Cyclosporine and Hepatitis C

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Received September 24, 2012; revised October 27, 2012; accepted November 7, 2012

ABSTRACT
End stage liver disease from hepatitis C is a leading indication for liver transplantation. Recurrent hepatitis C after liver transplantation may lead to cirrhosis and graft failure in up to 25% of recipients five years after liver transplantation. Antiviral therapy is challenging after liver transplantation due to increased side effects including cytopenias and decreased efficacy compared to the nontransplant population. Tacrolimus and cyclosporine are the most common immunosuppressants used to prevent graft rejection. Tacrolimus is more potent than cyclosporine and may be preferred to cyclosporine. However, cyclosporine may have activity against hepatitis C and may have a theoretical advantage to tacrolimus in hepatitis liver transplant recipients. Cyclosporine may inhibit NS5B and NS5A protein complexes and increase endogenous interferon activity. Cyclophilin inhibitors without immunosuppressive properties are under development and represent a novel mechanism for inhibiting HCV replication.

Keywords: Immunosuppression; Liver; Transplant

1. Introduction
Hepatitis C virus (HCV) infects an estimated 170 million people worldwide and is the single most common indication for orthotopic liver transplantation (OLT) in Europe and the United States [1,2]. Indeed, end stage liver disease from hepatitis C is the indication for liver transplantation in approximately 40% of liver transplant candidates. Although up to 40% of patients who develop acute HCV infection may spontaneously clear infection; in contrast, after liver transplantation HCV reinfection post-transplant is universal [3-6]. An estimated 20% of liver transplant recipients with chronic HCV infection develop cirrhosis 4 - 5 years post-liver transplant [6,7]. The accelerated course of hepatitis C may be due age of the donor liver, gender, and viral properties, as well as immunosuppression [3,5,7-9].

2. Hepatitis C Replication and Calcineurin Inhibitors
HCV is a hepatotropic member of the flaviviridae family. The virus is transmitted parenterally and rarely through vertical and sexual transmission [4]. HCV possesses a positive strand RNA genome of about 9.6 kb in length. The genome encodes a polyprotein of about 3000 amino acids that is then cleaved by host and viral proteases to generate functional viral proteins, namely: core, two envelope proteins (E1 & E2), and seven nonstructural proteins (p7, NS2, NS3, NS4A, NS4B, NS5A, and NS5B) [4,10]. The core and envelope proteins compose of an N-terminal triad and are the major constituents of the extracellular virion and involved in viral assembly and entry. The nonstructural proteins NS3/NS4A, NS5A, NS5B are involved in viral replication and the target of direct acting antiviral agents [10]. The NS3/4A protein, a serine protease, and NS5B protein, RNA-dependent RNA polymerase are thought to play a central role in viral genome replication. NS2 is a protease that mediates the cleavage of the polyprotein at the NS2-3 junction. The p7 proteins form an ion channel thought to be critical for secretion of infection virus. [10].

The lack of a robust animal model hindered the development of therapy against HCV, but the creation of the HCV replicon model overcame many of these limitations. HCV attaches to the host cell surface by utilizing at least four receptors, CD81, scavenger receptor class B type I (SR-BI), claudin-1 (CLDN1), and occludin (OCLN) [4]. HCV attaches itself to cells by low-affinity attachment to glycosaminoglycans and lectins. Then LDL receptors induce endocytosis and within the endocytic vesicle HCV is exposed to low pH, that induces conformational changes and high affinity to receptors such as, CD81/SR-B1, TLR3 [11]. These steps are essential to the release of the virus in the cytoplasm of host cells.

Prior to the approval of cyclosporine (CsA) in the United States in 1983, survival rates for liver transplant
recipients were less than 50%. Steroids and azathioprine were the mainstay of therapy to prevent rejection. The development of calcineurin inhibitors (CNIs) revolutionized transplant medicine and after the introduction of cyclosporine survival rates increased to more than 70%. Currently most patients undergoing OLT will receive either one of two such immunosuppressants, either tacrolimus (Tac) or (CsA) [1,9]. Calcineurin (CN) is a calcium-dependent serine/threonine phosphatase that promotes T cell activation, by the expression of several cytokine genes, the most important interleukin-2 (IL-2), but also interleukin-4 (IL-4). Cyclosporine A was first isolated from the fungus Tolypocladium inflatum and has powerful immunosuppressive effects. Cyclosporins are cyclic peptides that have broad spectrum of biological activity. CsA binds to cyclophilin (Cyp) to form a binary complex that binds CN and inhibits its activity. Tacrolimus (Tac) is an immunosuppressant that attaches to FK binding proteins which then decreases calcineurin activity ultimately leading to immunosuppression [5]. Tac is thought to be more potent than CsA, resulting in doses that are typically 25- to 50-fold less than CsA [12]. Although more potent, Tac-based and CsA-based regimen were thought to be comparable in terms of patient and graft survival in both OLT and living donor liver transplant [13,14]. Thus, both CsA-Cyp and Tac-FK complexes are both potent inhibitors of CN activity that ultimately lead to immunosuppression via suppression of IL-2 synthesis.

