donor kidney transplantation for hepatitis C-negative recipients with concomitant direct-acting antiviral therapy. *Am J Transplant*. Oct 2018;18(10):2496-2505.
- **36.** Potluri VS, Goldberg DS, Mohan S, et al. National Trends in Utilization and 1-Year Outcomes with Transplantation of HCV-Viremic Kidneys. *Journal of the American Society of Nephrology : JASN.* Oct 2019;30(10):1939-1951.
- **37.** Sibulesky L, Kling CE, Limaye AP, Johnson CK. Is Kidney Donor Profile Index (KDPI) Valid for Hepatitis C Aviremic Kidneys? *Ann Transplant*. Nov 6 2017;22:663-664.
- **38.** Sibulesky L, Kling CE, Blosser C, et al. Are we underestimating the quality of aviremic hepatitis C-positive kidneys? Time to reconsider. *Am J Transplant*. Feb 16 2018.

- **39.** Anesi JA, Goldberg DS. Maximizing Utilization of the Donor Pool by Appropriate Classification of Hepatitis C Antibody-Positive Donors. *Am J Transplant*. Nov 2017;17(11):2757-2758.
- **40.** Sibulesky L, Limaye AP. Hepatitis C NAT status in the UNOS database. *Am J Transplant*. Jun 2019;19(6):1870.
- 41. Cannon RM, Locke JE, Orandi BJ, et al. Impact of donor hepatitis C virus on kidney transplant outcomes for hepatitis C positive recipients in the direct acting antiviral era: time to revise the kidney donor risk index? *Transplantation*. Sep 9 2019.

Author Contribution:

VKA participated in study design and manuscript writing.

BS participated in study design, data analysis and manuscript writing.

NA participated in manuscript writing.

CH participated in manuscript writing.

MPM participated in manuscript writing.

RP participated in manuscript writing.

AA participated in data analysis.

BT participated in study design, data analysis and manuscript writing.

Figure Legends:

Figure 1: Flow chart of deceased donors registered in the UNOS database between January 1, 2015, and December 31, 2018, in the US.

Figure 2: Disposition of deceased donor kidneys based on HCV Ab and NAT status between January 1, 2015, and Dec 31, 2018, in the U.S.

Figure 3: KDPI distribution by HCV status and disposition (transplant vs. discard) between January 1, 2015, and December 31, 2018, in the U.S. (dashed and solid

vertical lines indicate median KDPI for transplanted and discarded kidneys, respectively).

Figure 4: Geographic distribution by the UNOS Region for hepatitis C virus (HCV) infected donors and transplants with HCV infected kidneys.

Figure 5: Geographic distribution of hepatitis C virus (HCV) infected (NAT+) kidney transplantation by the organ procurement organization (OPO) donation service area (DSA) between 2015 and 2018.

Figure 6: Geographic distribution of hepatitis C virus (HCV) infected (NAT+) kidney transplantation by transplant center between 2015 and 2018.

Figure 7: Forecasting number of potential HCV NAT+ kidneys by the year 2023 (computed using time series trend adjusted exponential smoothing method).

Table 1: Characteristics of deceased donors by HCV status between January 1, 2015 and December 31, 2018 in the U.S.

