

Prevalence of hepatitis C virus-resistant association substitutions to direct-acting antiviral agents in treatment-naïve hepatitis C genotype 1b-infected patients in western China

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Background: Direct-acting antivirals (DAAs) against hepatitis C virus (HCV) are potent and highly efficacious. However, resistance-associated substitutions (RASs) relevant to DAAs can impair treatment effectiveness even at baseline. Moreover, the prevalence of baseline RASs in HCV genotype 1b-infected patients in western China is still unclear.

Materials and methods: Direct sequencing of the HCV NS3, NS5A, and NS5B regions was performed in baseline serum samples of 70 DAAs treatment-naïve HCV 1b-infected patients in western China. The sequences were analyzed with MEGA version 5.05 software. Evolutionary patterns of RASs and amino-acid covariance patterns in the *NS3*, *NS5A*, and *NS5B* genes were analyzed by MEGA and Cytoscape (version 3.2.1), respectively.

Results: The presence of at least one RAS in the NS3 region (C16S, T54S, Q80R/L, A87T, R117H, S122G, V132I, V170I) was observed in 85.48% (53 of 62) of patients, RASs in the NS5A region (L28M, R30Q, Q54H, P58S/T, Q62H/R, Y93H) were observed in 42.42% (28 of 66) of patients, and RASs in the NS5B region (N142S, A300T, C316N, A338V, S365A, L392I, M414L, I424V, A442T, V499A, S556G) were observed in 100% (44 of 44) of patients. Evolutionary patterns of RASs and amino-acid covariance patterns for the *NS3*, *NS5A*, and *NS5B* genes are reported.

Conclusion: The prevalence of RASs relevant to DAAs detected in the NS3, NS5A, and NS5B regions of HCV 1b from DAA treatment-naïve patients is high. Therefore, more attention should be paid to RASs associated with DAAs in the upcoming DAA-treatment era in China.

Keywords: hepatitis C virus, unstructured proteins, resistance-associated substitutions, direct-acting antiviral agents.

Introduction

Hepatitis C virus (HCV) infection has greatly increased the burden of global disease.¹ The recent estimates of global disease burden show that the seroprevalence of HCV is 2.5%, equating to >177.5 million infections.² HCV is associated with liver cirrhosis, hepatocellular cancer, and liver failure, which has become an important cause of HCV-related deaths, resulting in 350,000 cases of patient death per year worldwide.^{3,4} As HCV is a RNA virus, its genome is highly variable, and the sequences are classified into seven recognized genotypes (GTs) and a large number of subtypes.⁵ HCV sequences belonging to different GTs show a 30%–35% difference at the nucleotide level.⁵ GT1 is estimated to account for more HCV cases than any other GTs, at 83.4 million cases (46.2%), and more than a third of GT1 infections occur in East Asia.⁶ Additionally, 56.8% of HCV infections in China are of subtype 1b.⁷

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Pegylated-interferon and ribavirin have been the standard of care for HCV-infected patients. However, severe side effects of this treatment lead to about 50% of patients being unable to tolerate therapy, and the sustained virological response rate after this treatment is only approximately 50% for HCV GT1 infections.⁸ Remarkably, direct-acting antivirals (DAAs) with the ability directly to inhibit the HCV unstructured proteins NS3/4A protease, NS5A protein, and NS5B polymerase, which are important for HCV replication in hepatocytes, have resulted in very high sustained virological response rates of more than 90% in Phase II and Phase III studies after short-term treatment.⁹ However, due to the low fidelity of HCV polymerase, high HCV-replication rate, and strong selective pressures on the virus, multiple variants differing by up to a few percent exist even within individual isolates, ie, HCV replicates as a quasispecies.^{10,11} Some quasispecies may carry the baseline resistance-associated substitutions (RASs), which convey varying degrees of resistance to DAAs. Indeed, RASs have been observed in both in vitro and clinical trials.¹² In particular, the clinically relevant baseline RASs that reduce susceptibility to DAAs and play a role in treatment failure in postmarketing studies or clinical practice play a potent role in treatment failure.^{13,14} DAAs have not been approved in China; however, this approval is imminent. Therefore, it is necessary to have an understanding of the prevalence of RASs in treatment-naïve HCV GT 1b-infected patients in western China.

Materials and methods

Patients

Serum samples were obtained from 70 patients who had been chronically infected with HCV 1b and never been treated for HCV with any DAAs. Patients were recruited between 2015 and 2016 from the Second Affiliated Hospital of Chongqing Medical University and were diagnosed according to the 2016 European Association for the Study of the Liver recommendations on treatment of HCV.¹⁵ HCV GT and subtype were determined by reverse-transcription polymerase chain reaction (PCR) and sequencing of the HCV core region. Patients enrolled in the study were not coinfecting with HBV or HIV, and all were from the Chinese Han population. The study was approved by the ethics committee of the Second Affiliated Hospital of Chongqing Medical University, and written informed consent was obtained from each study participant.

Amplification and sequencing of HCV NS3, NS5A, and NS5B genes

Viral RNA was extracted from 140 μ L serum samples using the MiniBest viral RNA extraction kit (Takara, Kusatsu,

Japan). cDNA was synthesized from HCV RNA using a PrimeScript RT reagent kit with genomic DNA eraser (Takara) following the manufacturer's instructions. Cycling conditions were incubation at 42°C for 2 minutes to remove genomic DNA, followed by 37°C for 15 minutes to synthesize cDNA. Full-length HCV NS3, NS5A, and NS5B genes were amplified using PrimeStar HS (Takara) in a nested PCR. Primers are shown in Table S1. PCR products in the first PCR round were obtained using a predenaturation step at 95°C for 5 minutes, followed by 35 cycles of denaturation at 98°C for 10 seconds, annealing at 62°C for 20 seconds, and extension at 72°C for 1 minutes and 20 seconds for the NS3 and NS5A genes and 35 cycles of denaturation at 98°C for 10 seconds, annealing at 56°C for 20 seconds, and extension at 72°C for 2 minutes for the NS5B gene. For all genes, these steps were followed by a final 10-minute extension step at 72°C. PCR volume contained 25 μ L of 2 \times reaction premix, 1 μ L each primer, 5 μ L cDNA template, and nuclease-free H₂O for a final volume of 50 μ L. Three microliters from the first round PCR product was used as a template in the second round of PCR, which was done with a similar reaction volume and conditions as the first round, but with an annealing-temperature increase to 64°C for the NS3 and NS5A genes and to 60°C for the NS5B gene.

