Liver transplantation has saved almost 500,000 life-years in the United States since 1987, with 30% of patients undergoing liver transplantation for hepatitis C virus (HCV)-related liver disease. Approximately 0.77% of the United States population harbors HCV RNA and twice as many patients have HCV-specific antibodies indicating prior infection. Until recently, the only treatment to eradicate HCV infection consisted of interferon (IFN) plus ribavirin, which was successful in only a minority of patients and had significant treatment-limiting side effects. Recurrence of HCV is universal after liver transplantation; for patients with sufficient follow-up, nearly all of those transplanted for HCV demonstrated biopsy proven cirrhosis within 5 years. Additionally, early HCV cholestatic recurrence, which limits graft survival, has historically affected up to 10% of liver transplants for HCV. The recurrence of HCV additionally resulted in significantly decreased graft and patient survival compared with liver transplantation for other indications, including hepatitis B virus (HBV), alcoholic liver disease (ALD), and nonalcoholic steatohepatitis (NASH). Liver transplantation for HCV thus prolonged recipients’ lives but ultimately did not cure them of liver disease.
MATERIALS AND METHODS

Database, Inclusion Criteria, and Data Encoding

This study used data from the Scientific Registry of Transplant Recipients (SRTR). The SRTR data system includes data on all donor, waitlisted candidates, and transplant recipients in the United States, submitted by members of the Organ Procurement and Transplantation Network (OPTN). The Health Resources and Services Administration, US Department of Health and Human Services provides oversight to the activities of the OPTN and SRTR contractors. After institutional review board approval, SRTR Standard Analysis Files (June 2017 release) transplant records were linked with candidate, donor and follow-up data elements.

Records were identified for adult (age ≥18 years) deceased donor whole liver transplant recipients based on SRTR-defined primary diagnoses and classified as: (1) HCV (Ahn type C, Cirrhosis type C, and Alcohol cirrhosis with HCV), (2) HBV (Ahn type B ABSAg+ and cirrhosis type B HBSAg+), (3) NASH (cirrhosis fatty liver), (4) ALD (alcoholic cirrhosis), and (5) other. Antiviral era was classified based on transplant date and stratified using the approach of Flemming et al as: IFN (January 2003 to December 2010), PI (January 2011 to December 2013), and DAA (January 2014 to May 2017). An additional inclusion criterion was Model for End-stage Liver Disease (MELD) score at transplant 15 or greater and no prior transplant. A laboratory of MELD ≥15 was chosen as this is the threshold MELD at which the benefit of liver transplant becomes apparent, and the focus of the study was on patients with decompensated liver disease. The Liver Donor Risk Index (LDRI) was computed using previously reported methods.

Statistical Methods

The effect of antiviral era on the proportions of persons transplanted within each diagnosis category (HCV, HBV, NASH, ALD, other) was evaluated using the \( \chi^2 \) test. Pairwise \( z \)-tests were used to evaluate differences between diagnosis-specific between-era proportions. Among persons transplanted for HCV, HBV, NASH and ALD, survival analysis methods tested the main and interaction effects of diagnosis and era on graft survival (event = retransplantation or death) over the first 3 years after liver transplantation. Observations were censored: (1) on the date of retransplantation or death if these events occurred within 3 years or (2) at the 3-year follow-up date if they occurred >3 years after liver transplantation. This approach equilibrated follow-up time between the antiviral therapy eras, for which total follow-up time necessarily differed.

RESULTS

A total of 53,788 patients with decompensated end-stage liver disease as defined by a laboratory MELD score of 15 or greater were transplanted during the study period of whom 61.9% were transplanted for HCV, HBV, NASH, or ALD. The primary diagnosis for deceased donor liver transplantation changed significantly over the 15-year period \( P < 0.001 \) (Table 1). The percentage of recipients transplanted for compensated HCV progressively decreased from 35.3% in the IFN era to 23.6% in the DAA era (all column-wise, \( P < 0.05 \)). This corresponded with a tripling of transplantation for recipients with compensated NASH, from 5.8% in the IFN era to 16.5% in the DAA era (all column-wise, \( P < 0.05 \)). There was also an era-related increase in the transplantation of patients with compensated ALD (15.6% IFN era versus 24.0% DAA era) (all column-wise, \( P < 0.05 \)), whereas the percentage of patients transplanted for compensated HBV remained relatively stable (2.9% IFN era versus 2.1% DAA era). Era-related changes in recipient characteristics extended beyond primary diagnosis at transplantation. Recipient age at transplant and diabetes progressively increased over time (all pairwise, \( P < 0.05 \)); Laboratory MELD score at transplant increased between the IFN and PI eras and remained stable thereafter. The percentage of patients receiving...
life support at time of transplant doubled from 6.2% in the IFN era to 12.5% in the DAA era. Correspondingly, over a quarter of patients (25.8%) required pretransplant dialysis in the DAA era compared with 14.1% in the IFN era.

