Multidisciplinary Approach to treat Liver Transplant Recipients with Hepatitis C using Sofosbuvir based Therapy

Abstract

Introduction: The safety and efficacy of the direct acting antivirals in liver transplant recipients is unknown.

Materials and Methods: Retrospective cohort study on 44 liver transplant recipients, ≥18 years old who received Sofosbuvir (SOF) based antiviral therapy (AVT) at our hospital from January 2012 through May 2016. Multidisciplinary (MDT) approach involving hepatologists, transplant pharmacists and a patient access coordinator was adapted. SVR at 12 and 24 weeks, safety and compliance of SOF based AVT were reported. We also reported on improvement in APRI and CPT score at SVR.

Results: 35 patients (79.5%) were treated with Ledipasvir (LDV) /SOF 7 (16 %) with SOF/RBV and 2 (4.5%) with SOF/Simeprevir (SIM). Most patients were HCV genotype 1 (n=37; 84.1%) and treatment experienced (n= 24; 55%). Median time between liver transplant (LT) and AVT was 2.7 years with majority (66%) treated ≥ 12 months after LT. Median pre-treatment HCV viremia was 3,800,000 (71,000-66,000,000) with a high viral load ≥ 800,000 copies/ml in 36 (82%) patients. SVR 12 and 24 were achieved in 43/44 (98%), despite a low (77.3%) rapid virological response. Statistically significant improvement was observed in pre-treatment and post- SVR median albumin levels, APRI and CPT scores. Fatigue (27%) was the most commonly reported adverse effect. Compliance rate was 100%.

Conclusion: With a MDT approach, using SOF based AVT, a very high SVR was achieved in liver transplant recipients with recurrent HCV infection. A MDT approach positively impacts medication acquisition, enhances patient education and compliance in post LT HCV management.

Keywords: Multidisciplinary; HCV; Liver transplantation; Antiviral therapy; Direct antiviral agents; Sofosbuvir

Abbreviations: SOF: Sofosbuvir; AVT: Antiviral Therapy; MDT: Multidisciplinary Team; LDV: Ledipasvir; SIM: Simeprevir; HCV: Hepatitis C Virus; LT: Liver Transplantation; IS: Immunosuppressive; SVR: Sustained Virological Response; GT: Genotype; DAA: Direct Acting Antiviral; CPT: Child-Pugh-Turcotte; TP: Transplant Pharmacist; PAC: Patient Access Coordinator; AASLD: American Association for the Study of Liver Diseases; RBV: Ribavirin; APRI: AST to Platelet Ratio Index; LTRs: Liver Transplant Recipients

Introduction

Hepatitis C virus (HCV) presents a major health care challenge and is the leading indication for liver transplantation (LT) in the United States [1,2]. Recurrence of HCV after LT is universal and contributes immensely to graft failure and early mortality due to immunosuppressive (IS) therapy related acceleration of fibrosis [3]. Treatment with AVT and eradication of virus is the only way to improve the outcomes related to recurrence of HCV.

Until 2011, pegylated interferon plus ribavirin was the standard of care for HCV treatment, resulting in a sustained virological response (SVR) in around 30%-50% of transplant recipients with HCV genotype 1 (GT) [4-11]. A major change in paradigm has been observed with the discovery of safe and effective direct acting antiviral (DAA) therapy in the field of HCV treatment [12]. These DAA have already been shown to change the landscape of AVT for HCV infection in non-transplant and post LT settings. Recently published randomized controlled studies on the use of sofosbuvir based AVT reported SVR rates ranging between 70% to 85% in post LT patients with child-pugh-turcotte (CTP) A, B and C patients [4,13,14]. Overall, the treatment was well tolerated, with few severe side effects. Similar results were reported in studies based on real world data such as TRIO and TARGET 2.0 trials [15-18]. A recent real world data-based study published from Canada on efficacy of SOF-based treatment reported overall SVR rates of 85% in all genotypes [19]. However, these results are not as encouraging as reported in non-transplant setting with SVR of ~95% [20-22]. Besides HCV genotype, and stage of fibrosis, compliance to therapy is a very important predictive factor of SVR [23]. Post LT patients are on multiple medications besides...
their immunosuppression including prophylaxis for bacterial and viral infections, bisphosphonate, antihypertensive, anti-diabetic medications, etc. They also can suffer from memory impairment due to their post- transplant course, previous history of drinking and IS use. Moreover, substance abuse and psychiatric illnesses are also common in hepatitis C patients [24]. All these factors may lead to issues with adherence to very expensive DAA for recurrent HCV infection resulting into lesser efficacy. Multidisciplinary team approach has been studied and reported to be successful in the era of interferon based AVT for patients with HCV infection in a non-transplant setting [25]. Improvement in patient compliance and the efficiency of antiviral treatment has already been reported in the non-transplant setting.