Direct sequencing of nested PCR products was performed using an automatic sequencer (ABI Prism 3100; Thermo Fisher Scientific, Waltham, MA, USA). Sequencing primers were the second-round PCR primers for each gene. Correction of the base sequences was performed with reference to direct-sequencing chromatograms. Representative chromatograms from the direct sequencing are shown in Figure S1.

Sequence analysis and phylogeny

Clinically relevant RASs to DAAs were summarized from the literature and are presented in Table S2. The NS3, NS5A, and NS5B nucleotide sequences were aligned with the HCV 1b reference sequence (GenBank accession AJ238799) and analyzed using BioEdit 7.0.9 software. A variant type was indicated by replacement of the consensus amino acid with a novel one, eg, Y93H and Y93N in the NS5A region were described as two variant types.

A phylogenetic tree of each gene was created using the maximum-likelihood statistical method with the Jones–Taylor–Thornton two-parameter model in MEGA version 5.05.¹⁶ A γ -parameter of 0.5 was used to model differences in substitution rates among base sites. The reliability of the cluster descending from that node was assessed by a bootstrap analysis with 1,000 replicates.

Structural bioinformatic modeling and amino-acid covariance analysis

Protein-homology modeling was performed with Swiss-Model (<https://swissmodel.expasy.org>), with an identity of more than 90% used to model the X-ray crystal structures of HCV NS3 protease, NS5A, protein and NS5B polymerase for GT 1b. Structural editing and analysis were done using PyMol (Schrödinger, New York, NY, USA). The three structures were loaded into PyMol to analyze the cartoon structures, the stick structures of RASs, and the drug-binding pocket.

Sequences consisting of the *NS3*, *NS5A*, and *NS5B* genes were aligned and translated into amino acid sequences using BioEdit 7.09. Then, all possible amino-acid covariant pairs were determined by employing the observed-minus-expected-squared (OMES) algorithm, which can exclude invariant positions and spurious correlations generated by random differences, as described in other work.¹⁷ To identify the covarying pairs, we calculated an OMES score for every possible pair. An OMES score of 0.5 was used as the cutoff for selecting covariant amino-acid pairs for analyses.¹⁸ Finally, amino-acid covariant networks were visualized by Cytoscape (version 3.2.1).¹⁹

Statistical analysis

Concentration trends of data were expressed as medians (minimum–maximum). Categorical variables were summarized as numbers. Categorical variables were compared between groups using the χ^2 test. Continuous data were compared between groups using Student's *t*-test. $P < 0.05$ was considered statistically significant. Statistical analyses were performed using SPSS 17 software (SPSS, Chicago, IL, USA).

Results

Characteristics of patients

For all 70 patients, the *NS3*, *NS5A*, and *NS5B* genes were successfully amplified and sequenced in 62, 66, and 44 patients, respectively. In total, 66 patients were included for further analysis in this study. The clinical and virological characteristics of these patients are shown in Table 1.

Prevalence of RASs relevant to NS3 protease inhibitors

In total, RASs conferring resistance to NS3 protease inhibitors (PIs) were detected in 53 of 62 (85.48%) sequences, as shown in Figure 1A. Among these, S122G (resistance to simeprevir/asunaprevir) was the most common, observed in 85.48% of sequences, followed by V170I-associated

Table 1 Baseline characteristics of patients with HCV 1b infection in western China

Characteristics	Values
Age (years), median (range)	47 (18–78)
Male (female)	36 (30)
Alb (g/L), median (range)	44.4 (16.8–54.9)
ALT (IU/L), median (range)	62 (5–230)
AST (IU/L), median (range)	49.5 (14–586)
γ -GGT (IU/L), median (range)	53 (9–1,955)
Total bilirubin (mg/dL), median (range)	0.72 (0.37–3.52)
Direct bilirubin (mg/dL), median (range)	0.34 (0.16–2.68)
Plt count ($\times 10^9/L$), median (range)	174 (56–315)
AST/Plt, median (range)	0.31 (0.07–7.23)
Hemoglobin (g/dL), median (range)	149 (77–178)
HCV RNA (\log_{10} IU/mL), median (range)	6.39 (3.00–7.73)

Abbreviations: HCV, hepatitis C virus; Alb, albumin; Plt, platelet.

resistance to telaprevir (25.81%) and T54S-associated resistance to telaprevir and boceprevir (14.52%). The other RASs were observed at frequencies lower than 10%, such as Q80R/L (resistance to simeprevir, faldaprevir, and asunaprevir), V132I and R117H (resistance to boceprevir), C16S (resistance to telaprevir), and A87T (resistance to telaprevir/boceprevir). No RASs showing high-level resistance to DAAs were detected. It is noteworthy that R155, V156, and D168, which are important variants resistant to PIs, were not detected in this study. In addition, some substitutions like Y56F, R123K, and S174A were also detected in this study.

Moreover, among the 62 patients, 25 (40.32%) had a mixture of variants that contained multiple NS3-resistance variants. In detail, 18 (29.03%) patients carried two RAS mixtures, such as V170I + S122G or T54S + S122G, and five (8.06%) patients carried three RAS mixtures, such as Q80R + T54S + S122G or Q80R + S122G + V170I. Importantly, two mixtures of four RASs (Q80R + T54S + V170I + S122G and C16S + R117H + V170I + S122G) were also each detected in one patient.

