Title
Curing Hepatitis C in Liver Transplant Recipients Is Associated with Changes in Immunosuppressant Use.

Permalink
https://escholarship.org/uc/item/96r5g42v

Journal
Journal of clinical and translational hepatology, 4(1)

ISSN
2225-0719

Authors
Saab, Sammy
Rheem, Justin
Jimenez, Melissa
et al.

Publication Date
2016-03-15

DOI
10.14218/jcth.2016.00001

Peer reviewed
Curing Hepatitis C in Liver Transplant Recipients Is Associated with Changes in Immunosuppressant Use

Sammy Saab*1,2, Justin Rheem3, Melissa Jimenez1, Sherona Bau2, Gina Choi1,2, Francisco Durazo1,2, Mohammed El Kabany1,2, Steven Han1,2, Alexander Farid1, Naadir Jamal1, Jonathan Grotts1, David Elashoff1,4 and Ronald W. Busuttil2

1Departments of Medicine at the University of California at Los Angeles, Los Angeles, California, USA; 2Department of Surgery at the University of California at Los Angeles, Los Angeles, California, USA; 3Department of Medicine at Harbor-University of California at Los Angeles Medical Center, Torrance, California, USA; 4Department of Biostatistics at the University of California at Los Angeles, Los Angeles, California, USA

Abstract

**Background and Aims:** All-oral interferon-free antivirals are highly effective in treating recurrent hepatitis C (HCV) infection in liver transplant (LT) recipients. The aim of the study was to assess immunosuppression needs after achieving a sustained viral response (SVR). **Methods:** We compared immunosuppression needs before and after achieving an SVR in adult LT treated recipients for recurrent HCV infection with all-oral direct acting agents. **Results:** We identified 52 liver LT treated recipients who achieved an SVR. The median (25th and 75th percentile interquartile range [IQR]) age was 62 years (57.75, 65). Most recipients received tacrolimus (TAC) for their immunosuppressant regimen. After achieving SVR, there was a statistically significant decrease in daily dose of TAC adjusted per weight (p > 0.05). However, there was a statistically significant decrease in daily dose of TAC adjusted per weight, serum levels of TAC, and the product of glomerular filtration rate and TAC. No statistically significant differences in cyclosporine unadjusted/adjusted per weight daily dose or serum levels were noted. **Conclusions:** Immunosuppression needs were increased for those patients treated with TAC but not cyclosporine. LT recipients prescribed TAC require close monitoring after treatment completion to avoid potential risk of acute rejection.

© 2016 The Second Affiliated Hospital of Chongqing Medical University. Published by XIA & HE Publishing Inc. All rights reserved.

**Keywords:** Liver transplantation; Immunosuppressant; Hepatitis C; Direct acting agents.

**Abbreviations:** ALT, alanine aminotransferase; AP, alkaline phosphatase; AST, aspartate Transaminase; BMI, body mass index; CKD-EPI, Chronic Kidney Disease Epidemiology Collaboration Formula; CSA, cyclosporine; GFR, glomerular filtration rate; HCV, Hepatitis C; HIV, human immunodeficiency virus; IQR, interquartile range; LT, liver transplant; MELD, Model for End-Stage Liver Disease; mTOR, mammalian target of rapamycin; SVR, sustained viral response; TAC, tacrolimus; TDD, total daily dose; UCLA, University of California, Los Angeles Medical Center.

Received: 5 January 2016; Revised: 19 February 2016; Accepted: 24 February 2016

*DOI: 10.14218/JCTH.2016.00001.

*Correspondence to: Sammy Saab, Pfleger Liver Institute, UCLA Medical Center, 200 Medical Plaza, Suite 214, Los Angeles, CA 90095, USA. Tel: +1-310-206-6705, Fax: +1-310-206-4197, E-mail: SSaab@mednet.ucla.edu