Recent advancements in discovery of various DAA may allow more patients to have access to life altering medications. However, many hurdles exist including high cost of treatment, drug-drug interaction and availability of long-term data in transplantation [26-28]. Once the medication is approved, there exists a litany of drug interactions in the transplant population [28]. Moreover, logistic and compliance issues abound above may also contribute to outcomes of the DAA based treatment in recurrent HCV patients. In order to tailor a SOF based AVT regimen in patients with recurrent HCV infection, we employed a MDT approach from the initial encounter, approval of medication to the monitoring of drug safety, interaction, compliance and ultimately completion of AVT. Our MDT approach included involvement of Hepatologists, Transplant Pharmacists (TP) and Patient Access Coordinator (PAC) in care of post LT HCV patients.

Materials and Methods

Study Population and Design

This is a retrospective, single center cohort study approved by the Institutional Review Board of our institution. All adult (age ≥18) liver transplant recipients (LTRs) (treatment experienced and naïve) with recurrent HCV infection treated with SOF-based AVT between January 2012 to May 2016 were included. Recurrent HCV infection was diagnosed based on HCV viremia in the post LT setting. Based on the American Association for the Study of Liver Diseases (AASLD) guidelines, a liver biopsy is not a prerequisite to qualify for treatment.

The patients in the current study were treated with SOF based AVT with or without Ribavirin (RBV) for 12 or 24 weeks. Regimens of SOF based AVT included LDV/SOF (n=35), SOF/RBV (n=7) and SOF/SIM, (n=2). Ribavirin dose was adjusted according to weight of patient and their renal function. Starting dose of RBV was 600 mg daily and increased to 1000 mg if patient weighs <75 kg or 1200 mg in patients weighing ≥75 kg. This dose could be reduced based on the tolerance and level of hemoglobin. No changes were made to the ongoing IS regimen. Patients with chronic kidney disease (GFR < 30 mL/min), HIV/HCV co-infection, HBV/HCV co-infection, prisoners and pregnant females were excluded.

Multi-disciplinary approach

Patients were assessed by a Hepatologist in the transplant clinic for SOF based AVT. Based on the eligibility criteria, patients underwent required laboratory testing in order to prepare for the prior authorization required for AVT. Patients were assigned treatment based on the ASL/D/ Infectious diseases society of America (IDSA) guidelines and practitioner expertise in a non-randomized process. All patients eligible for SOF based DAA were referred to a TP and PAC. The purpose of referral to pharmacist was to further emphasize on the compliance of AVT, reviewing insurance approval procedure, identifying drug-drug interactions, and educate the patient on possible side effects of treatment. The PAC ensured that appropriate labs were ordered after explaining their necessity to patients and obtained their consent to draw. After appropriate labs were obtained, the prescription for the medication(s) was sent to The Ohio State University Specialty Pharmacy in order to initiate the prior authorization process. At this point, a specialized prior authorization technician filled out the proper forms with resulted laboratory values and reason for HCV treatment. If initially denied, the technician collaborated with the Hepatologist and TP to compose a letter outlining the need for HCV treatment. Throughout this time period, the PAC kept communication and support open to the patient while they waited for approval. After treatment approval, the TP at the Specialty Pharmacy would contact the patient to reinforce education provided during initial assessment visit. On each follow up clinic visit, TP monitored for response to treatment, possible drug interactions, need for medication assistance program referral and/or therapy modification.