Prevalence of RASs relevant to NS5A inhibitors

In total, these RASs were detected in 36 of 66 (54.55%) sequences, as shown in Figure 1B. In detail, the most commonly observed RAS was Q54H (resistance to daclatasvir, 42.42%). The next-most common RASs, P58S/T (resistance to daclatasvir) and Q62H/R (resistance to daclatasvir), were detected in 6.06% and 9.09% of sequences, respectively. Lastly, L28M (resistance to daclatasvir/ombitasvir) and R30Q (resistance to daclatasvir) were identified in only one patient each. Notably, the key RAS Y93H/N, which is associated with resistance to daclatasvir/ledipasvir/ombitasvir,

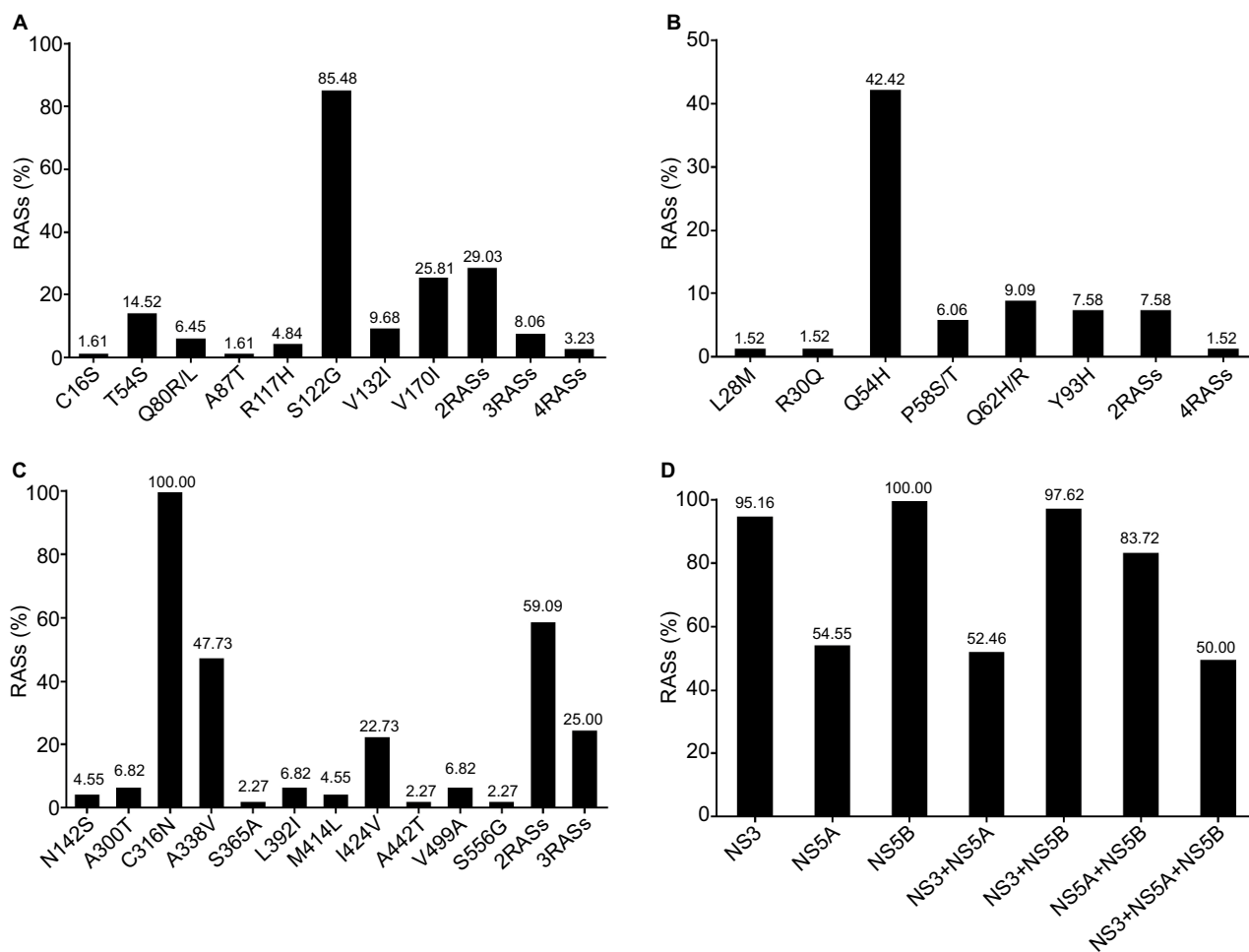


Figure 1 Prevalence of clinically relevant RASs in various regions against different DAA classes.

Notes: Frequency of clinically relevant RASs in the NS3 region (A), NS5A region (B), and NS5B region (C). Total frequency of clinically relevant RASs against different DAA classes (D).

Abbreviations: RASs, resistance-associated substitutions; DAA, direct-acting antivirals.

was detected in 7.58% of sequences and associated with a high level of resistance. But another key RAS, L31V/M, associated with resistance to daclatasvir/ledipasvir, was not observed in this study. In addition, five different mixtures of RASs were observed (Q54H + P58S, Q54H + Q62H, Q62H + Y93H, Q54H + Y93H, and R30Q + Q54H + P58S + Q62R) in six patients.

Prevalence of RASs relevant to NS5B polymerase inhibitors

The prevalence of RASs associated with resistance to NS5B nucleotide analogue inhibitors (NIs) and non-NIs (NNIs) was also differentially distributed, as shown in Figure 1C. Among RASs relevant to NIs, key RASs for sofosbuvir and mericitabine correlate with variants at positions S282 and L159. In this study, no sequence containing L159F and S282T/R was found. However, the prevalence values of

N142S (resistance to sofosbuvir) and A300T (resistance to R7128) were 4.55% and 6.82%, respectively. Compared to RASs relevant to NIs, RASs associated with resistance to NNIs were more commonly observed. Remarkably, C316N (resistance to dasabuvir) was found in all sequences in this study. In addition, A338V (resistance to beclabuvir) and I424V (resistance to beclabuvir) occurred in 47.73% and 22.73% of sequences, respectively. S365A (resistance to nesbuvir), L392I (resistance to beclabuvir), and A442T (resistance to beclabuvir) represented a minority of sequences, with frequencies of 2.27%, 6.82%, and 2.27%, respectively. Notably, the key RASs M414L (resistance to dasabuvir) and S556G (resistance to dasabuvir) were present in 4.55% and 2.27% of sequences, respectively. The RAS V499A (resistance to beclabuvir) was presented in 6.82% of sequences. There were also some substitutions detected, such as S368F, A421E, and C451Y. In addition, 37 of 44 (84.09%) sequences

contained more than one RAS: 26 (59.09%) patients had two RAS mixtures, such as C316N + A338V, C316N + I424V, and C316N + A300T. Eleven (25%) patients had mixtures of three or more RASs, such as C316N + S556G + A338V and C316N + S556N + V499A.