							p-values±	
						Ab-,N	AT-vs	Ab+,NAT-vs
	All groups	Ab-, NAT-	Ab+, NAT-	Ab±, NAT+	All ways	Ab+, NAT-	Ab±, NAT+	
								Ab±, NAT+
n [¥]	38702	35861	973	1868				
Age (years), Mean ± SD	40.0 ± 17.3	40.0 ± 17.7	41.6 ± 12.8	37.3 ± 11.1	< 0.001 ***	0.026*	< 0.001 ***	< 0.001 ***
Gender, n (%)					< 0.001 ***	< 0.001 ***	< 0.001 ***	< 0.001 ***
Female	15366 (39.7)	14243 (39.7)	463 (47.6)	660 (35.3)				
Male	23336 (60.3)	21618 (60.3)	510 (52.4)	1208 (64.7)				
Race, n (%)					< 0.001 ***	< 0.001 ***	< 0.001 ***	0.863
White	25543 (66.0)	23266 (64.9)	788 (81.0)	1489 (79.7)				
Black	6143 (15.9)	5886 (16.4)	79 (8.1)	178 (9.5)				
Hispanic	5295 (13.7)	5047 (14.1)	87 (8.9)	161 (8.6)				
Other	1721 (4.4)	1662 (4.6)	19 (2.0)	40 (2.1)				
BMI (kg/m²), Mean ± SD	28.0 ± 7.3	28.0 ± 7.4	28.2 ± 6.4	26.7 ± 5.4	< 0.001 ***	0.067	< 0.001 ***	< 0.001 ***
Blood type, n (%)					< 0.001 ***	0.022*	0.038*	0.761
0	18579 (48.0)	17152 (47.8)	499 (51.3)	928 (49.7)				
Α	14300 (37.0)	13235 (36.9)	357 (36.7)	708 (37.9)				
В	4522 (11.7)	4224 (11.8)	99 (10.2)	199 (10.7)				
AB	1299 (3.4)	1248 (3.5)	18 (1.8)	33 (1.8)				
Region of recovery, n (%)					< 0.001 ***	< 0.001 ***	< 0.001 ***	0.281
1	1393 (3.6)	1190 (3.3)	65 (6.7)	138 (7.4)				
2	4828 (12.5)	4238 (11.8)	217 (22.3)	373 (20.0)				
3	5965 (15.4)	5538 (15.4)	143 (14.7)	284 (15.2)				

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4	4141 (10.7)	3961 (11.0)	62 (6.4)	118 (6.3)				
5	5817 (15.0)	5538 (15.4)	103 (10.6)	176 (9.4)				
6	1544 (4.0)	1480 (4.1)	26 (2.7)	38 (2.0)				
7	3103 (8.0)	2977 (8.3)	48 (4.9)	78 (4.2)				
8	2735 (7.1)	2597 (7.2)	37 (3.8)	101 (5.4)				
9	1700 (4.4)	1566 (4.4)	59 (6.1)	75 (4.0)				
10	3382 (8.7)	3030 (8.4)	111 (11.4)	241 (12.9)				
11	4094 (10.6)	3746 (10.4)	102 (10.5)	246 (13.2)				
DCD, n (%)					< 0.001 ***	< 0.001 ***	< 0.001 ***	0.969
No	31705 (81.9)	29181 (81.4)	862 (88.6)	1662 (89.0)				
Yes	6997 (18.1)	6680 (18.6)	111 (11.4)	206 (11.0)				