Cyclosporine or tacrolimus are the most commonly used CNI’s used after liver transplantation. The different molecular processes and properties leading to calcineurin inhibition may thus provide a theoretical basis for which each CNI may interact with viral components. Furthermore, the difference between Tac and CsA potency has influenced how they were used in the past. Indeed, Tac was previously preferred in patients with HCV infection due to its higher potency, thus further limiting the use of steroids [6]. This view started to change, however, due to early reports suggesting acceleration of HCV recurrence in Tac-based regimens. However, both CNI’s are toxic to pancreatic islet cells, but Tac is more toxic possibly as a result of increased concentration in islet cells of tacrolimus compared to cyclosporine. Furthermore, HCV is a risk factor for diabetes. Although immunosuppression accelerates chronic hepatitis C, and viral loads are known to increase with corticosteroid treatment [2,6,12,15]. The best regimen that balances the permissive effects of immunosuppressants versus graft survival has yet to be determined.

**In Vitro Data of Cyclosporine and Hepatitis C**

Studies suggest that CsA inhibits HCV viral replication and increases endogenous interferon. In vitro data using the HCV subgenomic replicon system demonstrate an inhibitory effect of CsA on HCV protein expression and replication [8-11]. Cyclophilins interact with the NS5 protein complex. For example cyclophilin B interacts with the NS5B polymerase to directly stimulate RNA binding activity. By binding to cyclophilins, CsA may disrupt the CycB-NS5 complex formation and thus HCV replication in vitro [16]. However, the cyclophilins contributions and roles in the HCV life cycle remain to be determined.

Due the central role IFN-α plays in the control of HCV infection and the ability of HCV to suppress IFN-α production, it has been hypothesized that CsA may have an effect on IFN expression. Indeed, some authors have demonstrated that HCV replication was suppressed by CsA with associated elevated levels of endogenous IFN-α. More specifically, Liu and colleagues report that CsA treatment of HCV JFH-1-infected Huh7 hepatocytes restored endogenous levels of IFN-α and also enhanced its expression but others found no evidence that either CsA or Tac interferes with IFN-α-induced gene expression or antiviral activity [5,15]. In addition to effects of viral replication, CsA decreases the activation of intrahepatic fibroblasts induced by IL-4, thus reducing fibrosis in recurrent HCV [17]. It is thought that this may account for the observation in which Tac therapy was associated with higher probability of developing cirrhosis [11,17]. CsA may enhance the production of TGF beta but studies have not consistently shown this.

By binding to host proteins, CsA is thought to mediate its antiviral effects. Other investigators further report that CypA is the isofrom that serves as the essential cofactor for HCV replication and that the interaction of cyclophilins with NS5A seems to be the most critical [18]. Due to the high mutation rate of the HCV genome and relative stability of host cellular factors, new avenues for HCV therapy has involved targeting host proteins to further reduce drug-resistant HCV strains [16]. Currently, at least three CsA derivatives have shown anti-HCV effects by binding cyclophilins, namely: DEBIO-025, SCY635, and NIM811. Unlike CsA, however, these compounds possess a structural modification that interferes with CN but not Cyp binding [2]. Thus, these compounds are thought to be devoid of immunosuppressive activity that antagonize immune mechanisms for attaining SVR and may further simplify analysis of clinical trials by eliminating the permissive effects of immunosuppression on HCV replication.

Identifying the optimal immunosuppressive regimen is critical because recurrent HCV is a significant problem after liver transplant and treatment of post-transplant chronic HCV patients is plagued by lower rates of SVR (about 20%), side effects and the potential for increased
graft loss with use of alpha interferon (IFN) [5,15]. Thus, there remains potential for development of novel therapy for HCV for post-liver transplant recipients.

