Relation of pretreatment sequence diversity in NS5A region of HCV genotype 1 with immune response between pegylated-INF/ribavirin therapy outcomes

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SUMMARY. Hepatitis C virus (HCV) infection frequently persists despite substantial virus-specific immune responses and the combination of pegylated interferon (INF)-α and ribavirin therapy. Major histocompatibility complex class I restricted CD8+ T cells are responsible for the control of viraemia in HCV infection, and several studies suggest protection against viral infection associated with specific HLAs. The reason for low rates of sustained viral response (SVR) in HCV patients remains unknown. Escape mutations in response to cytotoxic T lymphocyte are widely described; however, its influence in the treatment outcome is ill understood. Here, we investigate the differences in CD8 epitopes frequencies from the Los Alamos database between groups of patients that showed distinct response to pegylated α-INF with ribavirin therapy and test evidence of natural selection on the virus in those who failed treatment, using five maximum likelihood evolutionary models from PAML package. The group of sustained virological responders showed three epitopes with frequencies higher than Non-responders group, all had statistical support, and we observed evidence of selection pressure in the last group. No escape mutation was observed. Interestingly, the epitope VLSDFK TWL was 100% conserved in SVR group. These results suggest that the response to treatment can be explained by the increase in immune pressure, induced by interferon therapy, and the presence of those epitopes may represent an important factor in determining the outcome of therapy.

Keywords: genotype 1, hepatitis C virus, hepatitis, immune response, NS5A.

INTRODUCTION

Hepatitis C Virus (HCV), a positive-strand RNA virus of the Flaviviridae family, is a major cause of chronic liver disease. An estimated 130 million individuals are infected worldwide [1], and about 70% of infections become chronic, linked with high rate of mutations in the HCV genome that leads to the occurrence and selection of escape variants, which contribute with persistence of infection and chronic cases that are often associated with cirrhosis and hepatocellular carcinoma [2–4].

The NS5A is a nonstructural HCV protein associated with several functions such as transcriptional repressor and associated via the interferon (INF)-sensitive determining region (ISDR), with influence on INF activity.[5]. The ISDR inhibits the cellular RNA-activate protein kinase R (PKR), which is an effector of INF antiviral activity. The NS5A might function as viral transactivator [6]. The NS5A is the major nonstructural protein associated with INF resistance; however, no specific explanation about which variables that influence the therapy outcome or how is the influence of immune pressure on this region was well described.

Furthermore, only about 30–40% of the patients treated with the current therapy with α-INF present sustained virological response (SVR), and the reasons why therapy is effective in only about half of cases are unknown [7]. The cytokine is involved in innate immune responses against
viral infections and Th1 polarization [8,9]. In addition, it was reported that co-administration of INF-α with HIV-1 DNA vaccine increases cellular and humoral immune response [10].

There is growing evidence that INF-α therapy is associated with the increase in HCV-specific T-cell immunity in chronic hepatitis C [11,12], nevertheless, the influence of immune system on the outcome of infection has not been understood.

Several studies suggest an important role of the adaptive T-cell response for control of acute hepatitis C virus (HCV) infection [13], and the association between the expression of particular major histocompatibility complex (MHC) class I alleles and the natural course of HCV infection has been described, suggesting an important role of virus infection control [14]. Moreover, an strong CD8+ T-cell immunity in acute resolving hepatitis C is matched by strong and sustained CD4+ T-cell proliferation to multiple recombinant structural and nonstructural viral proteins [15]. This virus-specific CD8+ T-cell response in acute resolving HCV infection is also vigorous and multispecific, targeting up to eight to twelve epitopes [16,17].

In studies on HCV infection, it was found strong association between viral persistence and escape mutations [18]. However, the determinants of viral escape, as well as its relative contribution to T-cell failure and HCV persistence, remain not well characterized.

The aim of this study is to evaluate the influence of immune response on entire NS5A region of viral quasispecies before the INF therapy in pretreatment patients infected with genotype 1 virus and its consequences on viral protein evolution regarding the presence of mapped epitopes.