Abbreviations: Ab: Antibody; BMI: Body Mass Index; DCD: Donation after Cardiac Death; ECD: Extended Criteria Donor; HCV: Hepatitis C Virus; KDPI: Kidney Donor Profile Index; NA: No data Available; NAT: Nucleic Acid Testing; PHS: Public Health Service; SD: Standard Deviation.

±ALL-ways comparisons p-value from chi-squared test for categorical variables and Kruskal-Wallis rank test for numerical variables, both adjusted by Holm's method for multiple pairwise testing

Table 1: Characteristics of deceased donors by HCV status — continued from previous page

			p-values [±]						
									Ab+, NAT- vs.
n [*]		All groups	Ab-, NAT-	Ab+, NAT-	Ab±, NAT+	All ways	Ab+, NAT-	Ab±, NAT+	
									Ab±, NAT+
ECD, n (%)						< 0.001 ***	0.007**	< 0.001 ***	< 0.001 ***
No		30263 (78.2)	27774 (77.4)	795 (81.7)	1694 (90.7)				
Yes		8439 (21.8)	8087 (22.6)	178 (18.3)	174 (9.3)				

^{*}n: number of records in each group. Missing/unknown values in any particular variable are ignored when reporting summary statistics.

Diabetes (any type), n (%)					< 0.001 ***	0.642	< 0.001 ***	0.002**
No	33862 (88.1)	31282 (87.8)	855 (88.8)	1725 (93.4)				
Yes	4581 (11.9)	4351 (12.2)	108 (11.2)	122 (6.6)				
Hypertension, n (%)					< 0.001 ***	0.843	< 0.001 ***	< 0.001 ***
No	25087 (65.3)	23026 (64.7)	631 (65.6)	1430 (77.7)				
Yes	13313 (34.7)	12572 (35.3)	331 (34.4)	410 (22.3)				
PHS increased risk, n (%)					< 0.001 ***	< 0.001 ***	< 0.001 ***	< 0.001 ***
No O)	28879 (74.6)	28225 (78.7)	294 (30.2)	360 (19.3)				
Yes	9815 (25.4)	7628 (21.3)	679 (69.8)	1508 (80.7)				
Cause of death, n (%)					< 0.001 ***	< 0.001 ***	< 0.001 ***	0.624
Anoxia	15937 (41.2)	14068 (39.2)	643 (66.1)	1226 (65.6)				
Cerebrovascular	10651 (27.5)	10196 (28.4)	173 (17.8)	282 (15.1)				
Head Trauma	10945 (28.3)	10488 (29.2)	137 (14.1)	320 (17.1)				
Other	1169 (3.0)	1109 (3.1)	20 (2.1)	40 (2.1)				
KDPI (%), Mean ± SD	53.6 ± 29.6	52.9 ± 30.0	68.5 ± 21.9	60.7 ± 21.3	< 0.001 ***	< 0.001 ***	< 0.001 ***	< 0.001 ***