Efficacy and safety assessment

Blood work including complete blood counts, creatinine, electrolytes, liver function tests and INR were monitored at baseline and then at every 4 week after the start of SOF based AVT. Complete blood count was monitored every other week in patients on RBV and the dose was adjusted according to hemoglobin levels. HCV RNA PCR was checked using ELISA immunoassay at week 4, week 12 or end of treatment (limit of detection, up to 12 IU/ml). SVR was considered as an undetectable HCV viral load 12 and 24 weeks after the end of treatment. Levels of calcineurin inhibitor were also monitored during and after the treatment as per our transplant center protocol.

Study outcomes

Sustained virological response at weeks 12 and 24 in LTRs with recurrent hepatitis C was reported. Secondary outcomes included side effects, drug-drug interactions and compliance. We also reported improvement in AST to platelets ratio index (APRI) score, responses at 4 weeks & at the end of treatment, and improvement in post AVT albumin levels.

Statistical analysis

Patient characteristics are presented as median and range for continuous variables and number and percent for categorical variables. The percentage of patients achieving SVR-12 and -24 is reported. Pre- and post -AVT variables including AST to platelets ratio index score, and albumin were compared, using Wilcoxon sign rank tests. Pre-AVT and post SVR CTP score were also compared using McNemar’s tests for paired data.

Results

Baseline characteristics

A total of 133 LTs were performed at our institute during the
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study period. Of these 59 (44%) underwent LT for HCV cirrhosis related complications; 44 with recurrent HCV infection were treated with SOF-based AVT. Of 37 GT-1 patients 35 (79.5%) were treated with SOF/LDV and two (4.5%) received SOF/SIM. Seven (16%) with GT-3 treated with SOF/RBV and. The majority of patients were placed on 24 weeks of therapy, however, due to insurance constraints, 2 patients in the SOF/LDV group and 1 patient in the SOF/SIM group were treated for 12 weeks with the addition of RBV. Baseline characteristics of the patient population are summarized in Table 1. Majority (79.5%) were males with a median age of 61 years (range, 41-70 years). Median HCV viral level was 3,800,000 (71,000-66,000,000) copies /ml and 36 (82%) had a high baseline viral load of ≥ 800,000 copies/ml. Thirty-seven (84%) were genotype 1a and of these 29 (78%) had GT-1A infection. Severity of liver disease was assessed with the help of APRI and CTP score. Median APRI score of our cohort was 0.78 (range: 0.21-7.36). There were 40 (91%) and 4 (9%) patients with CTP-A and CTP-B, respectively. None of the patients had decompensated liver disease at initiation of AVT. There were 24 (54.6) treatment experienced and 20 (45.4%) treatment naïve patients in our cohort. Median time between LT and start of AVT was 2.7 years and 34 % got treatment within 12 months from LT.

Table 1: Baseline characteristics of patients.

| Characteristic          | Median (Range) or n (%) | n=44 |
|-------------------------|-------------------------|------|
| Age (years)             | 61 (41-70)              |      |
| Gender                  | Female/Male             | 9 (20.5)/ 35 (79.5) |
| Race                    | African American        | 8 (18.2) |
|                         | White                   | 36 (81.8) |
|                         | BMI                     | 29 (17-67) |
| Diagnosis               | HCV only                | 14 (31.8) |
|                         | HCV/ETOH                | 6 (13.6) |
|                         | HCV/HCC                 | 24 (54.6) |
|                         | Genotype-1              | 37 (84) |
|                         | IA & IB                 | 29 (78.4) & 8 (21.6) |
|                         | Genotype Non-1          | 7 (16) |
| Immunosuppression       | Tacrolimus              | 21 (47.7) |
|                         | Cyclosporine            | 23 (52.3) |
|                         | Mycophenolate Mofetil   | 40 (91%) |
| Previous Treatment      | Treatment naïve         | 20 (45.4) |
|                         | Treatment experienced   | 24 (54.6) |
|                         | SOF-based treatment     | 35 (79.5) |
|                         | SOF/LDP                 |      |