Prevalence of multiple RASs relevant to different DAA classes

Overall, the frequency of multiple RASs associated with resistance to different DAA classes was high, and varied for different RAS combinations (52.46%–97.62%, Figure 1D). The highest prevalence of multiple RASs was in the NS3 + NS5B regions, followed by the NS5A + NS5B, NS3 + NS5B, and NS3 + NS5A + NS5B regions. C316N was observed in all NS5B sequences, which may contribute to its high prevalence. We analyzed the results again after excluding C316N. Similar results at a lower frequency were observed, as presented in Figure S2.

Evolutionary patterns of RASs in NS3, NS5A, and NS5B genes

To study whether RASs occurring at baseline were correlated with ongoing random variation and/or some specific evolutionary patterns in transmission clusters, phylogenetic trees of the NS3, NS5A, and NS5B genes were created, as described in the Materials and methods section. The results showed that most baseline RASs in the NS3 and NS5B genes seemed to

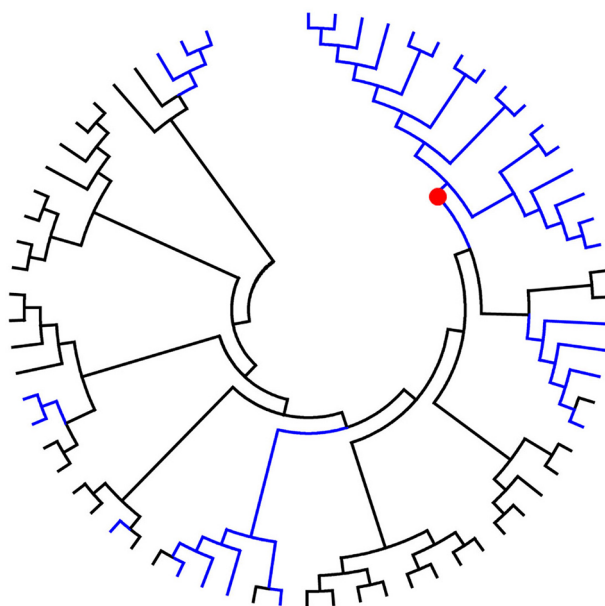


Figure 2 Phylogenetic tree for the hepatitis C 1b NS5A gene.
Notes: Black lines represent unresistant lineages; blue lines represent lineages carrying the resistance mutation; red dots represent the emergence of concentration-trend clades of unresistant isolates.

occur randomly, with no special evolutionary pattern occurring, as observed in Figure S3. However, a clade without RASs in the NS5A gene showed a prominent concentration trend, as shown in Figure 2. Therefore, clinical characteristic analyses were performed in patients with and without RASs in the NS5A gene to investigate the possible reason for this phenomenon. However, there were no significant differences between the two groups, as shown in Table S3.

Structural bioinformatic modeling and amino-acid covariance patterns of NS3, NS5A, and NS5B genes

In order to investigate the structures of the different viral unstructured proteins and reveal the mutations and drug-binding pockets, protein-structure bioinformatic modeling was performed on the HCV NS3, NS5A, and NS5B proteins. The HCV NS3 protease crystal structure and the stick structures of such RASs as R117L, Q80R, and S122G are shown in Figure 3A. The NS5A protein-crystal structure and the stick structures of RASs Q54H, Q62H, Y93H, and P58S are presented in Figure 3B. The NS5B polymerase-crystal structure and stick structures of RASs A338V, A300T, N142S, C316N, and M414L are shown in Figure 3C. Drug-binding pockets in different viral proteins are also shown in Figure 3.

Next, to understand the covariance relationship among the different regions, a covariance-network analysis was undertaken. As shown in Figure 4A, there were a considerable number of amino-acid covariance positions (nodes) and interacting lines (edges), constituting a huge covariance network. These nodes included both RASs and non-RASs. Most of them were found in the NS5B region, such as R120, M414, and V322. Further analysis focused on the RAS-covariance relationships only, and a similar but smaller network was observed, as shown in Figure 4B. RASs at positions R117 in the NS3 region, R30 in the NS5A region, and N142, L392, M414, I424, and V499 in the NS5B region interacted with one another.

Discussion

In this study, clinically relevant RASs associated with resistance to DAAs were frequently observed in treatment-naïve HCV 1b-infected patients in western China. RASs in the NS5B region were the most frequently observed (the C316N variant was detected in 100% of sequences), followed by those in the NS3 region and NS5A region (85.48% and 42.42%, respectively).

In the NS3 region, S122G, which has been confirmed to provide a low level of resistance, was the most commonly