Significance codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 ' ' 1

Abbreviations: Ab: Antibody; BMI: Body Mass Index; DCD: Donation after Cardiac Death; ECD: Extended Criteria Donor; HCV: Hepatitis C Virus; KDPI: Kidney Donor Profile Index;

NA: No data Available; NAT: Nucleic Acid Testing; PHS: Public Health Service; SD: Standard Deviation.

[¥]n: number of records in each group. Missing/unknown values in any particular variable are ignored when reporting summary statistics.

[±]ALL-ways comparisons p-value from chi-squared test for categorical variables and Kruskal-Wallis rank test for numerical variables; pairwise comparisons p-value from chi-squared test for categorical variables and Wilcoxon rank test for numerical variables, both adjusted by Holm's method for multiple pairwise testing

Table 2: Characteristics of transplanted deceased donor kidneys by HCV status between January 1, 2015 and December 31, 2018 in the U.S.

							p-values [±]	
						Ab-,	NAT- vs.	Ab+, NAT-
	All groups	Ab-, NAT-	Ab+, NAT-	Ab±, NAT+	All ways	Ab+, NAT-	Ab±, NAT+	Ab±, NA
n	56833	54232	968	1633				
Age (years), Mean ± SD	36.3 ± 15.9	36.3 ± 16.2	37.2 ± 10.8	32.9 ± 8.4	< 0.001 ***	0.163	< 0.001 ***	< 0.001 *
Gender, n (%)					< 0.001 ***	< 0.001 ***	0.199	< 0.001 *
Female	21664 (38.1)	20617 (38.0)	462 (47.7)	585 (35.8)				
Male	35169 (61.9)	33615 (62.0)	506 (52.3)	1048 (64.2)				
Race, n (%)					< 0.001 ***	< 0.001 ***	< 0.001 ***	0.4
White	38147 (67.1)	35927 (66.2)	842 (87.0)	1378 (84.4)				
Black	8020 (14.1)	7910 (14.6)	34 (3.5)	76 (4.7)				
Hispanic	8148 (14.3)	7930 (14.6)	68 (7.0)	150 (9.2)				
Other	2518 (4.4)	2465 (4.5)	24 (2.5)	29 (1.8)				
BMI (kg/m²), Mean ± SD	27.5 ± 7.1	27.6 ± 7.1	28.1 ± 6.3	26.1 ± 5.1	< 0.001 ***	< 0.001 ***	< 0.001 ***	< 0.001 *
Blood type, n (%)					< 0.001 ***	0.010 **	0.005 **	0.9
0	27269 (48.0)	25915 (47.8)	503 (52.0)	851 (52.1)				
A	21079 (37.1)	20184 (37.2)	347 (35.8)	548 (33.6)				
В	6629 (11.7)	6308 (11.6)	105 (10.8)	216 (13.2)				
AB	1856 (3.3)	1825 (3.4)	13 (1.3)	18 (1.1)		+		
DCD, n (%)	1000 (510)	1025 (511)	15 (110)	10 (111)	< 0.001 ***	< 0.001 ***	< 0.001 ***	0.2
No	45931 (80.8)	43590 (80.4)	854 (88.2)	1487 (91.1)				
Yes	10902 (19.2)	10642 (19.6)	114 (11.8)	146 (8.9)				
ECD, n (%)	,	(/	(-/	- (/	< 0.001 ***	< 0.001 ***	< 0.001 ***	0.00
No	49968 (87.9)	47446 (87.5)	909 (93.9)	1613 (98.8)				
Yes	6865 (12.1)	6786 (12.5)	59 (6.1)	20 (1.2)				
Diabetes (any type), n (%)	,	(,	,	- ,	< 0.001 ***	0.431	< 0.001 ***	< 0.001
No	52910 (93.6)	50398 (93.5)	909 (94.5)	1603 (98.8)				
Yes	3600 (6.4)	3527 (6.5)	53 (5.5)	20 (1.2)				
Hypertension, n (%)	, , ,	(/	(/	- (/	< 0.001 ***	0.441	< 0.001 ***	< 0.001
No	42259 (74.9)	40073 (74.4)	731 (76.2)	1455 (90.0)				
Yes	14166 (25.1)	13776 (25.6)	228 (23.8)	162 (10.0)				
PHS increased risk, n (%)					< 0.001 ***	< 0.001 ***	< 0.001 ***	< 0.001
No	43016 (75.7)	42554 (78.5)	243 (25.1)	219 (13.4)				
Yes	13809 (24.3)	11670 (21.5)	725 (74.9)	1414 (86.6)				
Cause of death, n (%)		,	• •	. ,	< 0.001 ***	< 0.001 ***	< 0.001 ***	0.4
Anoxia	23267 (40.9)	21417 (39.5)	691 (71.4)	1159 (71.0)				
Cerebrovascular	12864 (22.6)	12622 (23.3)	113 (11.7)	129 (7.9)				
Head Trauma	18908 (33.3)	18446 (34.0)	147 (15.2)	315 (19.3)				
Other	1794 (3.2)	1747 (3.2)	17 (1.8)	30 (1.8)				
KDPI (%), Mean ± SD	44.8 ± 27.0	44.4 ± 27.3	58.7 ± 19.7	49.8 ± 16.8	< 0.001 ***	< 0.001 ***	< 0.001 ***	< 0.001