Original Research

Introduction

Liver transplantation (LT) is the definitive treatment for patients with advanced liver disease and hepatocellular carcinoma.1–3 Initial attempts at LT in the 1960s were limited by graft failure from cellular rejection.4–6 Advances in immunosuppressive therapy have led to substantial improvement in patient and graft survival over the past several decades.7 However, the use of immunosuppressive therapy is also associated with long term complications, such as hypertension, diabetes, and renal insufficiency.8–13 The goal of immunosuppressive therapy is to maximize protection from rejection while minimizing the adverse effects by using the lowest dose possible.14 The most common indication for LT in the United States is hepatitis C.15 However, recurrent HCV infection is universal, and its progression to cirrhosis can lead to graft loss in a significant number of LT recipients.16–18 Successful treatment of recurrent hepatitis C virus (HCV) infection in LT recipients can reduce the risk of subsequent HCV-related complications and graft loss.14,17,19,20 The use of interferon to treat recurrent HCV was associated with concerns for graft loss because of the perceived enhanced immune modulating effects of interferon.21–23 Furthermore, graft failures were found to occur after viral suppression was achieved.21 The introduction of all-oral interferon-free antiviral therapy has revolutionized the treatment of HCV infection. These treatments are safe and highly effective in LT recipients with recurrent HCV infection, thus broadening the group of patients eligible for treatment.24–26 A direct consequence of avoiding interferon in the treatment of recurrent HCV is the increased number of recipients that would be eligible for the interferon-free therapy. However, a possible unexpected consequence of achieving viral suppression may be improved metabolism of immunosuppressant medications, aside from direct drug-drug interactions. For instance, the results of two recent studies of direct acting agents highlight changes in immunosuppressant dosing after antiviral therapy was completed.26,27 Calcineurin inhibitors are all metabolized by the liver (cytochrome P-450 3A4 system) and increased performance of hepatic metabolism function highly influences the bioavailability of these medications.28 With improved liver function after HCV treatment in LT recipients, patients’ needs for immunosuppression may change.
The hypothesis of this study is that LT recipients who achieve a sustained viral response (SVR) will require greater amounts of immunosuppression. Understanding this phenomenon will help to tailor strategies that will protect LT recipients from potential graft rejection after they are cured of HCV.

**Methods**

We performed a retrospective cohort study of all adult (age >18 years) LT recipients who achieved SVR after being treated with all-oral HCV antiviral therapy for recurrent HCV infection at the University of California, Los Angeles Medical Center (UCLA) between October 2011 and October 2014. A SVR was defined as an undetectable HCV RNA 12 weeks after treatment completion. Exclusion criteria included active infection, biliary obstruction, and allograft rejection on biopsy within 6 months of starting antiviral therapy.

This study was approved by the UCLA Institutional Review Board. Patient consent was not required. Data were obtained by medical chart review and retrieved by the UCLA Liver Transplant database. Demographic data (age, gender, and race/ethnicity), history of LT, history of previous HCV therapy, nonliver related medical history (active cardiopulmonary disease, hemodialysis, stroke, nonliver malignancy, and diabetes), co-existent liver disease (nonalcoholic fatty liver disease, hepatitis B, and autoimmune hepatitis), and stage of fibrosis, categorized as Mild (stage 0–2) and Moderate to Severe (stage 3–4), were recorded. Fibrosis was defined by liver biopsy, laboratory testing, and radiological imaging. Hematologic, biochemical data, immunosuppressant levels, and HCV RNA levels were collected prior to initiation of antiviral therapy and after achieving a SVR.

**Immunosuppressive needs**

We ascertained immunosuppressant needs through a variety of methods: 1) total daily dose of the immunosuppressant, 2) the dose of immunosuppressant per the recipient body weight (kg), 3) product of immunosuppressant and glomerular filtration rate (GFR) (Chronic Kidney Disease Epidemiology Collaboration Formula, CKD-EPI, mL/min/1.73 m²). The last method was chosen since it is our custom to tailor immunosuppressant dose to GFR to avoid renal toxicity. We compared laboratory values and computations for 3 months before starting HCV therapy and 3 months after therapy was completed.

**Statistics**

Data are presented as medians with 25th and 75th percentile interquartile range (IQR) for continuous variables and number within group with percent of group for discrete variables. The within patient differences in labs and medication regimens were compared before and after treatment using the two-sided Wilcoxon Rank Sum test. A multivariable linear regression model with percent change in immunosuppressant serum level as the dependent variable and percent change in alanine aminotransferase (ALT) and weight-adjusted, total daily dose as the independent variables was used to assess changes in serum level with respect to medication and liver function. \( p < 0.05 \) was considered statistically significant. The R Statistical Computing Environment was used for data analysis (R Core Team 2015).