Many characteristics of liver transplant recipients varied significantly by diagnosis (column-wise, \( P < 0.05 \)) (Table 2). The HCV recipients had the lowest laboratory MELD at transplant (25.5) and highest percentage of recipients with an HCC exception (12.2%). The HCV recipients also received the highest-quality organs as evidenced by the lowest LDRI score of 1.47. Conversely, NASH recipients were older at transplant and more often had comorbidities of diabetes (5.5%), hypertension (38.8%), and CAD (5.4%). The NASH recipients were larger (body mass index average, 32.3) and more likely to have had a prior abdominal operation (50.2%) or portal vein thrombosis (7.6%). Interestingly, nearly half of all NASH recipients were female but females comprised only about one quarter of the other recipients.

Unadjusted Kaplan Meier estimated probability of graft survival was compared between each era, stratified by (separately within) each diagnosis (Figure 1). A significant era-related improvement in graft survival was identified in HCV recipients (all log-rank, \( P \leq 0.001 \)), with 1- and 3-year point estimates increasing from 83.3% and 71.9% in the IFN era to 90.9% and 79% in the DAA era. Era-related improvement in graft survival was also noted for ALD recipients when comparing the PI or DAA era to IFN era. No differences were seen in graft survival between the IFN and DAA eras for

| TABLE 1. Cohort characteristics |
|--------------------------------|
| **Temporal trends in deceased donor liver transplantation by primary diagnosis** |
| **IFN (2003-2010)** | **PI (2011-2013)** | **DAA (2014-2017)** | **All eras** |
| Primary diagnosis | | | Between era, \( P \) |
| HCV | 9973 (35.3) | 3557 (31.9) | 3394 (23.6) | 16924 (31.5) | <0.001 |
| HBV | 824 (2.9) | 232 (2.1) | 302 (2.1) | 1358 (2.5) |
| NASH | 1637 (5.8) | 1251 (11.2) | 2370 (16.5) | 5258 (9.8) |
| ALD | 4423 (15.6) | 1898 (17.0) | 3442 (24.0) | 9763 (18.2) |
| Other | 11414 (40.4) | 4212 (37.8) | 4859 (33.8) | 20485 (38.1) |
| Survival analysis cohort (primary diagnosis HCV, HBV, NASH, or ALD) |
| n | 16857 | 6938 | 9508 | 33303 |
| Age, y | 53.8 (7.9) | 55.9 (8.1) | 56.4 (8.9) | 55.0 (8.3) | <0.001 |
| Male, % | 74.0 | 70.6 | 69.3 | 71.9 | <0.001 |
| Life support, % | 6.2 | 9.6 | 12.5 | 8.7 | <0.001 |
| LDRI score | 1.5 (0.4) | 1.5 (0.4) | 1.5 (0.4) | 1.5 (0.4) | 0.36 |
| MELD at transplant | 25.1 (7.7) | 27.3 (8.1) | 27.6 (8.4) | 26.3 (8.1) | <0.001 |
| HCC exception, % | 7.5 | 9.1 | 8.6 | 8.1 | <0.001 |
| Diabetes, % | 23.4 | 26.4 | 30.1 | 26.0 | <0.001 |
| Dialysis, % | 14.1 | 21.9 | 25.8 | 19.0 | <0.001 |

Unless noted, table entries are \( n \) (column %) or mean (SD).

Column-wise z tests of proportions or post hoc Bonferroni-adjusted comparisons: \( P < 0.05 \), a versus a and b versus b.