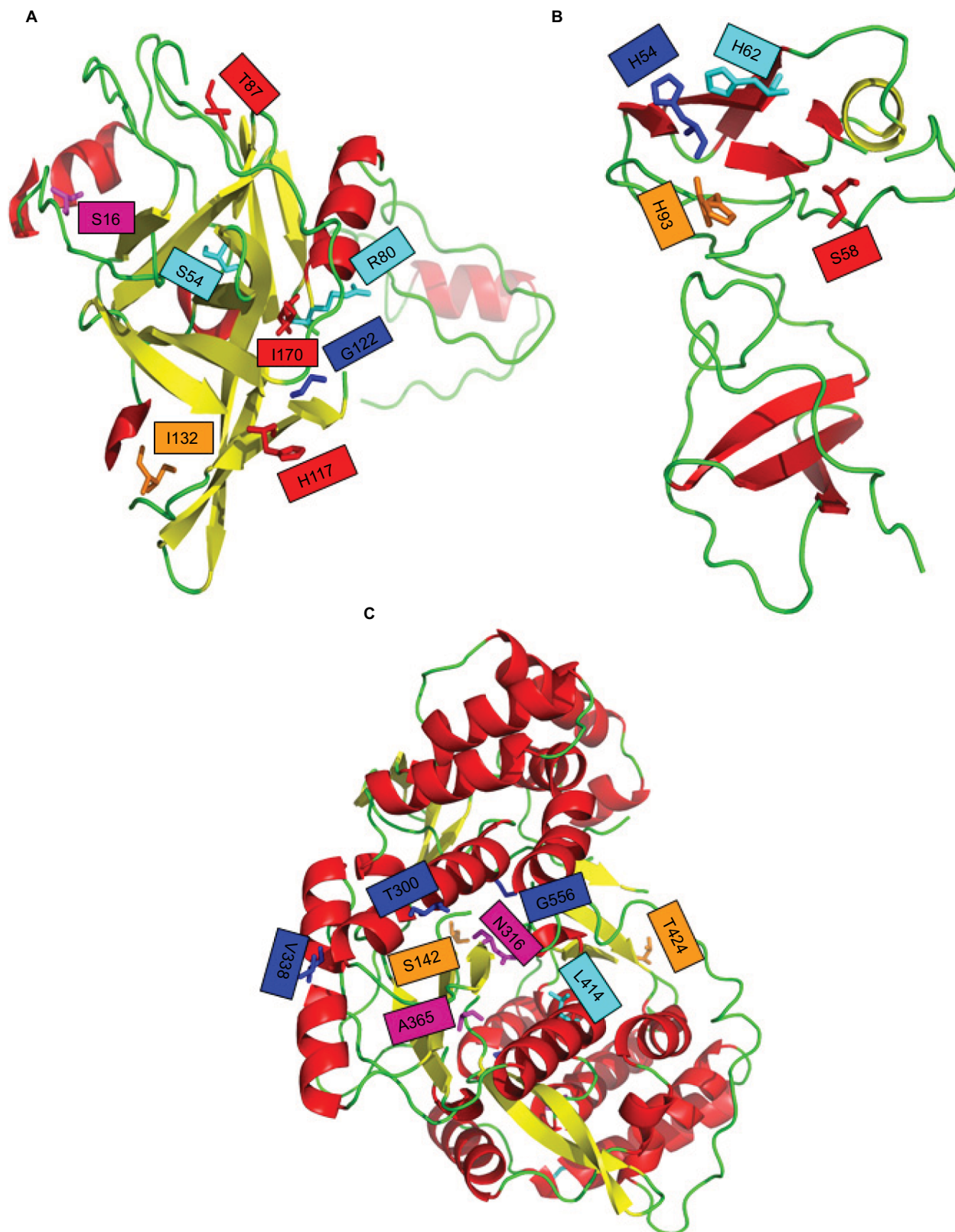


Figure 3 Structures and drug-binding pockets in different HCV unstructured proteins.

Notes: Crystal structures and stick views of RASs in the drug-binding pockets of the HCV NS3 protease (A), NS5A protein (B), and NS5B polymerase (C). Color of crystal structures determined by secondary structure of proteins, with α -helices colored red, β -sheets colored yellow, random coils and other structures colored green for NS3 and NS5B, α -helices colored yellow, and β -sheets colored red in NS5A. The stick structures of RASs are identified by color and amino acid sites.

Abbreviations: HCV, hepatitis C virus; RASs, resistance-associated substitutions.

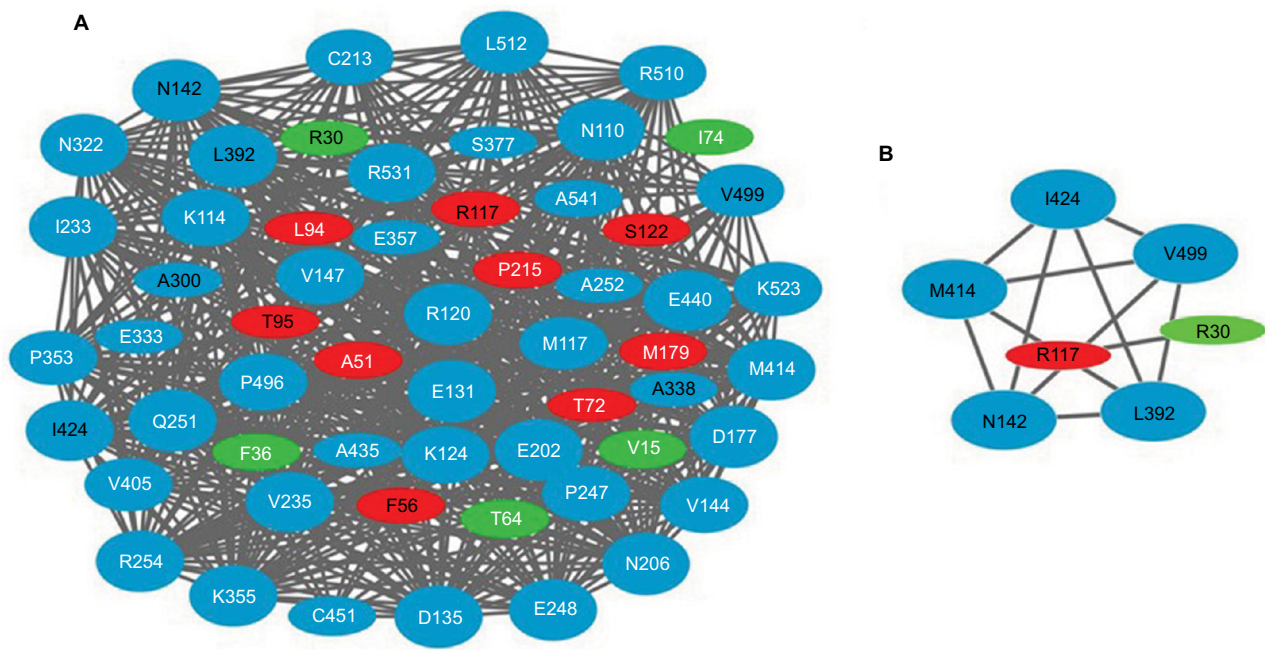


Figure 4 Amino-acid covariance networks for NS3, NS5A, and NS5B sequences in HCV 1b.