**Results**

The demographics of the study patient cohort are presented in Table 1. Twenty-five (48.1%) recipients had at least one nonhepatic comorbidity. Few patients had concomitant fatty liver, and no recipients were co-infected with hepatitis B or human immunodeficiency virus (HIV). Twenty percent of recipients had a diagnosis of hepatocellular carcinoma prior to LT. Most patients were infected with genotype 1 (78.8%). The rest of patients were infected with genotypes 2, 3, and 4 (5.8%, 11.5% and 3.8%, respectively). Pretreatment lab results were available a median (IQR) of 5 (3, 7.25) months. The median (IQR) time from SVR12 to collection of three laboratory values was 5 months (4, 7.25).

Tacrolimus (TAC)-based immunosuppression was used in 75% of LT recipients. The median (IQR) total daily dose, weight-adjusted total daily dose, and serum TAC values were 2 (1.5, 4) mg, 0.03 (0.02, 0.05) mg/kg, and 8.1 (6.57, 9.1) ng/mL, respectively. Of the recipients treated with cyclosporine, the median (IQR) total daily dose, weight-adjusted total daily dose, and serum cyclosporine was 150 (133.33, 200) mg, 1.93 (1.37, 2.81) mg/kg, and 112 (79, 128.67) ng/mL, respectively. Fifteen percent of recipients treated with TAC had moderate to severe fibrosis, and 39% of recipients treated with cyclosporine had moderate to severe fibrosis. Most recipients were taking mycophenolate before and after achieving a SVR.

The median (IQR) aspartate aminotransferase (AST) and ALT before achieving a SVR were 58.83 (40.66, 80.92) and 62.66 (43.67, 88.42), respectively. There was a statistically significant improvement in all assessed liver associated tests in LT recipients who achieved SVR12 \( (p < 0.05) \) (Table 2). There was also a statistically significant improvement in platelet count \( (p = 0.005) \). Body mass index (BMI) increased with SVR12. The median (IQR) BMI was 26.8 (23.69, 29.78) before treatment initiation and 27.58 (23.94, 31.0) after SVR \( (p = 0.012) \). There was no change in renal function as measured by GFR after achieving SVR \( (p = 0.099) \).

Changes in immunosuppression need were assessed in multiple ways (Tables 3 and 4). Most recipients treated with TAC had a decrease in their weight adjusted total daily dose of TAC and TAC serum levels (Table 3). There was no statistically significant change in the unadjusted total dose of TAC \( (p = 0.174) \), but there was an observed significant decrease in weight adjusted total dose of TAC \( (p = 0.001) \) and serum levels of TAC after achieving SVR \( (p\text{-value} < 0.001) \) (Table 4, Figure 1a). Moreover, the product of TAC serum level and GFR was significantly lower after SVR \( (p < 0.001) \). In contrast, there was no significant changes in unadjusted or adjusted total dose of cyclosporine, serum levels of cyclosporine, or the product of cyclosporine serum level and GFR \( (p > 0.05) \) (Table 4, Figure 1b). There was an observed interaction between serum change of immunosuppression levels and total daily dose immunosuppression with the percent change of median ALT values (Figure 2). In a multivariate analysis, there was an observed relationship between percent change in serum levels and percent change in weight-adjusted, total daily dose \( (p = 0.027) \), but not in percent change in median ALT values \( (p = 0.226) \). When restricting the population to only patients taking TAC, the relationship was significant between change in serum levels and weight-adjusted total daily dose \( (p = 0.014) \) and change in serum levels and change in median ALT values \( (p = 0.022) \).
The results of our study demonstrate that LT recipients’ immunosuppression needs may increase after achieving a SVR. The goal of immunosuppression in LT is to minimize clinically significant side effects and to prevent rejection.\textsuperscript{28–32} Immunosuppression regimens are tailored based on multiple factors, including renal and hepatic function, and may require frequent adjustments by providers to tailor them to the individual patient. Therefore, the clinical implication of our study is that LT recipient’s status post-recurrent HCV treatment may be at increased risk of rejection.