ALD, alcoholic liver disease; DAA, direct-acting antiviral; HBV, hepatitis B virus; HCC, hepatocellular carcinoma; HCV, hepatitis C virus; IFN, interferon; LDRI, Liver Donor Risk Index; MELD, Model for End-stage Liver Disease; NASH, nonalcoholic steatohepatitis; PI, protease inhibitors. PVT, portal vein thrombosis.

| TABLE 2. Survival analysis cohort characteristics by diagnosis |
|--------------------------------|
| **HCV (n = 16924)** | **HBV (n = 1358)** | **NASH (n = 5258)** | **ALD (n = 9763)** |
| Age, y | 54.7 (7.1) | 51.5 (10.7) | 58.9 (8.3) | 53.7 (9.2) | <0.001 |
| Male sex | 12521 (74.0) | 1035 (76.2) | 2856 (54.3) | 7546 (77.3) | <0.001 |
| Laboratory MELD at transplant | 25.5 (8.1) | 29.1 (8.8) | 26.0 (7.8) | 27.3 (8.9) | <0.001 |
| HCC exception | 2064 (12.2) | 120 (8.8) | 215 (4.1) | 312 (3.2) | <0.001 |
| Recipient life support | 1224 (7.2) | 222 (16.3) | 444 (8.4) | 1013 (10.4) | <0.001 |
| Recipient diabetes | 3532 (21.2) | 296 (22.3) | 2861 (55.0) | 1841 (19.1) | <0.001 |
| Recipient dialysis | 2776 (16.4) | 232 (17.1) | 1113 (21.2) | 2221 (22.7) | <0.001 |
| Recipient hypertension | 3039 (23.4) | 188 (13.9) | 1211 (23.8) | 1607 (25.9) | <0.001 |
| Recipient coronary artery disease* | 260 (2.5) | 22 (3.0) | 126 (5.4) | 145 (2.8) | <0.001 |
| Prior abdominal surgery | 5500 (33.6) | 313 (24.5) | 2586 (50.2) | 2660 (28.0) | <0.001 |
| Portal vein thrombosis | 650 (4.0) | 46 (3.6) | 393 (7.6) | 381 (4.0) | <0.001 |
| Body mass index (kg/m²) | 28.6 (5.6) | 27.5 (5.6) | 32.3 (6.0) | 28.2 (5.6) | <0.001 |
| LDRI | 1.47 (0.38) | 1.51 (0.41) | 1.53 (0.42) | 1.52 (0.41) | <0.001 |

Table entries are mean (SD) or \( n \) (%). Post hoc pairwise comparisons (\( P \leq 0.001 \)) and z tests of column percentages (\( P < 0.05 \)) a versus a, b versus b, and c versus c.

*Data were populated for fewer than 60% of cases; all other variables were populated for at least 97% of cases.

ALD, alcoholic liver disease; HBV, hepatitis B virus; HCC, hepatocellular carcinoma; HCV, hepatitis C virus; IFN, interferon; LDRI, Liver Donor Risk Index; MELD, Model for End-stage Liver Disease; NASH, nonalcoholic steatohepatitis.
either HBV or NASH recipients. Similarly, unadjusted Kaplan-Meier estimated probability of graft survival was compared between each of the 4 diagnoses, stratified by the 3 eras (Figure 2). In the IFN era, HCV recipients had the lowest 1- and 3-year graft survivals of any diagnosis (log-rank, \( P < 0.001 \) versus other diagnoses). However, by the DAA era, NASH recipients had lower graft survival (87.5% 1 year, 77.9% 3 years) compared with both HCV (90.0% 1 year, 79% 3 years) and ALD (90.7% 1 year, 78.9% 3 years) recipients (log-rank \( P \leq 0.01 \)).

Multivariable modeling identified increasing recipient age, presence of recipient diabetes, increasing laboratory MELD score, need for life support, and HCC exception as independent predictors of graft failure over the first 3 years (Table 3). The donor characteristics of increasing LDRI score and diabetes were also associated with an increased risk of graft failure over the first 3 years. Although there were indications of both era- and diagnosis-related differences in graft survival, a statistically significant era by diagnosis interaction effect was identified (\( P < 0.001 \)) precluding interpretation of these findings in isolation. Specifically, the likelihood of graft loss for a given diagnosis varied between eras as illustrated by the significantly decreased risk of graft loss for HCV recipients in the DAA era when compared with ALD recipients in the IFN era (HR 0.705; 95% CI 0.593-0.838; DAA HCV interaction contrast, Table 3). Interestingly, although not reaching statistical significance, there was a trend toward an increased risk of graft loss for NASH patients in both the PI and DAA eras when compared with the IFN era ALD reference group.