Notes: Amino-acid covariance networks for all variant positions (**A**) and clinically relevant resistance-associated substitutions (RASs) (**B**). The size of the nodes is determined by the number of edges that they contact. Red nodes are within the NS3 region, green nodes in the NS5A region, and blue nodes in the NS5B region. White font indicates non-RASs, and black font indicates RASs. Numbered amino acid residue position is provided relative to the first amino acid of the NS3, NS5A, or NS5B region.

observed RAS. S122G is a major RAS relevant to simeprevir and asunaprevir. The occurrence of this RAS in this study was much higher than that observed in an open-label trial in Japan (85.48% vs 36.7%).²⁰ The higher number of patients included in their study may be a reason for this discrepancy. Another key RAS, Q80K, was observed in none of the 62 HCV 1b sequences, while 6.45% of sequences were detected to contain Q80R/L, which is commonly observed in GT 1b-infected patients in whom treatment has failed.²¹ In addition, the low genetic barrier to resistance and considerable cross-resistance have been implicated as being involved in RASs at positions V36, T54, R155, and A156.¹² The first-generation PIs boceprevir and telaprevir are not recommend for HCV treatment,¹⁵ but in our study only the T54S variant was found, in 14.52% of sequences. Substitutions at positions R155 and D168, which have been confirmed to reduce the activity of simeprevir, asunaprevir, paritaprevir, and vaniprevir, were not found in this study either.^{12,22}

Current NS5A inhibitors are characterized by broad genotypic coverage; however, low barriers to resistance make RASs against NS5A inhibitors of wide concern.²³ The prevalence of RASs relevant to NS5A inhibitors in this study was higher than that reported by a previous study (42.42% vs 29.6%).²⁴ This discrepancy might be because more clinically relevant RASs were identified in this study than in prior work. RASs at positions 28, 30, 31, and 93 have been frequently

observed in both laboratory and clinical trials,²⁵ but in this study the most commonly occurring variant was Q54H, which confers low-level resistance to daclatasvir.²⁶ Furthermore, L31M and Y93H are regarded as key resistance variants, and the simultaneous presence of these two substitutions at baseline will significantly affect the outcome of NS5A-based antiviral therapy.²⁷ The prevalence of Y93H, which confers high-level resistance to DAAs, in this study was 7.58%, which is similar to findings from other reports (3.8%–14.1%).^{27,28} Nevertheless, none of the sequences contained L31M, which confers a low–medium level of resistance to daclatasvir and ledipasvir, and this finding is different from previously reported frequencies (2.1%–6.3%).²⁸ This discrepancy may be a result of differences in sensitivity between population sequencing and ultradeep sequencing for detection of viral populations. Although no Y93H + L31M variants were found in this study, other concomitant variants, such as R30Q and P58S, should be given more attention. These two variants are regarded as secondary-resistance variants, and their presence will enhance the resistance conferred by primary variants.²⁹ In addition, the variant combination of L28M and L31F has been shown to have a higher level of resistance to ombitasvir than either of the variants separately did.³⁰ In this study, L28M was detected without the presence of L31F.

NS5B polymerase inhibitors, including the NNI dasabuvir and the NI sofosbuvir, are potent agents with a high resistance

barrier.³¹ Additionally, no cross-resistance has been observed with currently approved NS5B inhibitors, with one exception, C316N, which is associated with low-level resistance to dasabuvir and sofosbuvir.^{32,33} Baseline polymorphisms at position 316 vary by geographical location and subtype. C316 is highly conserved in GT 1a (99.89%), but polymorphic in GT 1b (81.83%), based on frequencies.³² In addition, HCV 1b isolates harboring C316N at baseline are more frequently observed in Asia compared to Europe and the US (91.6% vs 32% vs 5%, respectively).³⁴ Remarkably, the polymorphic C316N was detected in 100% of the 44 sequences in this study, in agreement with the results of another study in China.³⁵ This indicated that C316N might confer enhanced fitness for chronic HCV 1b patients in China and that more attention should be paid to this substitution in Chinese HCV 1b patients. The other key RASs that may lead to failure of sofosbuvir treatment are L159F, S282T, L320F, V321A, and L159F + V321A, which are observed in 2.2%–4.4% of subjects.³² However, none of these RASs was detected in this study, and it may be that mutations in these positions will greatly influence viral fitness. Moreover, M414L and S556G, which can lead to the treatment failure of dasabuvir, were commonly observed in this study, similarly to the results of previous studies.¹²

In this study, we found that a clade without RASs in the NS5A gene showed a more prominent concentration trend than others. However, we did not find any factors that were responsible for this observation. A larger-population study may be needed to clarify this phenomenon. In addition, the covariance relationships between RAS and non-RAS nodes or between RASs indicated a complex interacting relationship in the HCV genome at baseline. The RASs in the NS3 region, the NA5A region, and the NS5B region affected one another, which may indicate that an RAS found in one of the three regions affects RASs in the other two regions. Therefore, this finding indicates that more attention should be paid to multisegment combined drug resistance when using DAAs in future.

To our knowledge, this study is the first to report the prevalence of RASs in western Chinese chronic HCV 1b patients. Moreover, the latest and most comprehensive panel of clinically relevant RASs was included to analyze all DAA-resistance regions (the NS3, NS5A, and NS5B regions). However, there are several limitations to this study. First, the number of patients was limited, which may have led to relatively high prevalence. Second, some NS5B genes were unsuccessfully amplified, which reduced the sample size of this study. Third, direct sequencing, not ultradeep pyrosequencing, was used in this study, which hindered our

ability to investigate the prevalence of minority variants (<20%), though RASs detected in this study represent the main variants.

In conclusion, the prevalence of RASs clinically relevant to DAAs was high in treatment-naïve HCV 1b-infected patients in western China, especially for variants S122G, Q54H, and C316N. Further studies are needed to clarify the role of these RASs so that we can improve our strategy for managing and using DAAs in China in future.

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Author contributions

PH and HR conceived the study, ZL and ZC conducted the experiments, ZL collected the data, and ZL, ZC, and HL analyzed the data. All authors contributed toward data analysis, drafting and revising the paper and agree to be accountable for all aspects of the work.

Disclosure

The authors report no conflicts of interest in this work.