Our results are consistent with prior observations made by others. Kwo \textit{et al.} found that eight out of 34 LT recipients required adjustments of their immunosuppressant shortly after completing their antiviral therapy.\textsuperscript{26} Although there is a significant drug-drug interaction between the 3D regimen, TAC, and cyclosporine, and the immunosuppressant trough levels were mostly stabilized several weeks after initiation of the 3D regimen, these observations were made after completing therapy.\textsuperscript{33} Likewise, Chartlon \textit{et al.} described 25 of 229 LT recipients who required dose adjustments after completing a course of antiviral therapy, but the direction of the change was not characterized.\textsuperscript{27} We speculate that improvement in liver function with interferon-free therapy may explain the increased need for immunosuppression. In patients with recurrent HCV post-transplantation, treatment with interferon-free therapy resulted in early post-treatment liver function improvement, as evidenced by decreases in bilirubin and Model for End-Stage Liver Disease (MELD) scores and increases in albumin.\textsuperscript{34,35} Improved metabolic function of the liver may explain our results. Calcineurin inhibitors, mammalian target of rapamycin (mTOR) inhibitors, and antimetabolites are metabolized by the liver (cytochrome P-450 3A4 system), and performance of hepatic metabolism function highly influences the bioavailability of these agents.\textsuperscript{28} An important finding of our study is the differential effect of curing HCV on

| Variable                                      | All   | Tacrolimus | Cyclosporine |
|------------------------------------------------|-------|------------|--------------|
| Number of patients                            | 52    | 39         | 13           |
| Age, median (interquartile range)              | 62 (58,65) | 62 (58,65) | 60 (55,63)   |
| Gender                                         |       |            |              |
| Male (%)                                       | 42 (80.77) | 31 (79.49) | 11 (84.62)   |
| Hispanic (%)                                   | 18 (34.62) | 12 (30.77) | 6 (46.15)    |
| Non-Hispanic White (%)                         | 16 (30.77) | 11 (28.21) | 5 (38.46)    |
| Other (%)                                      | 18 (34.62) | 16 (41.03) | 2 (15.38)    |
| Body Mass Index (kg/m\(^2\), median (interquartile range) | 26.8 (23.69, 29.78) | 25.73 (23.23, 29.55) | 28.59 (25.58, 30.94) |
| Months since liver transplant, median (interquartile range) | 32.9 (13.78, 74.42) | 29.18 (13.09, 60.82) | 59.24 (39.63, 179.60) |
| Mycophenolate                                  |       |            |              |
| Before Treatment                               | 36 (69.23) | 29 (74.36) | 7 (53.85)    |
| After Treatment                                | 35 (67.31) | 26 (66.67) | 9 (69.23)    |
| Sirolimus                                      |       |            |              |
| Before Treatment                               | 1 (1.92) | 0 (0)      | 1 (7.69)     |
| After Treatment                                | 2 (3.85) | 1 (2.56)   | 1 (7.69)     |
| Hepatitis C Treatment Experienced (%)          | 23 (44.2) | 14 (35.9)  | 9 (69.2)     |
| Antiviral Therapy                              |       |            |              |
| Sofosbuvir/Ribavirin (%)                       | 31 (59.6) | 23 (59)    | 8 (61.5)     |
| Sofosbuvir/Simeprevir (%)                      | 21 (40.4) | 16 (41)    | 5 (38.5)     |
| Non-hepatic comorbidities                      |       |            |              |
| Cardiac disease (%)                            | 5 (9.6) | 5 (12.8)   | 0 (0)        |
| Hemodialysis (%)                               | 3 (5.77) | 2 (5.13)   | 1 (7.69)     |
| Cerebral vascular stroke (%)                   | 1 (1.92) | 1 (2.56)   | 0 (0)        |
| Non-liver malignancy (%)                       | 5 (9.6) | 4 (10.3)   | 1 (7.69)     |
| Diabetes (%)                                   | 17 (32.69) | 13 (33.33) | 4 (30.77)    |
| Lymphoma (Non-PTLD) (%)                        | 1 (1.92) | 0 (0)      | 1 (7.7)      |

Discussion

The results of our study demonstrate that LT recipients’ immunosuppression needs may increase after achieving a SVR. The goal of immunosuppression in LT is to minimize clinically significant side effects and to prevent rejection.\textsuperscript{28–32} Immunosuppression regimens are tailored based on multiple factors, including renal and hepatic function, and may require frequent adjustments by providers to tailor them to the individual patient. Therefore, the clinical implication of our study is that LT recipient’s status post-recurrent HCV treatment may be at increased risk of rejection.