MATERIALS AND METHODS

Patients
For this study, eleven patients infected chronically with HCV RNA genotypes 1a or 1b from the Hepatology Department of the São José do Rio Preto University Hospital were included. All patient characteristics, therapy protocols and Ethics Committee were described by Jardim et al. [19]. In this analysis, the patients were classified into two groups, according to their response to therapy: three patients (2–4) were classified as SVR. Eight patients (5–12) were non-responders (NR) to the therapy. SVR was defined as absence of HCV RNA in plasma by qualitative PCR 6 months after the end of the therapy. NR was defined as continued presence of HCV RNA in plasma during treatment and 6 months after and HCV RNA-negative at the end of the therapy but experienced a rebound in HCV viraemia afterwards. The patient 1 was excluded from analysis because the NS5A region was not fully sequenced. The SVR and NR groups have similar age means (39.6 and 39.7, respectively); however, most of the patients were men (one woman and two men in SVR group and one woman and seven men in NR group). The SVR group has two patients infected with 1a genotype virus and one with 1b. The NR group has four 1a virus genotype-infected patients and four infected with 1b genotype virus. The Nucleotide sequence data reported in this article are available in GenBank under the accession numbers EU309511 to EU309673.

Cloning and sequencing HCV genome
For each patient, 15 clones of RT-PCR-amplified fragment of the virus were sequenced, except for patients 3 and 5, who have lower numbers of sequenced clones (14 clones). The sequence length was 1344 nucleotides, representing all NS5A regions. Thus, 163 full-length NS5A sequences were analysed in this work. All protocols of RNA extraction, RT-PCR, NS5A amplification, cloning and sequencing the HCV RNA from patients are described elsewhere [19].

Sequence analysis
Sequences were aligned with Muscle. [20] and the alignments were visually edited using GeneDoc [21]. The epitope mapping was performed using the CD8+ epitopes from HCV Immunology database [22]. For this, all the 21 NS5A epitopes were used to evaluate the frequency of each epitope between two groups. To determine potential positive selection, we used two approaches. A maximum likelihood approach (ML) using codeml program of PAML4 package [23] was employed for the entire NS5A region. For each epitope with statistical difference between both groups, we estimate the dN/dS ratio, dS and dN using the yn00 program of PAML4 package.

The dN/dS, dN and dS estimation
For dN/dS ratio, dN and dS epitope analysis, all values were measured using the method of Yang and Nielsen [24]. For each epitope, the nucleotide sequence alignments were performed and used for the analysis. To calculate the dN/dS values for all epitope quasispecies between two groups, all pairwise comparisons for an epitope were performed, and values were averaged overall.

Maximum likelihood analysis
Maximum likelihood topologies were estimated using PhyML v2.44 [25] and used in ML analysis implemented in codeml. The site models were used to detect positive selection without averaging the dN/dS ratio throughout the phylogenetic tree. The M0 (one ratio), M1a (neutral), M2a (selection), M7 (beta) and M8 (beta&K) codon-based ML substitution models were used to measure the dN/dS ratio and to test the evidence of positive selection by comparing the likelihood values of the different nested models using the. The Bayes Empirical Bayes approach was performed if the LRT suggests
evidence of positive selection to determine which individual codon is subjected to positive selection.

**Statistic analysis**

Statistical difference in the parameters was determined between different patients’ groups by Mann–Whitney non-parametric test for numerical variables. The comparisons of presence/absence of epitopes among the patient groups were made using the G statistics. We also applied the conservative Bonferroni’s correction for multiple comparisons.

The LRT statistics ($2\Delta l$) was performed and compared against $\chi^2$, with critical values to be 5.99 and 9.21 at 5% and 1% significance levels. The first pair M1a (neutral) vs M2a (selection) and M7 (beta) vs M8 (beta&W) were used for verify evidence of positive selection as suggested by Yang [23].

**RESULTS**

**Mapping of CD8+ HCV-specific epitopes in both infected patient groups in NS5A protein**

First, we performed a screening of epitopes from HCV Immunology Database in NS5A translated protein sequence from SVR group ($N = 44$), as well as for protein sequence from the NR group ($N = 119$). Thus, 21 epitopes overall were used for the mapping, representing all from NS5A region, and these results are presented in Table 1.