SOF/RBV                  | 7 (15.9) |
SOF/SIM                  | 2 (4.5)  |
Cirrhosis                | 3 (6.8)  |
HCV PCR > 800K copies/ml | 36 (81.8) |
Inflammation grade >1    | 11 (25)  |
Fibrosis stage ≥ F1      | 16 (36.4) |
Median time between LT and AVT (yrs) | 2.7 (0.5 - 17) |
Time from LT to AVT <12months | 15 (34.1) |

**Treatment response**

Overall 43 (97.7%) patients out of 44 treated achieved SVR-12 and -24. Of these 32 (72.7%) achieved undetectable HCV RNA at week-4 and 43 (97.7%) had end of treatment response. No patients experienced virological breakthrough while on SOF-based regimens or relapsed. The only patient who did not respond to a 24 weeks of SOF/RBV combination had HCV GT-3 infection and was treatment naïve. There were no differences in SVR rates for subgroups based on age (<60 vs ≥60 years), gender (male vs female), ethnicity (White vs African American), prior treatment status (treatment naïve vs treatment experienced) and HCV viremia (<800K vs ≥800K) levels (Figure 1). There was statistically significant improvement in median APRI score (0.78, range: 0.21-7.36 to 0.32, range: 0.12-1.15, p<0.001), median albumin (3.9 grams/L range: 2.6-4.8 to 4.0 grams/L, range: 3.0-5.0, p=0.01), and CTP score between pre-AVT and after achieving SVR (Table 2).

![Figure 1: Sustained Virological response is not dependant on age, gender, race, HCV viremia and prior treatment response.](https://example.com/figure1.png)
Table 2: Changes in pre AVT and post SOF-based treatment parameters.

| Lab Measure             | Pre SOF (n=43) Median (Ranges) | 24 weeks Post SOF (n=43) Median (Ranges) | p-value* |
|-------------------------|--------------------------------|------------------------------------------|----------|
| Hemoglobin (gm/L)       | 12.8 (8.5-15.4)                | 12 (8.6-16.9)                            | 0.94     |
| INR**                   | 1.0 (0.9-1.6)                  | 1.0 (0.9-1.2)                            | <0.001   |
| Creatinine (mg/dl)      | 1.31 (0.62-2.60)               | 1.22 (0.70-2.40)                         | 0.13     |
| ALT (IU)                | 46 (11-322)                    | 17 (6-51)                                | <0.001   |
| AST (IU)                | 48 (14-377)                    | 20 (12-46)                               | <0.001   |
| Alkaline Phosphate (IU) | 89 (8-308)                     | 80 (45-190)                              | 0.01     |
| Creatinine (mg/dl)      | 0.8 (0.3-2.8)                  | 0.7 (0.1-1.8)                            | 0.11     |
| Albumin (g/L)           | 3.9 (2.6-4.8)                  | 4.1 (3.0-5.0)                            | 0.001    |
| CTP Score               | 5 (5-7)                        | 5 (5-6)                                  | 0.02     |
| APRI Score              | 0.78 (0.21-7.36)               | 0.32 (0.12-1.15)                         | <0.001   |
| HCV PCR                 | 3,800,000 (71,000-66,000,000)  | 6 (0-286)                                | <0.001   |

*Sign rank tests for paired differences

**Median INR was equal but distribution was higher for Pre SOF.

Safety

Treatment was very well tolerated with majority n=28 (63.6%) reporting no side effects (Table 3). Sixteen patients (36%) reported adverse events, all of which were mild and did not require treatment discontinuation. In those reporting adverse events, the most common were fatigue (27%), headache (9%) and insomnia (9%). None of the patients had graft rejection or died and all were able to complete the treatment.