3. Clinical Studies of Calcineurin Inhibitors and Hepatitis C

Initial comparisons by Wiesner, et al. suggest that long-term outcome was superior for Tac-based therapy in patients infected with HCV when compared to a regimen based on CsA [12,19] (Table 1). The secondary immunosuppressants used in the trial, however, varied considerably and was not controlled for in each arm. Indeed, steroid doses of the Tac-based arm were rapidly tapered to 5 mg/day by 3 months, and no other immunosuppressives

<table>
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<tr>
<th>Study (ref)</th>
<th>Design</th>
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<tr>
<td>Castells LL et al. [8]</td>
<td>Multicenter retrospective study in 14 Spanish transplant centers comparing SVR and relapse rates of patients treated with PEG-IFN and Ribavirin while either on CsA or Tac based regimen</td>
<td>410</td>
<td>• CsA arm achieved 48% SVR compared to 37% in Tac arm (p = 0.37) mainly due to the lower relapse rate in CsA group</td>
<td>• Study is retrospective and any inherent biases may not have been accounted for</td>
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<td>Berenguer M et al. [12]</td>
<td>Randomized single-center prospective study comparing Tac vs. CsA based regimen in HCV positive recipients undergoing OLT. Unless otherwise severe, treatment with PEG-IFN and Ribavirin was implemented after the 1st yr</td>
<td>253</td>
<td>• Rates of severe HCV, bridging fibrosis or cirrhosis, and cholestasis-like hepatitis in the first year were not significant between the two arms • Patient survival was also found to be similar at 1, 3, 5, and 7 years post OLT. • SVR were not significantly different between the two arms</td>
<td>• Relatively small sample size with a calculated statistical power of 6% • CsA trough levels (ng/mL): 1st mo: 250 - 350 2nd - 3rd mo: 150 - 250 4th m - 1yr: 100 - 150 &gt;1yr: 100 • Tac trough levels (ng/mL): 1st mo: 10 - 15, 2nd - 3rd mos: 5 - 10 &gt;3 mos: 3 - 10</td>
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<td>Flisiak et al. [22]</td>
<td>Multi-center, randomized, double-blind, placebo-controlled trial designed to determine whether DEBIO-025 in combination with PEG-IFN-α2a is safe and efficacious at dosages of 200 mg, 600 mg, and 1000 mg. No CsA or Tac arm</td>
<td>90</td>
<td>• Dose of 600 and 1,000 mg/day combined with PEG-IFN-α2a induced a significant drop in HCV viral load in genotypes 1 &amp; 4 when compared to those treated with PEG-IFN-α2a alone</td>
<td>• Small sample size since the study is in Phase II clinical trial • It indirectly supports the in vitro data suggesting that CsA have antiviral effects. However, the study does not completely rule out unforeseen mechanisms that may have arisen due to structural modification of the parent compound • Although PEG-IFN and Ribavirin was the standard therapy at the time, Ribavirin was not added to the dosing scheme to simply analysis • Secondary immunosuppressants were not controlled for each arm • CsA group received an overall greater dose of steroids • Maintenance trough levels declined gradually; 5-yr post transplant 80% of individuals have the following trough levels (ng/mL): CsA: 75 - 328 Tac: 2.5 - 14.8</td>
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<td>Wiesner [19]</td>
<td>Multicenter RCT comparing the efficacy and safety of Tac and CsA in primary OLT recipients</td>
<td>529 113 with HCV</td>
<td>• When stratified by pre-OLT HCV exposure, 5 yr patient survival rate in Tac arm was significantly longer than CsA arm (p = 0.041) • Re-OLT for recurrent HCV slightly lower in Tac arm (n = 3; 2.6%) compared to CsA arm (n = 7; 3.9%) • Median time to recurrent HCV requiring re-OLT is earlier in Tac (142 ± 34 days) than in CsA (787 ± 805 days)</td>
<td>• Secondary immunosuppressants were not controlled for each arm • CsA group received an overall greater dose of steroids • Maintenance trough levels declined gradually; 5-yr post transplant 80% of individuals have the following trough levels (ng/mL): CsA: 75 - 328 Tac: 2.5 - 14.8</td>
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<td>Ghibrial et al. [20]</td>
<td>Retrospective analysis that compared effect of CsA vs. Tac based regimen after a primary OLT</td>
<td>340 total patients in both arms, however, only 10 patients were re-transplanted due to recurrent HCV.</td>
<td>• Subgroup analysis reported is too small to draw any firm conclusions • CsA and Tac Trough levels not reported</td>
<td>• Subgroup analysis reported is too small to draw any firm conclusions</td>
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administered. The CsA arm, on the other hand, received greater doses of steroids. Thus, the observed differences may not have been related to the type of CN-inhibitor, but rather due to drug dose, higher steroid doses in the CsA arms, and/or overall levels of immunosuppression [6,12]. Other early studies not primarily designed to compare HCV recurrence rates between CsA and Tac, such as those reported by Ghobrial et al. and a systematic review, however, suggested cyclosporine was associated with lower rates of HCV recurrence and better responses to antiviral therapy (20,21) (Table 1). The median time of HCV recurrence was significantly earlier in Tac-based immunosuppression than those based on CsA. A follow up study by the latter group, however, showed that the histological outcome of both groups after 7 years were similar [6]. More recently, a multicenter retrospective analysis suggests that CsA based therapy is associated with higher response rates to antiviral therapy in subjects on CsA based regimens compared to tacrolimus based therapy [8]. In this study, sustained virological response (SVR) and relapse rates among 410 patients who were treated with Peg-Interferon and ribavirin post-OLT were assessed. The study showed that patients in the CsA arm achieved a 48% SVR and only 37% in the Tac arm (p = 0.037), which was largely due to the lower relapse rate with CsA group (p < 0.001). Multivariate analysis demonstrated that there were no significant differences in factors associated with SVR between both arms. A meta-analysis of published data between 1993 and 2005 found that patient and graft survivals in post-OLT HCV positive patients were independent of the CN inhibitor used [21]. In an attempt to address this question, Berenguer et al. designed a single-center prospective study to assess the recurrence of HCV based on the initial CNI prescribed [11]. The authors reported the rate of severe hepatitis C, those with bridging fibrosis or cirrhosis and those that developed cholestasis-like hepatitis in the first-year, were similar in both CsA and Tac arm. Thus, it is not surprising that SVR is not significantly different between the two CNIs with subsequent treatment with PEG-IFN and Ribavirin. Furthermore, patient survival at 1, 3, 5, and 7 years were similar in the CsA and Tac based regimen. The major limitation of this study, however, is the relatively small sample size (n = 253). Data from retrospective studies provide the rationale for performing larger well-designed randomized prospective studies that compare hepatitis C recurrence rates after liver transplant between tacrolimus and cyclosporine based regimens [3].