Our results are consistent with prior observations made by others. Kwo \textit{et al.} found that eight out of 34 LT recipients required adjustments of their immunosuppressant shortly after completing their antiviral therapy.\textsuperscript{26} Although there is a significant drug-drug interaction between the 3D regimen, TAC, and cyclosporine, and the immunosuppressant trough levels were mostly stabilized several weeks after initiation of the 3D regimen, these observations were made after completing therapy.\textsuperscript{33} Likewise, Chartlon \textit{et al.} described 25 of 229 LT recipients who required dose adjustments after completing a course of antiviral therapy, but the direction of the change was not characterized.\textsuperscript{27} We speculate that improvement in liver function with interferon-free therapy may explain the increased need for immunosuppression. In patients with recurrent HCV post-transplantation, treatment with interferon-free therapy resulted in early post-treatment liver function improvement, as evidenced by decreases in bilirubin and Model for End-Stage Liver Disease (MELD) scores and increases in albumin.\textsuperscript{34,35} Improved metabolic function of the liver may explain our results. Calcineurin inhibitors, mammalian target of rapamycin (mTOR) inhibitors, and antimetabolites are metabolized by the liver (cytochrome P-450 3A4 system), and performance of hepatic metabolism function highly influences the bioavailability of these agents.\textsuperscript{28} An important finding of our study is the differential effect of curing HCV on
TAC and cyclosporine levels. One possible explanation could be that the serum concentration of cyclosporine is 15-fold higher than that of TAC (112 ng/ml vs 8.1 ng/ml, respectively) and that cyclosporine, but not TAC, has a pharmacokinetics inhibitory effect on hepatic CYP metabolic enzymes.36 In addition, restoration of host immune response by interferon-free therapy may also explain this phenomenon.

In chronic hepatitis C infection, HCV renders the host’s virus-specific CD8+ T cells dysfunctional and transient by several mechanisms, such as viral escape and T-cell exhaustion.37,38 Previously, interferon-based therapies failed to recover HCV-specific CD8+ T cell functions.39,40 However, in a recent study with IFN-free therapies (faldaprevir and deleobuvir), proliferative HCV-specific CD8+ T cells were rapidly restored in the majority of patients (n/N = 19/22, 86.4%).3 In contrast, HCV-specific CD8+ T cells did not increase in patients with treatment failure. This finding suggests that adaptive immunity after successful HCV treatment is restored and may require immunosuppression adjustment to prevent graft rejection.

In addition, successful HCV therapy can lead to removal of the immunosuppressive effect from HCV itself. In a few LT recipients, immunosuppressive medication may not be required after sometime due to a phenomenon called “transplant tolerance.”29,41 Its mechanism is not completely understood, but the current hypothesis is that HCV infection induces immunoregulation and facilitates “transplant tolerance,” leading to successful discontinuation of immunosuppression.41 In contrast, a successful eradication of the immunosuppressive effect of HCV may lead to a heightened

### Table 2. Changes in clinical and laboratory parameters before and after achieving a sustained viral response. Data are presented as a median with interquartile range in parenthesis for continuous variables

| Parameter          | Before                  | After                   | Within patient difference | P-value |
|--------------------|-------------------------|-------------------------|---------------------------|---------|
| **Clinical**       |                         |                         |                           |         |
| BMI (kg/m²)        | 26.8 (23.685, 29.777)   | 27.58 (23.94, 31.003)   | 0.46 (−0.153, 1.588)      | 0.012   |
| **Laboratory**     |                         |                         |                           |         |
| Hemoglobin (g/dL)  | 12.84 (11.047, 14.803)  | 13.435 (11.793, 14.72)  | 0.48 (−0.592, 1.462)      | 0.111   |
| Platelet Count (×10^12/μL) | 121.335 (77.875, 147.92) | 136.835 (104.082, 159.67) | 11.835 (−5.165, 34.915) | 0.005   |
| SCL (mg/dL)        | 1.13 (0.908, 1.355)     | 1.115 (0.918, 1.36)     | 0.03 (−0.1, 0.15)         | 0.412   |
| Albumin (g/dL)     | 4.07 (3.73, 4.3)        | 4.37 (4.17, 4.508)      | 0.215 (0.07, 0.515)       | < 0.001 |
| AST (U/L)          | 58.83 (40.665, 80.918)  | 24.665 (18.67, 30.415)  | −27.335 (−54.165, −14.082)| < 0.001 |
| ALT (U/L)          | 62.665 (43.668, 88.418) | 21.665 (16, 33.415)     | −34.835 (−65.08, −15.668) | < 0.001 |
| Bilirubin, total (mg/dL) | 0.95 (0.73, 1.178)     | 0.7 (0.53, 0.948)       | −0.215 (−0.4, −0.06)      | < 0.001 |
| AP (U/L)           | 96.5 (73.75, 134.502)   | 84.665 (63.75, 117.418) | −14.835 (−28.338, −0.077) | 0.001   |
| GFR (mL/min/1.73 m²)| 65 (53, 80.498)         | 62 (48.502, 78.33)      | −1.67 (−8.33, 3.25)       | 0.099   |

Abbreviations: BMI, body mass index; SCL, creatinine; AST, aspartate aminotransferase; ALT, alanine aminotransferase; AP, alkaline phosphatase; GFR, glomerular filtration rate.