Table 3: Adverse events on SOF based anti-viral therapy.

| Side Effects       | n=44 (%) |
|--------------------|----------|
| Fatigue            | 12 (27.3)|
| Headache           | 4 (9.1)  |
| Nausea             | 3 (6.8)  |
| Insomnia           | 4 (9.1)  |
| Anemia             | 3 (6.8)  |
| Decompensation     | 0        |
| Treatment Discontinuation | 0   |
| No Adverse Effects | 28 (63.6)|

Compliance

With the help of our multidisciplinary team approach all patients were able to complete the SOF based AVT. None dropped out, reported missing a single dose of the medication or was lost to follow-up. Therefore our study medication compliance was 100%.

Discussion

In this study, we describe our experience of treating recurrent HCV infection in LTRs including mainly genotypes 1 and 3 with SOF-based regimens adapting a multi-disciplinary approach. Very high SVR rates were achieved with the help of SOF-based AVT and treatment was very well tolerated with 100% compliance rate. During treatment, there were no antiviral therapy related serious adverse effects or significant drug-drug interactions noted. Only one patient was unable to achieve SVR belonging to HCV genotype 3, which has emerged as the most challenging of all HCV genotypes to treat despite the introduction of newer direct-acting antiviral therapies.

Previous studies have shown that overall graft and patient survival are lower in liver transplant recipients with HCV infection than among recipients without HCV infection, due to its universal recurrence and rapid progression to cirrhosis. In the present study, the SVR rates were comparable among all treatment regimens, and the addition of RBV had no impact on treatment or SVR rates. SVR at 24 weeks in this entire cohort was 98%. This is comparable to preliminary results of a phase 2 study presented at the 2014 AASLD meeting, using a fixed dose combination of LDV/SOF with RBV for 12 or 24 weeks post-LT, patients with genotype 1 or 4 with recurrent HCV post transplantation [14]. Our results are consistent with a recently published trial which evaluated combination of ronivir with paritaprevir, ombitasvir, plus
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References

1. WHO (2017) Hepatitis C: fact sheet no. 164. World Health Organization, Geneva.

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With the help of albumin, APRI and CTP score in our cohort after successful completion of AVT. Study by Pellicelli et al also showed improvement in CTP and MELD scores after completion of therapy, which is consistent with our findings [30].

Furthermore, most factors previously associated with inferior response such as age, gender, ethnicity, viral load and previous treatment history don't hold the same importance with SOF-based regimens as they did for interferon-based therapy. Our findings suggest that most baseline characteristics and lack of viral response during AVT may no longer be relevant predictive factors when treating patients with SOF-based regimens. A very recent study by Welzel et al. [36] also showed that on-treatment HCV RNA quantification is of limited clinical use in patients with advanced liver disease and/or liver transplantation and does not predict SVR12 [36]. Faisal et al. [19] have reported similar results in their observational study, which evaluated the efficacy, safety and tolerability of regimens containing sofosbuvir in the treatment of HCV recurrence in all genotypes outside of clinical trials in all Canadian transplant centers [19].

In our study, doses of calcineurin inhibitors were monitored during the treatment period to maintain therapeutic levels and no dose adjustment was required. No deaths or episodes of graft rejection occurred in this population. The most common side effects were fatigue, headache and insomnia, seen in 27%, 9% and 9%, respectively; no patients discontinued treatment due to adverse events. There was no need for blood products for patients on RBV.