4. Future Directions

The approved protease inhibitors for genotype 1 patients in the US and Europe, telaprevir and boceprevir, used in combination with peginterferon and ribavirin have substantially improved sustained viral response rates in the nontransplant population. These agents significantly increase tacrolimus and cyclosporine levels and ongoing clinical trials will study pharmacokinetics, drug interactions, and efficacy. DEBIO-025 is a cyclophilin under development without immunosuppressant properties but has in vitro and in vivo anti-HCV properties [22]. Flisiak et al. designed a 4-week, multicenter, randomized, double-blind, placebo controlled phase II trial to determine the antiviral properties of DEBIO-025 with PEG-IFNα-2a in nontransplant, treatment naïve patients with chronic HCV. The effect of DEBIO-025 at three different doses in combination with pegylated interferon α2a on HCV viral load of varying doses of DEBIO-025 in combination with 180 μg/week PEG IFN-α2a was compared to PEG IFN-α2a and ribavirin. The investigators found that a dose of 600 and 1000 mg/day when combined with PEG INF-α2a induced a statistically significant additive drop in HCV viral load in genotype 1 and 4 compared to PEG IFN-α2a. Furthermore, the safety profile of the combination therapy is similar to that observed in the standard of care arm.

Although in vitro data suggest the should be a benefit of cyclosporine over tacrolimus in hepatitis C liver transplant recipients, clinical studies have not shown that hepatitis C recurrence is significantly lower in recipients treated with cyclosporine. Furthermore, the development of DAA’s has changed the efficacy and side effect profile of treatment and the impact of these new classes of agents and interaction with CNI’s needs further study. Thus, cyclosporine can not be routinely recommended for hepatitis C recipients. Another exciting class of agents are cyclophilin inhibitors, and if proven to be safe and effective, CsA-derivatives may prove to be an important part of the regimen for liver transplant recipients infected with HCV.

REFERENCES