Abbreviations: Ab: Antibody; BMI: Body Mass Index; DCD: Donation after Cardiac Death; ECD: Extended Criteria Donor; HCV: Hepatitis C Virus; KDPI: Kidney Donor Profile Index; NA: No data Available; NAT: Nucleic Acid Testing; PHS: Public Health Service; SD: Standard Deviation.

n : number of records in each group. Missing/unknown values in any particular variable are ignored when reporting summary statistics.

ALL-ways comparisons p-value from chi-squared test for categorical variables and Kruskal-Wallis rank test for numerical variables; pairwise comparisons p-value from chi-squared test for categorical variables and Wilcoxon rank test for numerical variables, both adjusted by Holm's method for multiple pairwise testing

Table 3: Characteristics of discarded deceased donor kidneys by HCV status between January 1, 2015 and December 31, 2018 in the U.S.

							p-values [±]	
$\overline{}$							NAT- vs.	Ab-, NAT- vs.
	All groups	Ab-, NAT-	Ab+, NAT-	Ab±, NAT+	All ways	Ab+, NAT-	Ab±, NAT+	
n [¥]	13617	11992	491	1134				
	505.460	510:100	40.0 : 40.0	20.5 : 44.4				
Age (years), Mean ± SD	50.5 ± 16.0	51.9 ± 16.0	43.8 ± 12.9	38.6 ± 11.1	< 0.001 ***	< 0.001 ***	< 0.001 ***	< 0.001 ***
Gender, n (%)					< 0.001 ***	0.328	< 0.001 ***	< 0.001 ***
Female	6208 (45.6)	5573 (46.5)	245 (49.9)	390 (34.4)			3.332	
Male	7409 (54.4)	6419 (53.5)	246 (50.1)	744 (65.6)				
Race, n (%)	, ,	` '	` '	,	< 0.001 ***	< 0.001 ***	< 0.001 ***	0.832
White	9108 (66.9)	7786 (64.9)	394 (80.2)	928 (81.8)				
Black	2350 (17.3)	2221 (18.5)	40 (8.1)	89 (7.8)				
Hispanic	1565 (11.5)	1427 (11.9)	49 (10.0)	89 (7.8)				
Other	594 (4.4)	558 (4.7)	8 (1.6)	28 (2.5)				
BMI (kg/m²), Mean ± SD	29.4 ± 7.6	29.6 ± 7.8	28.1 ± 6.3	27.2 ± 5.5	< 0.001 ***	< 0.001 ***	< 0.001 ***	0.007**
Blood type, n (%)					< 0.001 ***	0.181	0.004 **	0.938
0	6401 (47.0)	5595 (46.7)	242 (49.3)	564 (49.7)				
A	5069 (37.2)	4439 (37.0)	188 (38.3)	442 (39.0)				
В	1621 (11.9)	1452 (12.1)	53 (10.8)	116 (10.2)				
AB	522 (3.8)	502 (4.2)	8 (1.6)	12 (1.1)				
DCD, n (%)			- · · · · · ·	,,	0.940	0.999	0.941	0.974
No	10802 (79.3)	9517 (79.4)	390 (79.4)	895 (78.9)				
Yes	2815 (20.7)	2475 (20.6)	101 (20.6)	239 (21.1)				
ECD, n (%)				,	< 0.001 ***	< 0.001 ***	< 0.001 ***	< 0.001 ***
No	7028 (51.6)	5596 (46.7)	385 (78.4)	1047 (92.3)				
Yes	6589 (48.4)	6396 (53.3)	106 (21.6)	87 (7.7)				
Diabetes (any type), n (%)		, , , , , , , , , , , , , , , , , , , ,		, ,	< 0.001 ***	< 0.001 ***	< 0.001 ***	0.008**
No	10293 (76.4)	8816 (74.2)	420 (87.3)	1057 (94.5)				
Yes	3182 (23.6)	3060 (25.8)	61 (12.7)	61 (5.5)				
Hypertension, n (%)					< 0.001 ***	< 0.001 ***	< 0.001 ***	< 0.001 ***
No	5519 (41.0)	4377 (36.8)	290 (60.2)	852 (76.9)				
Yes	7957 (59.0)	7509 (63.2)	192 (39.8)	256 (23.1)				
PHS increased risk, n (%)					< 0.001 ***	< 0.001 ***	< 0.001 ***	< 0.001 ***
No	10675 (78.4)	10241 (85.4)	184 (37.5)	250 (22.0)				
Yes	2936 (21.6)	1745 (14.6)	307 (62.5)	884 (78.0)				
Cause of death, n (%)					< 0.001 ***	< 0.001 ***	< 0.001 ***	0.795
Anoxia	5231 (38.4)	4259 (35.5)	278 (56.6)	694 (61.2)				
Cerebrovascular	5850 (43.0)	5536 (46.2)	121 (24.6)	193 (17.0)				
Head Trauma	2162 (15.9)	1868 (15.6)	78 (15.9)	216 (19.0)				
Other	374 (2.7)	329 (2.7)	14 (2.9)	31 (2.7)				
KDPI (%), Mean ± SD	76.8 ± 21.6	78.1 ± 21.5	74.2 ± 20.0	64.1 ± 19.5	< 0.001 ***	< 0.001 ***	< 0.001 ***	< 0.001 ***

Significance codes: 0 0.001 '**' 0.01 '*' 0.05 ' ' 1

Abbreviations: Ab: Antibody; BMI: Body Mass Index; DCD: Donation after Cardiac Death; ECD: Extended Criteria Donor; HCV: Hepatitis C Virus; KDPI: Kidney Donor Profile Index; NA: No data Available; NAT: Nucleic Acid Testing; PHS: Public Health Service; SD: Standard Deviation. n¥: number of records in each group. Missing/unknown values in any particular variable are ignored when reporting summary statistics.

*ALL-ways comparisons p-value from chi-squared test for categorical variables and Kruskal-Wallis rank test for numerical variables; pairwise comparisons p-value from chi-squared test for categorical variables and Wilcoxon rank test for numerical variables, both adjusted by Holm's method for multiple pairwise testing

Table 4: Characteristics of discarded deceased donor kidneys by HCV status between January 1, 2015 and December 31, 2018 in the U.S.