To our knowledge this is the first study reporting on role of multidisciplinary team approach including hepatologists, specialty pharmacist and a patient access coordinator in LTRs using DAA for recurrent HCV infection. An earlier study in Interferon based AVT era also assessed the importance of MDT comprising two hepatologists, two nurses, one pharmacist, one psychologist, one administrative assistant, and one psychiatrist. They reported increased efficiency and compliance in MDT approach as compared to historic control group [25]. The management of HCV infections is centered on pharmacotherapy, making the clinical pharmacist appropriately suited as part of the patient’s care team. Direct acting antiviral medications used to treat HCV-infected patients are effective in achieving SVR. However, the monitoring of adverse effects, significant drug interactions, and high risk of non-adherence and treatment discontinuation demands a transplant clinical pharmacist involvement in managing this expensive treatment [37]. Accumulating evidence suggests that a pharmacist as part of a multidisciplinary team can have beneficial effects on patient care. Pharmacist-provider team-based care is widely supported in the literature, demonstrating significant improvement in cardiovascular and renal outcomes [38-40]. Our findings suggest that role of a pharmacist based education and a patient access coordinator in facilitation on medication acquisition and improved patient outcomes, which reflects the importance of adopting a multidisciplinary approach in HCV management. We found that median time between LT and AVT therapy start was 2.7 years and 34% were able to get the treatment in less than 12 months after LT. Early acquisition of DAA was achieved with the help of our access coordinator which ultimately resulted in improved treatment efficacy and compliance.

There are a few limitations that should be considered in the interpretation of this report, including its small sample size, which is insufficient to allow subgroup comparisons and characterization of efficacy in certain populations (i.e. African Americans, HCV genotype 3). In addition, our population consisted of patients with well-compensated liver disease, so we could not study the efficacy of DAA in patients with advanced disease. Lastly, we did not have a comparison cohort to compare the MDT approach and assess its role.

Conclusion

In conclusion, a multidisciplinary team approach was found to be helpful in improving patient compliance and increases the efficiency of SOF-based antiviral therapy. We found that a close cooperation among the healthcare providers in the care of post LT patients with recurrent HCV infection can ensure optimal treatment performance and high SVR rates.
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2. Scientific Registry of Transplant Recipients 2012 annual data report. Rockville, MD: Health Resources and Services Administration, 2012.

3. AASLD/IDSA/IAS-USA (2015) Recommendations for testing, managing, and treating hepatitis C.

4. Kwo PY, Mantry PS, Coakley E, Te HS, Vargas HE, et al. (2014) An interferon-free antiviral regimen for HCV after liver transplantation. New Engl J Med 371(25): 2375-2382.

5. Gane EJ, Agarwal K (2014) Directly acting antivirals (DAAs) for the treatment of chronic hepatitis C virus infection in liver transplant patients: “a flood of opportunity.” Am J Transplant 14(5): 994-1002.

6. Gordon FD, Kwo P, Ghalib R, Crippin J, Vargas HE, et al. (2012) Peginterferon-a-2b and ribavirin for hepatitis C recurrence postorthotopic liver transplantation. J Clin Gastroenterol 46(8): 700-706.

7. Xiao N, Shi S, Zhuang H (2010) A meta-analysis that compares the use of either peginterferon-a2a or peginterferon-a2b plus ribavirin for HCV infection. Hepat Med 28: 99-109.

8. Ferenci P, Formann E, Laferl H, Gschwantler M, Hackl F, et al. (2006) Randomized, double-blind, placebo-controlled study of peginterferon alfa-2a (40KD) plus ribavirin with or without amantidine in treatment-naïve patients with chronic hepatitis C genotype 1 infection. J Hepatol 44(2): 275-282.

9. Carrion JA, Navasa M, Garcia-Retortillo M, Garcia-Pagan JC, Crespo G, et al. (2007) Efficacy of antiviral therapy on hepatitis C recurrence after liver transplantation: a randomized controlled study. Gastroenterology 132(5): 1746-1756.

10. Charlton M, Gane E, Manns MP, Brown RS Jr, Curry MP, et al. (2015) Sofosbuvir and ribavirin for treatment of compensated recurrent hepatitis C virus infection after liver transplantation. Gastroenterology 148(1): 108-117.