Additional era-specific multivariable modeling was performed to further evaluate the effects of diagnosis within each era in the setting of the previously-described era by diagnosis interaction effect (Table 4). These effects were excerpted from...
3 separate era-specific models that adjusted for the recipient and donor characteristics included in Table 3. After adjusting for the recipient and donor characteristics, the era-specific models demonstrate that during the IFN era HBV, NASH, and ALD had a significantly decreased risk of graft failure compared with HCV. However, by the PI era only ALD had a significantly decreased risk of graft failure compared with HCV. In the most recent DAA era HCV primary diagnosis conferred no difference in risk of graft failure as compared with HBV, NASH, or ALD.

**DISCUSSION**

The development of effective treatment for HCV infection and the progressive obesity epidemic have dramatically altered the landscape of liver transplantation in the United States. Our
study examined the indication for liver transplantation over 3 eras spanning 2003 to the current time and confirms that the incidence of liver transplantation for complications of HCV-related end-stage liver disease is progressively declining while the incidence of transplantation for NASH is on the rise. Our study additionally aimed to evaluate outcomes over 3 years and found improved outcomes for patients with HCV over the eras, after accounting for era-related effects.

The effect of HCV antiviral treatment has also been examined in terms of the liver transplant waitlist. Flemming et al analyzed listing rates using the SRTR database from 2003 to 2015 for decompensated cirrhosis and HCV, HBV, or NASH in the DAA era compared with the IFN and PI eras and noted that waitlisting for HCV with decompensated cirrhosis declined by >30% in the DAA era. Similarly, European studies have reported delisting rates in patients with HCV since the advent of the current antiviral therapies. Perricone et al examined patients with HCV on the waitlist for liver transplant and found that 30% were delisted, with a <10% incidence of liver-related complications in the subsequent 2 years after delisting. Kwong et al examined the OPTN database to examine changes in waitlist mortality in listed patients with HCV. The study reported a decrease in waitlist mortality in the DAA era for patients with HCV, and the authors presumed this change to be at least in part due to the advent of DAA therapy. Correspondingly, Bowring et al examined the SRTR data and found an increase in the use of HCV livers (6.9% in 2010 to 16.9% in 2015). In summary, effective HCV antiviral therapy has led to a decrease in HCV liver transplants, a decrease in HCV patients being listed for liver transplant, an increase in HCV patients being removed from the liver transplant waitlist and an increase in the use of HCV donor livers.

The question that inevitably follows is whether HCV antiviral treatment has resulted in improved longer-term outcomes after liver transplantation. Cholankeril et al evaluated the main effects of HCV diagnosis in the UNOS database and comparably-defined transplant antiviral eras on 1-year survival after liver transplantation. They noted reduced short-term 1-year mortality and graft failure in liver transplant recipients with HCV in the DAA era compared with the pre-DAA era. Our study aimed to evaluate outcomes over 3 years and found improved outcomes for patients with HCV over the eras. Axelrod et al examined SRTR data from 2007 to 2016 and merged these data with national pharmaceutical claims to specifically examine the impact of posttransplant antiviral HCV treatment. The authors found that posttransplant antiviral therapy improved outcomes for HCV patients up to 3 years after liver transplant. As the use of HCV antiviral therapies becomes more widespread (and hopefully more accessible) the long-term effectiveness of these therapies and their effect on liver transplant outcomes will be further elucidated. Further investigations into the timing of treatment surrounding liver transplant and the treatment of HCV patients with liver disease but without cirrhosis are needed.