						p-values±		
+						Ab-	, NAT- vs	Ab-, NAT- vs
	All groups	Ab-, NAT-	Ab+, NAT	Ab±, NAT+	All ways	Ab+, NAT-	Ab±, NAT+	Ab±, NAT+
Disposition reason, n [¥] (%)					< 0.001 ***	< 0.001 ***	< 0.001 ***	0.079
AKI	784 (5.8)	722 (6.0)	26 (5.3)	36 (3.2)				
Anatomical abnormalities	772 (5.7)	723 (6.0)	23 (4.7)	26 (2.3)				
Biopsy	3943 (29.0)	3768 (31.4)	57 (11.6)	118 (10.4)				
CIT	294 (2.2)	262 (2.2)	5 (1.0)	27 (2.4)				
Diseased organ	400 (2.9)	385 (3.2)	1 (0.2)	14 (1.2)				
Donor quality	333 (2.4)	273 (2.3)	20 (4.1)	40 (3.5)				
Donor social history	19 (0.1)	4 (0.0)	3 (0.6)	12 (1.1)				
HCV	120 (0.9)	8(0.1)	40 (8.1)	72 (6.3)				
Infection	44 (0.3)	39 (0.3)	2 (0.4)	3 (0.3)				
No recipient located/list exhausted	5536 (40.7)	4496 (37.5)	298 (60.7)	742 (65.4)				
Non-renal cancer	102 (0.7)	101 (0.8)	7 (1.4)	1 (0.1)				
Organ trauma during recovery	521 (3.8)	499 (4.2)	2 (0.4)	15 (1.3)				
Pump	335 (2.5)	327 (2.7)	1 (0.2)	8 (0.7)				
Renal cancer	35 (0.3)	32 (0.3)	2 (0.4)	1 (0.1)				
Turned down in the OR	62 (0.5)	60 (0.5)	4 (0.8)	2 (0.2)				
Vascular disease	145 (1.1)	141 (1.2)	NA (NA)	3 (0.3)				
Warm ischemia time	66 (0.5)	62 (0.5)	NA (NA)	2 (0.2)				
Other	106 (0.8)	90 (0.8)	NA (NA)	12 (1.1)				

Significance codes: 0 0.001 '**' 0.01 '*' 0.05 ' ' 1

Abbreviations: Ab: Antibody; BMI: Body Mass Index; DCD: Donation after Cardiac Death; ECD: Extended Criteria Donor; HCV: Hepatitis C Virus; KDPI: Kidney Donor Profile Index; NA: No data Available; NAT: Nucleic Acid Testing; PHS: Public Health Service; SD: Standard Deviation. n[¥]: number of records in each group. Missing/unknown values in any particular variable are ignored when reporting summary statistics.

^{*}ALL-ways comparisons p-value from chi-squared test for categorical variables and Kruskal-Wallis rank test for numerical variables; pairwise comparisons p-value from chi-squared test for categorical variables and Wilcoxon rank test for numerical variables, both adjusted by Holm's method for multiple pairwise testing

Table 5: Kidney discards by HCV groups and PHS designations and predicting discard in recovered kidneys for transplantation using multivariable mixed-effects logistic regression models.

All Cohort†		HCV Ab	o–, NAT– (n=66,2	224)	HCV A	b+, NAT- (n=1,	,459)	HCV Ab±, NAT+ (n=2,767)			
(n = 70, 450)											
		PHS IR	PHS non IR	p-value	PHS IR	PHS non IR	p-value	PHS IR	PHS non IR	p-value	
PHS‡, n (%)	16739 (23.8)	13411 (20.3)	52787 (79.7)		1030 (70.7)	427 (29.3)		2298 (83.0)	469 (17.0)		
Age (years), Mean ± SD	39.0 ± 16.9	34.6 ± 13.3	40.3 ± 17.9	<0.001	34.5 ± 11.3	48.9 ± 11.2	<0.001	33.5 ± 8.8	43.8 ± 11.2	< 0.001	
KDPI %, Mean ± SD	51.0 ± 29.0	38.9 ± 25.7	53.4 ± 29.5	<0.001	57.8 ± 18.8	78.8 ± 19.1	<0.001	52.8 ± 17.6	69.7 ± 20.8	< 0.001	
Biopsy of either kidney, n (%)	37863 (53.8)	6361 (47.4)	29087 (55.1)	<0.001	552 (53.9)	329 (77.1)	<0.001	1215 (52.9)	313 (66.7)	< 0.001	
GS > 20%, n (%)	5266 (13.9)	671 (10.6)	4423 (15.2)	<0.001	38 (6.9)	46 (14)	<0.001	62 (5.1)	26 (8.3)	0.03	
IF moderate or severe, n (%)	3326 (8.8)	423 (6.7)	2774 (9.5)	<0.001	30 (5.4)	30 (9.1)	0.04	44 (3.6)	25 (8.0)	0.001	
Discard rate, n (%)	13617 (19.3)	1745 (13.0)	10241 (19.4)	<0.001	307 (29.8)	184 (43.1)	<0.001	884 (38.5)	250 (53.3)	< 0.001	
		Estimate	ed odds ratios fo	r discard and	d 95% confider	nce intervals§					
PHS designation	1.20 (1.15-	1.24 (1.15-	Reference	<0.001	1.24 (0.84-	Reference	0.28	1.04 (0.79-	Reference	0.78	
	1.29)	1.34)			1.83)			1.38)			
HCV designation	2.29 (2.15-	Refer	rence		2.07 (1.	78-2.40)	< 0.001	5.21 (4.	.62-5.89)	< 0.001	
7	2.43)										