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Supplementary material

Table S1 Amplification and sequencing primers for HCV NS3,¹ NS5A,² and NS5B³ genes in genotype 1b patients

Target gene	Primers	5'-3' sequences
NS3	NS3F out	CGAGACCTTGC GG TGGCAG
	NS3R out	CAGCCGYTCCGCTTGGTCC
	NS3F in	CATCACCTGGGGGGCAGACACC
	NS3F in	GTCAGTTGAGTGGCACTCATCAC
NS5A	NS5AF out	GTTTGGGACTGGATATGC
	NS5AR out	CGTCACGTAGTGGAAATC
	NS5AF in	ACCTGGCTCCAGTCCAAG
	NS5AF in	CCTCCACRTACTCCTCAG
NS5B	NS5BF out	CGYTGAGTCRTAYTCTCCATGC
	NS5BR out	GGGRCRGAGACASGCTGTGATA
	NS5BF in	CTCAGYGACGGGTCYTGGTC
	NS5BR in	GCTGTGATATATGTCTCCCC

Abbreviation: HCV, hepatitis C virus.

Table S2 Summary of HCV RASs to DAAs

Region	Residue	RAS(s)	DAA(s)	Reference(s)
NS3	C16	S	Tvr	4
	V36	A/M/L/G/I/C	Tvr, Boc, Asv, Ptv, Fdv	5-9
	Q41	R	Smv	10, 11
	F43	S/V/I/L	Boc, Tvr, Ptv, Smv	6, 10, 12
	T54	A, S, C, G, V	Tvr, Boc, Fdv, Asv	5, 9, 10, 12
	V55	A	Boc, Tvr	6, 9
		F, I, K, T	Boc, Asv	1, 13
	Y56	H	Ptv, Asv	5
		L	Asv	5, 10
	Q80	K, R, H, N, G, L	Smv, Asv/Ptv	1, 5, 6, 10
	A87	T	Tvr, Boc	14
	V107	I	Boc	5
	R109	K	Smv, Boc	15
	R117	H	Boc	14
	S122	A, I, R, T, G, N	Smv, Asv	5, 9, 16
	R123	T	Tvr, Boc	17
	V132I	I	Tvr	18
	R155	K/G/T/M/I/L/S/Q/P/N/W	Tvr, Boc, Smv, Vnv, Fdv, Asv, Ptv	5, 6, 9, 10, 19, 20
	A156	S/T/F/N/V/I/G/D	Tvr, Boc, Smv, Vnv, Fdv, Asv, Gzv	5, 6, 10, 19, 20
	V158	I, M	Boc, Tvr	5, 6
	D168	Q/A/Y/V/E/T/N/P/I/H/G/F/S/K	Smv, Vnv, Asv, Fdv, Ptv, Gzv	5, 6, 9, 10, 13, 21, 22
		A	Boc, Tvr	6
	V170	T	Boc, Smv	5
		I	Tvr	18
	S174	F/P	Tvr	14
	M175	L	Boc	5
	NS5A	L28	F, M, T	Dcv, Omv
P29		S, X	Dcv	5
R30		G, H, P, Q	Dcv	5, 10
L31		V/M/F/I	Dcv, Omv, Ldv	5, 6, 9
P32		L, S, X	Dcv, Ldv	5, 6, 10
Q54		H/N/L/Y	Dcv, Omv	18, 23, 24
P58		S/A/H/T	Dcv	5, 24
		D	Ldv	6
Q/E62		D/H	Dcv	5, 18
		R/A/P/S	Dcv	24, 25
A92		K/T	Dcv	5, 6, 24
Y93		H/N/C	Dcv, Ldv, Omv	5, 6, 23, 24, 26

(Continued)

Table S2 (Continued)

Region	Residue	RAS(s)	DAA(s)	Reference(s)
NS5B	H95	Q/R	Dsv	18, 27
	L159	F	Sof, Mcb	5, 28, 29
	N142	T/S	R1479, Sof	27, 30
	S282	T/R	Sof, Mcb	5, 6, 30
	M289	I/L/V	Sof	3, 6
	A300	T	R7128	31
	V321	A	Sof	5
	C316	H, N, Y, S	Dsv, Nbv, Sof	5, 6, 9
	L320	F, I, S	Sof, Mcb, PSI938	6, 28–30
	V321	A	Sof	5
	A338	V	Bcv	18, 32
	S365	A, L, T	Nbv	18, 32, 33
	S368	T	Dsv	6
	L392	I	GS9669	18
	N411	S	Dsv	6
	M414	L/T/I/V/Q	Dsv, A782759, Nbv	5, 6, 13
	A421	V	BMS791325	5, 13
	M423	I, T	Nbv	13, 34
	I424	V	Bcv	35
	M426	T/V/I	A782759, Nbv	3, 34
	A442	T	Bcv	5, 18, 19
	C445	F,	GS9190	15, 36
	E446	K, Q	Dsv	6, 9
	Y448	C, H	Dsv, GS9190	6, 37, 38
	V494	A, T, L, I	Nbv	13, 15
	P495	A, L, S	Bcv, BI207127	5, 27, 36, 39
	V499	A	Bcv	18, 40
	A553	I, T, V	Dsv	6, 10
	G554	S, D	Dsv	6, 10
	S556	G, N, R, C	Dsv	5, 6, 13, 41
G558	R	Dsv	6	
D559	G, S, N	Dsv	6, 9	
Y561	H	Dsv	6	

Abbreviations: HCV, hepatitis C virus; RASs, resistance-associated substitutions; Asv, asunaprevir; Boc, boceprevir; Ptv, paritaprevir; Smv, simeprevir; Tvr, telaprevir; Fdv, faldaprevir; Vnv, vaniprevir; Gzv, grazoprevir; Dcv, daclatasvir; Ldv, ledipasvir; Omv, ombitasvir; Sof, sofosbuvir; Dsv, dasabuvir; Bcv, beclabuvir; Mcb, mericitabine; Nbv, nesbuvir.