11. Ascha M, Ascha M, Zein N, Alkhouri N, Eghtesad B, et al. (2015) Treatment of recurrent hepatitis C genotype-4 post-liver transplantation with sofosbuvir plus simprevir. Int J Organ Transplant Med 6(2): 86-90.

12. Crespo G, Marino Z, Nacasa M, Foras X. Viral hepatitis in liver transplantation. Gastroenterology 142(6): 1373-1383.

13. Manns M, Foras X, Samuel D, Denning J, Aterburn S, et al. (2015) Ledipasvir/sofosbuvir with ribavirin is safe and efficacious in decompensated and post-liver transplant patients with HCV infection: preliminary results of the SOLAR-2 trial. Program and abstracts of the 50th Annual Meeting of the European Association for the Study of the Liver; Vienna, Austria. G02.

14. Charlton M, Everson GT, Flamme SL, Kumar P, Landis C, et al. (2015) Ledipasvir and sofosbuvir plus ribavirin for treatment of HCV infection in patients with advanced liver disease. Gastroenterology 149(3): 649-659.

15. Backus LI, Belperio PS, Shahoumian TA, Loomis TP, Mole LA (2015) Effectiveness of sofosbuvir-based regimens in genotype 1 and 2 hepatitis C virus infection in 4026 U.S. Veterans. Aliment Pharmacol Ther 42(5): 559-573.

16. Höner Zu Siederdissen C, Maasoumy B, Deterding K, Port K, et al. (2015) Eligibility and safety of the first interferon-free therapy against hepatitis C in a real-world setting. Liver Int 35(7): 1845-1852.

17. Jensen DM, O’Leary JG, Pockros P, Sherman K, Kwo P, et al. (2015) Safety and efficacy of sofosbuvir-containing regimens for hepatitis C: real-world experience in a diverse, longitudinal observational cohort. 65th Annual Meeting of the American Association for the Study of Liver Diseases. Boston, USA. p. 45.

18. Dieterich D, Bacon BR, Flamme SL, Kowdley K, Milligan S, et al. (2014) Evaluation of sofosbuvir and simprevir-based regimens in the TRICT network: academic and community treatment of a real-world, heterogeneous population. 65th Annual Meeting of the American Association for the Study of Liver diseases. Boston, USA. p. 46.

19. Faisal N, Bilodeau M, Aljudaibi B, Hirsch G, Yoshida EM, et al. (2016) Sofosbuvir-Based Antiviral Therapy Is Highly Effective In Recurrent Hepatitis C in Liver Transplant Recipients: Canadian Multicenter “Real-Life” Experience. Transplantation. 100(5): 1059-1065.

20. Lawitz E, Poordad F, Pang PS, Hyland RH, Ding X, et al. (2014) Sofosbuvir and ledipasvir fixed-dose combination with and without ribavirin in treatment-naïve and previously treated patients with genotype 1 hepatitis C virus infection (LONESTAR): an open-label, randomised, phase 2 trial. Lancet 383(9916): 515-523.

21. Afshar N, Zeuzem S, Kwo P, Chojkier M, Gitlin N, et al. (2014) Ledipasvir and sofosbuvir for untreated HCV genotype 1 infection. N Engl J Med 370(20): 1880-1898.

22. Reddy KR, Bourliere M, Sulkowski M, Omata M, Zeuzem S, et al. (2015) Ledipasvir and sofosbuvir in patients with genotype 1 hepatitis C virus infection and compensated cirrhosis: an integrated safety and efficacy analysis. Hepatology 62(1): 79-86.

23. Lai M, Afshar NH (2012) Clinical utility of interferon-free-28B testing in patients with genotype 1. Hepatology 56(1): 367-372.

24. Martin-Santos R, Diez-Quevedo C, Castelvil P, Navinés R, Miquel M, et al. (2008) De novo depression and anxiety disorders and influence on adherence during peginterferon-alpha-2a and ribavirin treatment in patients with hepatitis C. Alimentary Pharmacology and Therapeutics 27(3): 257-265.