Three of them have showed no matches in NS5A protein in both SVR and NR groups (epitopes: VLDSFKTWL, LPKLPGVPF and QLSAPSILKTCTANH), and one has no match in SVR group and low frequency in NR group (5.04% of the epitope KFPPALPIW). Five epitopes have demonstrated higher frequency rates in the NR group when compared with SVR group, and eight showed higher rates in SVR group, but without statistical significance.

Overall, the frequency of HCV-specific epitopes was lower in NR group, when compared with SVR group, but this difference was not statistically significant. Four epitopes showed statistical differences with the G test between patient response groups. Three of them have shown higher frequency rate in SVR group (epitopes: VLDSFKTWL, ARGSPPSVASSASQALSAP and SVASSASQSLA). However, the epitope VLDSFKTWL showed 100% frequency compared to 48% in NR group. Only the epitope LTDPSHITA had higher frequency rate in NR group, with 98% compared to 65% in SVR group. All epitopes that showed statistical significance were identified as HLA allele A-binding epitopes (alleles: A1, A2 and both).

**Analysis of selection pressure in patient groups**

Maximum likelihood methods were used to assess amino acid variation and identify targets of positive selection, and all values are presented in Table 2. The mean dN/dS value of NS5A region from SVR, NR and both groups was measured with M0 and showed no statistical support. The comparison of first pair M1a (neutral) and M2a (selection) showed no statistical support in NR and SVR groups; however, the second pair M7 (beta) and M8 (beta&W) analysed with the

### Table 1 Frequencies of epitopes in the HCV quasispecies NS5a protein sequences from the patient groups (Genotype 1 a/b)

<table>
<thead>
<tr>
<th>EPITOPE</th>
<th>Allele</th>
<th>SVR</th>
<th>NR</th>
</tr>
</thead>
<tbody>
<tr>
<td>VLSDFKTWL</td>
<td>A2</td>
<td>100</td>
<td>48.73*</td>
</tr>
<tr>
<td>VLSDFKTWL</td>
<td>A2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>SDFTWLKA</td>
<td>Patr-B*1601</td>
<td>65.90</td>
<td>47.89</td>
</tr>
<tr>
<td>DFKTWLQSRL</td>
<td>NA</td>
<td>34.09</td>
<td>50.42</td>
</tr>
<tr>
<td>LPKLPGVPF</td>
<td>B35</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>MWSSGFPIKAY</td>
<td>Patr-A*0601</td>
<td>63.63</td>
<td>49.57</td>
</tr>
<tr>
<td>LLREVEFSRFR</td>
<td>A2</td>
<td>34.09</td>
<td>49.57</td>
</tr>
<tr>
<td>HEYPVGSOL</td>
<td>B60, H-2k</td>
<td>63.63</td>
<td>48.73</td>
</tr>
<tr>
<td>PCEPEPDAVL</td>
<td>B35</td>
<td>31.81</td>
<td>47.05</td>
</tr>
<tr>
<td>EPEPDAVL</td>
<td>B35</td>
<td>31.81</td>
<td>47.05</td>
</tr>
<tr>
<td>LTDPSHITA</td>
<td>A1</td>
<td>65.90</td>
<td>98.31*</td>
</tr>
<tr>
<td>ARGSPPSVASSASQALSAP</td>
<td>A1/A2</td>
<td>65.90</td>
<td>25.21*</td>
</tr>
<tr>
<td>SVASSASQSLA</td>
<td>A1/A2</td>
<td>65.90</td>
<td>25.21*</td>
</tr>
<tr>
<td>QLSAPSILKTCTANH</td>
<td>NA</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>NHDSPDPAEL</td>
<td>B38</td>
<td>61.36</td>
<td>49.57</td>
</tr>
<tr>
<td>SPDAEIEANL</td>
<td>A2</td>
<td>65.90</td>
<td>49.57</td>
</tr>
<tr>
<td>ELIEANLW</td>
<td>A25</td>
<td>65.90</td>
<td>49.57</td>
</tr>
<tr>
<td>ILDSFDPLV</td>
<td>A2</td>
<td>31.81</td>
<td>10.92</td>
</tr>
<tr>
<td>LDSFDPLVA</td>
<td>Patr-B*0401</td>
<td>65.90</td>
<td>47.89</td>
</tr>
<tr>
<td>REISVPAEL</td>
<td>B60</td>
<td>31.81</td>
<td>49.57</td>
</tr>
<tr>
<td>KFPALPIW</td>
<td>A*2402</td>
<td>0</td>
<td>5.04</td>
</tr>
</tbody>
</table>