### Table 3. Changes in immunosuppressant parameters

| Variable                          | All (n = 52) | Tacrolimus (n = 39) | Cyclosporine (n = 13) |
|-----------------------------------|--------------|---------------------|-----------------------|
| **Total Immunosuppressant Daily Dose** |              |                     |                       |
| Decrease                          | 19 (36.5%)   | 16 (41%)            | 3 (23.1%)             |
| Equivalent                        | 18 (34.6%)   | 14 (35.9%)          | 4 (30.8%)             |
| Increase                          | 15 (28.9%)   | 9 (23.1%)           | 6 (46.2%)             |
| **Total Immunosuppressant Daily Dose/Weight (kg)** |              |                     |                       |
| Decrease                          | 34 (65.38%)  | 29 (74.36%)         | 5 (38.46%)            |
| Equivalent                        | 2 (3.85%)    | 1 (2.56%)           | 1 (7.69%)             |
| Increase                          | 16 (30.77%)  | 9 (23.08%)          | 7 (53.85%)            |
| **Immunosuppressant Serum Level** |              |                     |                       |
| Decrease                          | 35 (67.31%)  | 31 (79.49%)         | 4 (30.77%)            |
| Equivalent                        | 0 (0%)       | 0 (0%)              | 0 (0%)                |
| Increase                          | 17 (32.69%)  | 8 (20.51%)          | 9 (69.23%)            |
immunity in LT recipients. Hence, the immunosuppressants may need to be increased accordingly, following the treatment HCV infection.

There are a number of important limitations to our study. First, the study is retrospective, and laboratory test results were not collected using a systematic approach. Nevertheless, this is one of the largest analyses performed on HCV therapy among LT recipients that is applicable and generalizable across several different antiviral regimens. A prospective trial with different antiviral regimens and a systematic laboratory collection would be helpful to clarify this intricate relationship between HCV therapy and immunosuppression. Second, there is no ideal method to estimate immunosuppressant needs. Dosing of TAC and cyclosporine is not standardized and is adjusted based on multiple factors, including renal function, BMI, and individual provider judgment. We chose several novel parameters (total daily dose of the immunosuppressant, the dose of immunosuppressant per the recipient body weight, and product of immunosuppressant and GFR) to estimate immunosuppressant needs in our study. Nonetheless, the results were consistent across all the measured and calculated parameters.

In our study, curing HCV infection in LT recipients resulted in greater immunosuppression needs. Given the availability, tolerability, and success of all-oral interferon-free antiviral regimens, practitioners should be aware of this potential, unanticipated side effect to prevent graft loss. They may consider more frequent and closer monitoring of immunosuppression levels 12 weeks after SVR.

| Parameter                  | Before                  | After                  | Within patient difference | P-value |
|---------------------------|-------------------------|------------------------|---------------------------|---------|
| Tacrolimus                |                         |                        |                           |         |
| TAC TDD (mg)              | 2 (1.5, 4)              | 2.167 (1.583, 3.75)    | 0 (−0.667, 0)             | 0.174   |
| TAC TDD (mg)/Weight (kg)  | 0.03 (0.019, 0.049)     | 0.029 (0.019, 0.045)   | −0.002 (−0.009, 0)        | 0.011   |
| GFR X TAC (mL/min/1.73 m² × mg) | 507.5 (362.285, 615.85) | 348 (216.185, 458.5)   | −147.57 (−281.65, −103.235) | < 0.001 |
| FK Serum Level (ng/mL)    | 8.1 (6.57, 9.1)         | 5.13 (3.965, 6.685)    | −2.4 (−4.27, −1.25)       | < 0.001 |
| Cyclosporine              |                         |                        |                           |         |
| CSA TDD (mg)              | 150 (133.333, 200)      | 166.667 (133.333, 200) | 0 (0, 25)                 | 0.905   |
| CSA TDD (mg)/Weight (kg)  | 1.935 (1.368, 2.813)    | 2.1 (1.531, 2.841)     | 0.153 (−0.063, 0.302)     | 0.505   |
| CSA Serum Level (ng/mL)   | 112 (79, 128.67)        | 105.67 (77, 128.33)    | 5.76 (−15.34, 15)         | 0.78    |
| GFR X CSA (mL/min/1.73 m² × mg) | 6286.67 (5059, 6720.6)  | 5563.33 (5271, 6638.83) | 212 (−1513.66, 815.66)    | 0.944   |