Concurrent with the decline of HCV liver transplants is the rise of NASH liver disease. Pandemic obesity is thought to be driving the increase of nonalcoholic fatty liver disease and NASH cirrhosis in the United States and globally. The burden of NASH on liver transplantation is anticipated to rise sharply, with decompensated cirrhosis and HCC attributed to NASH each projected to increase by well over 100% in the United States by 2030. The NASH liver disease is often viewed as the liver manifestation of metabolic syndrome, and there is interplay with obesity, diabetes, hyperlipidemia and hypertension. These patients have more comorbidities compared to patients without NASH as described in our study. Recipient age, diabetes, coronary artery disease, and portal vein thrombosis are all known risk factors associated with inferior patient and graft survival postliver transplant. All of these risk factors are more prevalent in NASH recipients and may account for the lower 1- and 3-year graft survivals in the modern DAA era when compared with HCV, HBV, and ALD recipients. Although recipients with NASH have “equivalent” survival when compared with non-NASH recipients after adjustment for age and diabetes, these represent artificial corrections as over 50% of patients with NASH have diabetes, and the pathophysiology of NASH is such that it takes longer to develop into cirrhosis. Additionally, the same risk factors leading to the development of NASH cirrhosis, metabolic syndrome and obesity, remain largely untreated postoperatively with as many as 90% of patients transplanted

**TABLE 4.**

| Each model, $P < 0.001$ | Estimate | $P$ | Hazard ratio | $95\%$ CI | Lower bound | Upper bound |
|------------------------|----------|-----|--------------|-----------|-------------|-------------|
| IFN era model (n = 14775) | 0.002 | 0.000 | 0.621 | 0.489 | 0.789 |
| HBV (reference: HCV) | −0.476 | 0.000 | 0.621 | 0.489 | 0.789 |
| NASH (reference: HCV) | −0.245 | 0.02 | 0.782 | 0.640 | 0.956 |
| ALD (reference: HCV) | −0.226 | 0.01 | 0.797 | 0.670 | 0.949 |
| PI era model (n = 6668) | 0.06 | 0.000 | 1.218 | 0.541 | 1.266 |
| HBV (reference: HCV) | −0.189 | 0.38 | 0.828 | 0.529 | 1.012 |
| NASH (reference: HCV) | −0.313 | 0.06 | 0.731 | 0.479 | 0.907 |
| ALD (reference: HCV) | −0.417 | 0.01 | 0.659 | 0.313 | 0.731 |
| DAA era model (n = 9224) | 0.17 | 0.000 | 1.218 | 0.606 | 1.657 |
| HBV (reference: HCV) | 0.02 | 0.99 | 1.002 | 0.968 | 1.958 |
| NASH (reference: HCV) | 0.320 | 0.08 | 1.377 | 0.865 | 1.715 |
| ALD (reference: HCV) | 0.197 | 0.26 | 1.218 | 0.85 | 1.715 |

These effects were excerpted from 3 separate era-specific models that adjusted for the recipient and donor characteristics included in Table 3.

ALD, alcoholic liver disease; DAA, direct-acting antiviral; HBV, hepatitis B virus; HCV, hepatitis C virus; IFN, interferon; NASH, nonalcoholic steatohepatitis; PI, protease inhibitors.
for NASH redeveloping NAFLD and up to 40% developing NASH in some cohorts. The same factors that cause NASH predispose these patients to progressive cardiovascular disease postoperatively despite the highly restrictive selection process for cardiovascular disease and obesity in candidates needing liver transplantation, resulting in high decline rates for patients with NASH cirrhosis. In fact, cardiovascular disease remains one of the leading causes of mortality after liver transplant, and this is likely to rise as the percentage of liver transplants performed for NASH cirrhosis increases. Although vast resources and time have been dedicated to the treatment of HCV with great success, there is currently no pharmacologic agent approved for treatment of NASH or NASH recurrence, which will soon overwhelm the liver transplant system.

This study is limited by the unavoidable shorter duration of follow up for the most recent DAA period. For this reason, we limited the follow-up to the first 3 years for all antiviral era cohorts. Although short-term outcomes may now be similar among groups the long term outcomes may diverge beyond 3 years. The limitations related to the use of a large national database include the lack of granularity of the data, namely the use of antiviral therapy and the timing of administration of antiviral therapy. The data can thus demonstrate era-related effects, but are unable to definitively confirm outcomes being related to the use of specific antiviral therapies.

In the current era of DAA HCV therapy, liver transplantation for HCV is on the decline with improved outcomes. This is being counterbalanced by a rapid increase in transplantation for NASH, which now has the lowest overall 1- and 3-year graft survival of the diagnoses studied. Treatment and prevention of nonalcoholic fatty liver disease prior to the development of end-stage liver disease and prevention of recurrence posttransplant must be aggressively pursued.

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