NA, not available; SVR, sustained virological response group; NR, non-responders group.

*Represents different frequencies between patient groups with P-value <0.05 (corrected with the Bonferroni’s procedure); All frequencies are expressed in percentage.

### Table 2 Results of selective pressure between the patient groups

<table>
<thead>
<tr>
<th>Treatment response</th>
<th>dN/dS ratio</th>
<th>M2a vs M1a</th>
<th>M8 vs M7</th>
<th>Positive selected AA (position)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SVR</td>
<td>0.09</td>
<td>0.367</td>
<td>4.058</td>
<td>ND</td>
</tr>
<tr>
<td>NR</td>
<td>0.09</td>
<td>0.023</td>
<td>13.376*</td>
<td>L(2130), P(2319)</td>
</tr>
<tr>
<td>Both groups</td>
<td>0.1</td>
<td>0.001</td>
<td>15.49*</td>
<td>L(2130), P(2319)</td>
</tr>
</tbody>
</table>

ND, not detected; dN/dS, nonsynonymous/synonymous substitution ratio rate; AA, aminoacid; SVR, sustained virological response group; NR, Non-responders group.

*Represents statistically significant P-value <0.01, in likelihood ratio test.
LRT statistic suggests that M8 model was selected \((P < 0.01)\), providing evidence of varying selection pressure at individual sites across the NS5A region of NR group and in both groups. As LRT suggested that these groups are under positive selection, the Bayes empirical Bayes (BEB) method was used to calculate the posterior probabilities that each codon is from the site class of positive selection under model M8, thus, indicating that the aminoacid L in the position 2130, according to H77 reference strain, and P in the position 2319 as being positively selected, with a posterior probability >95%.

d\(N/dS\) ratio, \(dS\) and \(dN\) measure of significant epitopes

For the four epitopes that showed statistical differences between two groups, a measure pairwise analysis of \(dN/dS\) ratio, the proportion of synonymous and the proportion of nonsynonymous mutations of epitope region were calculated. The results are shown in Table 3. Three of them have higher \(dN/dS\) ratio in NR group, and one of them showed higher ratio in SVR group, all of them with statistical support. Interestingly, the epitopes showed higher ratios in groups where the frequency of them was lower. In \(dN\) measure, one epitope showed no difference between groups. Two of them showed \(dN\) value higher in NR group, and one had \(dN\) value higher in SVR group, with statistical support. The \(dS\) measure provides higher values in NR group of three epitopes with statistical support. One epitope had no difference in \(dS\) values in both groups.

<table>
<thead>
<tr>
<th>Epitope</th>
<th>Group</th>
<th>(dN/dS)</th>
<th>(dN)</th>
<th>(dS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>VLSDFKTWL</td>
<td>SVR</td>
<td>0.03</td>
<td>0.02</td>
<td>0.76</td>
</tr>
<tr>
<td></td>
<td>NR</td>
<td>0.04*</td>
<td>0.03*</td>
<td>0.91*</td>
</tr>
<tr>
<td>LTDPSHITA</td>
<td>SVR</td>
<td>0.018*</td>
<td>0.026*</td>
<td>0.79</td>
</tr>
<tr>
<td></td>
<td>NR</td>
<td>0.001</td>
<td>0.001</td>
<td>0.86*</td>
</tr>
<tr>
<td>ARGSPPSVASSASQLSAPS</td>
<td>SVR</td>
<td>0.02</td>
<td>0.01</td>
<td>0.29</td>
</tr>
<tr>
<td></td>
<td>NR</td>
<td>0.09*</td>
<td>0.01</td>
<td>0.34*</td>
</tr>
<tr>
<td>SVASSASQLSA</td>
<td>SVR</td>
<td>0.06</td>
<td>0.02</td>
<td>0.25</td>
</tr>
<tr>
<td></td>
<td>NR</td>
<td>0.12*</td>
<td>0.03*</td>
<td>0.25</td>
</tr>
</tbody>
</table>