25. Carrion JA, Gonzalez-Colominas E, Garcia-Retortillo M, Cañete N, Ciren L, et al. (2013) A multidisciplinary support program increases the efficiency of peglated interferon alfa-2a and ribavirin in hepatitis C. J Hepatol 59(5): 926-933.

26. Zhao YJ, Khoo AL, Lin L, Tang M, Koh CJ, et al. (2016) Cost-Effectiveness of Strategy-based Approach to Treatment of Genotype 1 Chronic Hepatitis C J Gastroenterol Hepatol 31(9): 1628-1637.

27. Lo Re V, Gowda C, Urick PN, Halladay JT, Binkley A, et al. (2016) Disparities in Absolute Denial of Modern Hepatitis C Therapy by Type of Insurance. Clin Gastroenterol Hepatol 14(7): 1035-1043.

28. The New York Times (2016) Costly Hepatitis C Drugs for Everyone?

29. Pellicelli AM, Montalbano M, Lionetti R, Durand C, Ferenci P, et al. (2014) Sofosbuvir plus daclatasvir for post-transplant recurrent hepatitis C: potent antiviral activity but no clinical benefit if treatment is given late. Dig Liver Dis 46(10): 923-927.

30. Pungpapong S, Aqel B, Leive M, Werner KT, Murphy JL, et al. (2015) Multicenter experience using simprevir and sofosbuvir with or without ribavirin to treat hepatitis C genotype 1 after liver transplant. Hepatology 61(6): 1880-1886.

31. Gutierrez JA, Carrion AF, Avalos D, O’Brien C, Martin P, et al. (2015) Sofosbuvir and simprevir for treatment of hepatitis C virus infection in liver transplant recipients. Liver Transpl 21(6): 823-830.

32. Jacobson IM, Gordon SC, Kowdley KV, Yoshida E, Rodriguez-Torres M, et al. (2013) Sofosbuvir for hepatitis C genotype 2 or 3 in patients without treatment options. New England J Med 368: 1867-1877.

33. Foster GR, Planko S, Brown A, Forton D, Nahass RG, et al. (2015) Efficacy of Sofosbuvir plus Ribavirin With or Without Peginterferon-Alpha in Patients With Hepatitis C Virus Genotype 3 Infection and Treatment-Experienced Patients With Cirrhosis and Hepatitis C Virus Genotype 2 Infection. Gastroenterology 149(6): 1462-1470.
34. Welzel TM, Reddy KR, Flamm SL, Denning J, Lin M, et al. (2016) On-treatment HCV RNA in patients with varying degrees of fibrosis and cirrhosis in the SOLAR-1 trial. Antivir Ther 21(6): 541-546.

35. Mohammad R, Bulboch M, Chan J, Deming P, Love B, et al. (2014) Provision of Clinical Pharmacist Services for Individuals With Chronic Hepatitis C Viral Infection. Pharmacotherapy 34(12): 1341-1354.

36. Odum L, Whaley A (2012) The Role of Team-Based Care Involving Pharmacists to Improve Cardiovascular and Renal Outcomes. Cardiorenal Med 2(4): 243-250.

37. Leung WY, So WY, Tong PC, Chan NN, Chan JC (2005) Effects of structured care by a pharmacist-diabetes specialist team in patients with type 2 diabetic nephropathy. Am J Med 118(12): 1414.

38. Bozovich M, Rubino CM, Edmunds J (2000) Effect of a clinical pharmacist-managed lipid clinic on achieving national cholesterol education program low-density lipoprotein goals. Pharmacotherapy 20(11): 1375-1383.

39. Joy MS, Dehart RM, Gilmartin C, Hachey D, Hudson J, et al. (2005) Clinical pharmacists as multidisciplinary health care providers in the management of CKD: a joint opinion by the Nephrology and Ambulatory Care Practice and Research Networks of the American College of Clinical Pharmacy. Am J Kidney Dis 45(6): 1105-1118.