Abbreviations: TAC, tacrolimus; CSA, cyclosporine; GFR, glomerular filtration rate; TDD, total daily dose.
Saab S. et al: Curing Hepatitis C affects post liver transplant immunosuppressant use

Conflict of interest
None

Author contributions
Generated the study concept and design, performed data analysis and interpretation, and drafted the manuscript (SS, JR), contributed to data acquisition (JR, MJ, AF, NJ, SB), critically revised the manuscript for important intellectual content (SS, GC, FD, MEK, SH, RWB), performed statistical analysis (JG, DE), provided administrative support (MJ), supervised the study (SS).

References
[1] Mazzaferrro V, Regalia E, Doci R, Andreola S, Pulvirenti A, Bozzetti F, et al. Orthotopic liver transplantation: the first 60 patients. Br Med J 1977;1:471–476. doi: 10.1136/bmj.1.6059.471.

[2] Watt KD, Pedersen RA, Kremers WK, Heimbach JK, Charlton MR. Evolution of causes and risk factors for mortality post-liver transplant: results of the NIDDK longterm follow-up study. Am J Transplant 2010;10:1420–1427. doi: 10.1111/j.1600-6143.2010.03126.x.

[3] Textor SC, Talier CJ, Canzanello VJ, Schwartz L, Augustine JE. Posttransplantation hypertension related to calcineurin inhibitors. Liver Transpl 2000;6:521–530. doi: 10.1016/s0090-0980(00)00773-7.

[4] Calne RY, Williams R. Orthotopic liver transplantation. 3rd. Fifteen years of clinical liver transplantation. Gastroenterology 1979;77:375–388.

[5] Kuo HT, Sampaio MS, Ye X, Reddy P, Martin P, Bunnapradist S. Risk factors for acute rejection after interferon therapy in liver transplant recipients. Liver Int 2015;35:2442–2447. doi: 10.1111/liv.12856.

[6] Kuo PY, Mantry PS, Cookley E, Te HS, Vargas HE, Brown R Jr. Jr. An interferon-free antiviral regimen for HCV after liver transplantation. N Engl J Med 2014;371:2375–2382. doi: 10.1056/NEJMoa1408921.

[7] Charlton M, Everson GT, Flamm SL, Kumar P, Landis C, Brown RS, et al. SOLAR-1 Investigators. Ledipasvir and Sofosbuvir Plus Ribavirin for Treatment of HCV Infection in Patients With Advanced Liver Disease. Gastroenterology 2015;149:649–659. doi: 10.1053/j.gastro.2015.05.010.

[8] Moini M, Schilsky ML, Tichy EM. Review on immunosuppression in liver transplantation. World J Hepatol 2015;7:1335–1366. doi: 10.4245/wjh.v7.i10.1355.

[9] Lan X, Liu MG, Chen HK, Liu HM, Zeng W, Wei D, Chen P. Efficacy of immunosuppression monotherapy after liver transplantation: a meta-analysis. World J Gastroenterol 2014;20:12330–12340. doi: 10.3748/wjg.v20.i34.12330.

[10] Manzia TM, Angelico R, Ciano P, Mugweru J, Owusu K, Sforza D, et al. Impact of immunosuppression minimization and withdrawal in long-term hepatitis C virus liver transplant recipients. World J Gastroenterol 2014;20:12217–12225. doi: 10.3748/wjg.v20.i34.12217.

[11] Landofo MC, Rimola A, O’Grady J, Sanchez-Fueyo A. Immunosuppression minimization vs. complete drug withdrawal in liver transplantation. J Hepatology 2013;59:872–879. doi: 10.1016/j.jhep.2013.04.003.

[12] Liu Z, Chen Y, Tao R, Xv J, Meng J, Yong X. Tacrolimus-based versus cyclosporine-based immunosuppression in hepatitis C virus-infected patients after liver transplantation: a meta-analysis and systematic review. PLoS One 2014;9: e107057. doi: 10.1371/journal.pone.0107057.