\(dN/dS\), nonsynonymous/synonymous substitution rate ratio; \(dN\), nonsynonymous substitution rate; \(DS\), synonymous substitution rate; SVR, sustained virological response group; NR, non-responders group. *Represents statistically significant \(P\)-value <0.05, in Mann–Whitney Test comparing SVR and NR groups.

Aminoacid change in epitopes with statistical difference

As shown in Table 4, no viral sequence variation was detected within epitope VLSDFKTWL in SVR group. However, in NR group, it was observed two aminoacid changes in position 3 of epitope (P3) serine was substituted by threonine with 37.8% and substituted by alanine with 12.6%. Sequence analysis of epitope LTDPSHITA has demonstrated aminoacid change in second position (P2), a threonine was substituted by serine in 34% of sequences of the SVR group. The aminoacid changes in NR group were <2%. The epitope ARGSPPSVASSASQLSAPS showed aminoacid change in position 8, where a valine was substituted by leucine with 33.9% in SVR group. Another aminoacid changes occurred in this epitope with 2.2% of frequency. Arginine was substituted by glycine at second position, and serine was substituted by leucine in position 12. In NR group, this epitope presented aminoacid change in position 8, where valine was substituted by leucine with 63% of frequency and valine to methionine with 12.6%. The epitope SVASSASQLS showed aminoacid variation in position 2, where valine was substituted by leucine with 33.9% of frequency in SVR group and 63% in NR group. Valine was substituted by methionine in the same position with 12.6% frequency in NR group. None of the aminoacids changes within epitopes had BEB analysis posterior probability evidence of positive selection.

<table>
<thead>
<tr>
<th>Epitope</th>
<th>SVR</th>
<th>LTDPSHITA</th>
<th>ARGSPPSVASSASQLSAPS</th>
<th>SVASSASQLS</th>
</tr>
</thead>
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<tr>
<td>VLSDFKTWL</td>
<td>. . . .</td>
<td>.S. . . .</td>
<td>. . . . . . . . . .</td>
<td>.L. . . .</td>
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<tr>
<td>LTDPSHITA</td>
<td>. . . .</td>
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<tr>
<td>ARGSPPSVASSASQLSAPS</td>
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<tr>
<td>SVASSASQLS</td>
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</table>

SVR, sustained virological response group; NR, non-responders group.
NS5A region [22] with significant differences between the patient groups, despite the small number of patient in our analysis. The epitopes VLSDFKTLW, ARGPSPSVSASS ASQLSAPS and SVAASSSASQLSA showed higher frequencies in sustained viral response (SVR) group when compared to the NR group. Thus, this data might indicate that those epitopes, specially the VLSDFKTLW, induced by INF, promote immune response against HCV and may drive the treatment outcome. Despite the epitope LTDPHITA, which had shown higher aminiacid variation in SVR compared to NR group, it could be explained as an ineffective response against this epitope in NR group, therefore, this epitope could not have effect in the clearance of HCV.

It is important to note that epitope VLSDFKTLW had 100% of matches in protein sequences of SVR group and 48% in NR group. However, the NR group has two distinct treatment failure patients: complete non-response patients, defined as continued presence of HCV RNA in plasma during treatment and 6 months after the end of the treatment. Patients who were HCV RNA-negative at the end of the therapy but experienced a rebound in HCV viraemia afterwards were classified as end of treatment response (ETR). About 75% of patients (three of them) with sequence variation presented complete nonresponse to the treatment, and just 25% (one patient) showed ETR. This result suggests that the presence of this epitope plays an important role in specific CD8+ response against HCV control, and mutations in this region could affect in the treatment outcome. However, because of our small number of patients, future studies are needed to provide more information about the course of HCV infection and t-cell epitope dynamics during therapy.