Abbreviations: Abbreviations: Ab = Antibody; GS = Glomerulosclerosis; HCV = Hepatitis C virus; IF = Interstitial fibrosis; KDPI = Kidney profile risk index; NAT

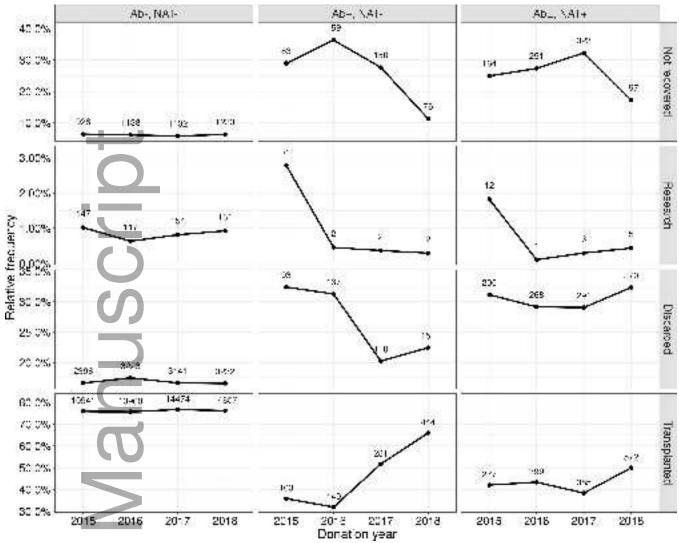
†All cohort includes the recovered kidneys, either transplanted or discarded, excludes the ones used for research.

‡PHS status is missing for 26 patients in the HCV Ab-, NAT- group and 2 patients in the HCV Ab+, NAT- group.

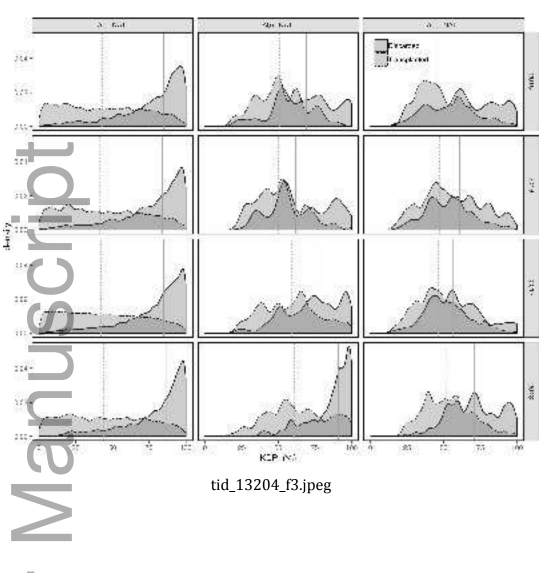
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⁼ Nucleic acid testing; PHS IR = Public health service increased risk; SD: Standard Deviation.

§Multivariable two-level mixed-effects logistic regression models (the second level of the model defining different intercepts for each United Network for Organ Sharing (UNOS) Region in a random-intercept model to account for regional variations in discards) were estimated to predict discard. The models were adjusted f or previously identified donor factors in the literature, including donor age >50 or not, either kidney biopsied, if biopsied glomerulosclerosis >20% or not, cytomegalovirus (CMV) status, cause death due to cerebrovascular accident (CVA), donation after cardiac death (DCD) status, height, weight, history of tattoo, KDPI, either kidney pumped, cold ischemia time, hepatitis B core antibody status, hepatitis B surface antigen status, history of diabetes, history of hypertension, history of cocaine use, history of IV drug use (IVDU), terminal creatinine >1.5 mg/dl or not, ABO blood type, and race (White, African American, Hispanic, Asian, and others), and transplant year.



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