[13] Badi P, Dutta S, Coakley E, Cohen D, Ding B, Podsadecki T, et al. Pharmacokinetics and dose recommendations for cyclosporine and tacrolimus when coadministered with ABT-450, ombravir and dasabuvir. Am J Transplant 2015;15:1313–1322. doi: 10.1111/ajt.13111.

[14] Reddy KR, Everson G, Flamm S, Denning J, Arterburn S, Brandt-Sarf T, et al. Ledipasvir/Sofosbuvir with Ribavirin for the Treatment of HCV in Patients with Post Transplant Recurrence: Preliminary Results of a Prospective, Multicenter Study. Hepatology 2014;60:2024–2030. doi: 10.1002/hep.27234.

[15] Crespo G, Marino Z, Navasa M, Forns X. Viral hepatitis in liver transplantation hypertension related to calcineurin inhibitors. Liver Transpl 2000;6:865–879. doi: 10.1011/lcj.2011.02111.

[16] Berenguer M, Prieto M, Rayón JM, Mora J, Pastor M, Ortiz V, et al. Natural history of clinically compensated hepatitis C virus-related graft cirrhosis after liver transplantation. Hepatology 2000;32:852–858. doi: 10.1001/hjhep.2000.17924.

[17] Crespo G, Marino Z, Navasa M, Forns X. Viral hepatitis in liver transplantation hypertension related to calcineurin inhibitors. Liver Transpl 2000;6:865–879. doi: 10.1011/lcj.2011.02111.

[18] Levitsky J, Fiel MI, Norvell JP, Wang E, Watt KD, Curry MP, et al. Risk for immunosuppressed graft dysfunction in liver transplant recipients with recurrent HCV infection treated with pegylated interferon. Gastroenterology 2012;142:1132–1139. doi: 10.1053/j.gastro.2012.01.030.

[19] Berenguer M, Palau A, Aguilera V, Rayón JM, Juan FS, Prieto M. Clinical benefits of antiviral therapy in patients with recurrent hepatitis C following liver transplantation. Am J Transplant 2008;8:679–687. doi: 10.1111/j.1600-6143.2007.02126.x.

[20] Carrió JA, Navasa M, García-Retortillo M, García-Pagan JC, Crespo G, Brugueras M, et al. Efficacy of antiviral therapy on hepatitis C recurrence after liver transplantation: a randomized controlled study. Gastroenterology 2007;132:1746–1756. doi: 10.1053/j.gastro.2007.03.041.

[21] Saab S, Kalmaz D, Gajjar NA, Hiatt J, Durazo F, Han S, et al. Outcomes of acute rejection after interferon therapy in liver transplant recipients. Liver Transpl 2004;10:869–867. doi: 10.1002/lt.20157.

[22] Selzner N, Guindi M, Renner EL, Berenguer M. Immune-mediated complications of the graft in interferon-treated hepatitis C positive liver transplant recipients. J Hepatol 2011;55:207–217. doi: 10.1016/j.jhep.2010.11.012.

[23] Levitsky J, Fiel MI, Norvell JP, Wang E, Watt KD, Curry MP, et al. Risk for immunosuppressed graft dysfunction in liver transplant recipients with recurrent HCV infection treated with pegylated interferon. Gastroenterology 2012;142:1132–1139. doi: 10.1053/j.gastro.2012.01.030.
Abdel-Hakeem MS, Bédard N, Badr G, Ostrowski M, Sékaly RP, Bruneau J, et al. Comparison of immune restoration in early vs. late alpha interferon therapy against hepatitis C virus. J Virol 2010;84:10429-10435. doi: 10.1128/JVI.01094-10.

Badr G, Bédard N, Abdel-Hakeem MS, Trautmann L, Willems B, Villeneuve J-P, et al. Early interferon therapy for hepatitis C virus infection rescues polyfunctional, long-lived CD8+ memory T cells. J Virol 2008;82:10017-10031. doi: 10.1128/JVI.01083-08.

Bohne F, Londoño MC, Benítez C, Miquel R, Martínez-Llordella M, Russo C, et al. HCV-induced immune responses influence the development of operational tolerance after liver transplantation in humans. Sci Transl Med 2014;6:242ra81. doi: 10.1126/scitranslmed.3008793.