We investigated the evidence of positive selection in HCV NS5A sequences of patients using evolutionary models as described in the Methods section. We observed the same value of dN/dS ratio in both groups (dN/dS ratio = 0.09) and a similar value when we analysed them together. All of the three values showed evidence of purifying selection. However, the straightforward use of this ratio is rarely effective to detect positive selection, because the ratio over all sites is often <1 [23]. In the ML analysis of selection models, the results showed no evidence of positive selection in the SVR group, which suggests that immune response and the peg-INF therapy were effective to clear the HCV before the emergence of the escape mutations which improve the viral fitness, leading to the treatment failure. Nevertheless, in the NR group and in both groups together, we observed evidence of positive selection. This finding suggests that the failure of therapy is associated with the NS5A region variation, despite none of the two amino acids were shown to be under positive selection by the Bayesian analysis in the epitopes regions. The evidence of positive selection in regions out of epitopes indicates that the mutations in NS5A protein keep the viral replication in the presence of the drugs and compensate the viral fitness.

Several studies have reported that sequence variation in NS5A region is associated with IFN responsiveness, specifically the IFN sensitivity-determining region. RNA-activated protein kinase (PKR)-binding domain and V3 region [26–29]. Nevertheless, the role of CD8+ T cells in the control of HCV replication during INF therapy has been not well described. Our results show that the presence of epitope VLSDFKTLW is strongly associated with virus clearance during treatment, suggesting that the CD8+ T cells’ specificity to the HCV could increase and promote a strong immune response, being an important factor in the treatment success. Interestingly, this epitope was described in cytoplasmatic retention signal region [30], because this region was not been associated with immune response activity or therapy effectiveness.

In agreement with several other studies, we found evidence of an important role of specific CD8+ T-cells epitopes in control of viraemia and association with viral persistent in chronically evolving patients and escape mutations, even with the small sample size [31–33].

Because escape mutations could impair HLA binding or loss of processing during MHC presenting process, we then assessed the specific mutations in epitopes regions to verify which positions have sequence variation and compare to information about HLA epitopes in the SYFPEITHI database [34]. With the exception of the epitope SVAASSSASQLSA, which showed the mutation valine to leucine at second position of the epitopes, all other mutations were not associated with expected aminiacids in anchor positions or preferred residues. In this epitope, a valine mutates to leucine, which is an expected anchor aminoacid to this position in allele HLA A1. The epitope VLSDFKTLW showed no sequence variation in SVR group; however, in the NR group, the position 3 serine was substituted by threonine with 37.8% and by alanine with 12.6%. These results reinforce the association between the role of CD8+-specific immune response against HCV and the therapy, because those aminoacids are not expected as preferred residues as described in SYFPEITHI database from allele A2, indicating that are escape mutations. T-cell escape mutations resulting in complete abrogation of HLA binding or disruption of processing will not be presented on the surface of the cell and are thus invisible to immune system. It is important to note that, in addition to the hypothesis suggested previously, host factors, such as the genetic restriction of the immune response, T-cell receptor diversity or strong CD4+ T-cell help [35–37], may also contribute to the observed viral escape in NR group.

We decide to focus on the epitope regions and evaluate the dN/dS ratio, dS and dN for each epitope in both groups to verify statistical differences. For three epitopes that showed higher frequencies in SVR group, we found dN/dS ratio lower than NR, suggesting that immune system perform selective pressure on this epitopes. Despite the epitope found LTDPHITA showed dN/dS ratio higher in SVR group.
it could be explained by the higher dS value in NR group, suggesting unspecific immune response in patients from NR group, because this group showed no response to treatment.

In summary, mapping 21 NS5A epitopes in our eleven HCV chronically evolving patients under peg-INF therapy, we could show evidence of at least one T-cell HCV-specific epitope with association on the treatment outcome. In addition, evolutionary models were used to confirm the presence of immune pressure on the virus. Taken together, these results have important implications to understand the HCV/immune system dynamics during the treatment period, indicating that immune system plays an important role in HCV clearance during treatment period.

FINANCIAL DISCLOSURE

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