Table S3 Baseline characteristics of patients with and without NS5A RASs

Characteristics	Wild type (n=36)	Mutation type (n=16)	P
Age (years), median (range)	50 (18–70)	42 (35–58)	0.192
Male (female)	21 (15)	7 (9)	>0.05
Alb (g/L), median (range)	44.35 (36.4–54.9)	45.25 (24.6–48)	0.513
ALT (U/L), median (range)	61 (10–189)	66 (20–230)	0.505
AST (U/L), median (range)	54 (14–163)	47 (21–199)	0.657
γ -GGT (IU/L), median (range)	57 (15–207)	33 (9–97)	0.054
Total bilirubin (mg/dL), median (range)	0.85 (0.44–1.91)	0.63 (0.37–1.25)	0.597
Direct bilirubin (mg/dL), median (range)	0.34 (0.22–0.79)	0.33 (0.16–0.90)	0.746
Plt count ($\times 10^9/L$), median (range)	153 (57–240)	184 (56–315)	0.323
AST/Plt, median (range)	0.43 (0.12–1.63)	0.26 (0.1–1.80)	0.867
Hemoglobin (g/dL), median (range)	153 (85–171)	142.5 (77.4–178)	0.907
HCV RNA (\log_{10} IU/mL), median (range)	6.35 (3.00–7.5)	6.57 (4.37–7.23)	0.444

Abbreviations: HCV, hepatitis C virus; RASs, resistance-associated substitutions; Alb, albumin; Plt, platelet.

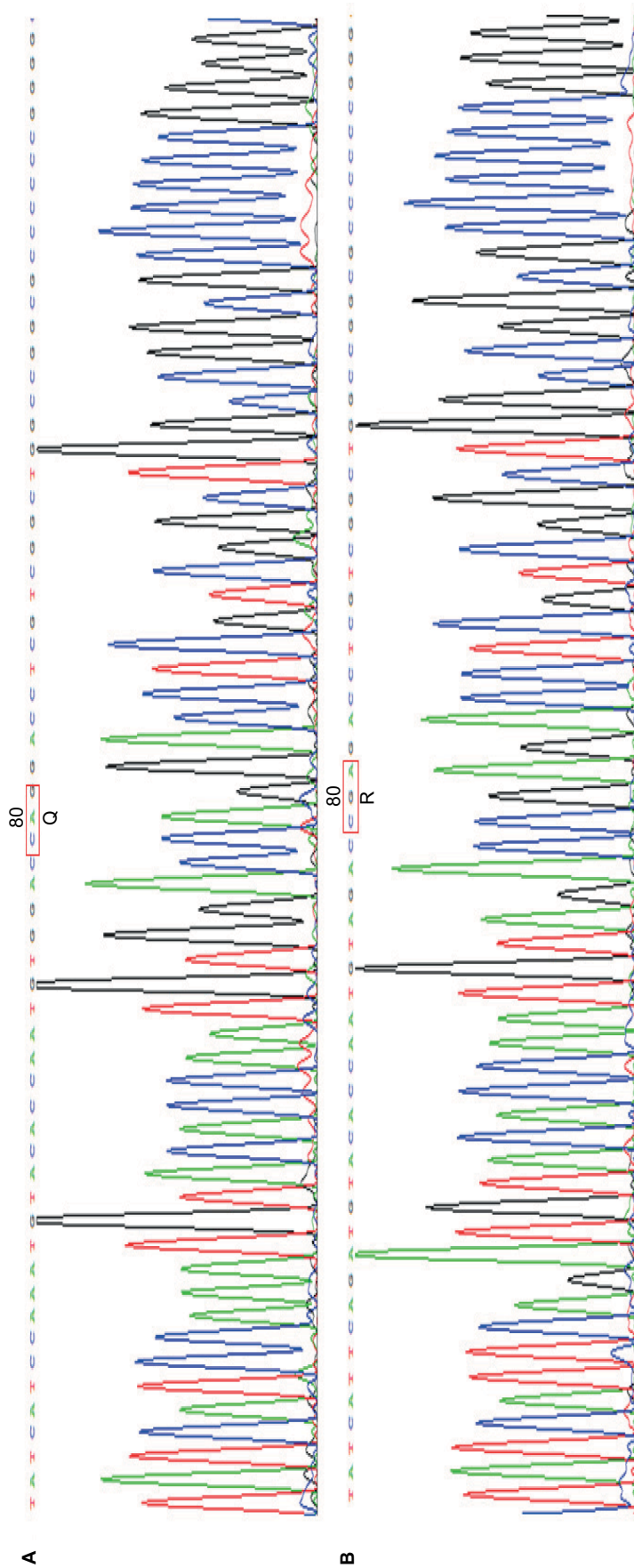


Figure S1 Chromatograms of Sanger sequencing.

Notes: Chromatograms of Sanger sequencing in part of NS3 segment with 80Q (**A**); with 80R (**B**). Amino acid site 80 occurring bases substitution leading to amino acid Q change to R which resistance to DAAs, thus, Q80R was described as a variant type.

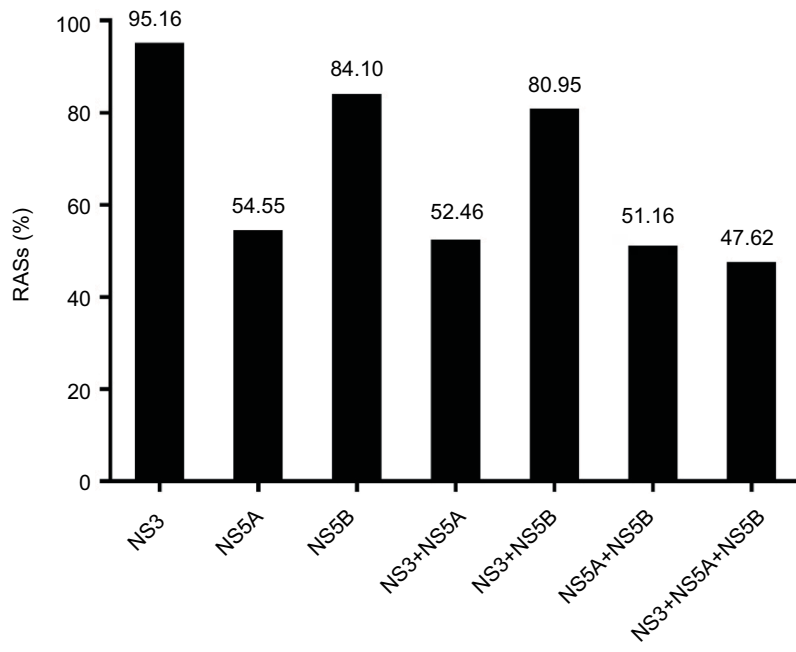


Figure S2 The prevalence of clinically relevant resistance associated substitutions (without C316N) in various DAA classes.

Abbreviation: RASs, resistance-associated substitutions.

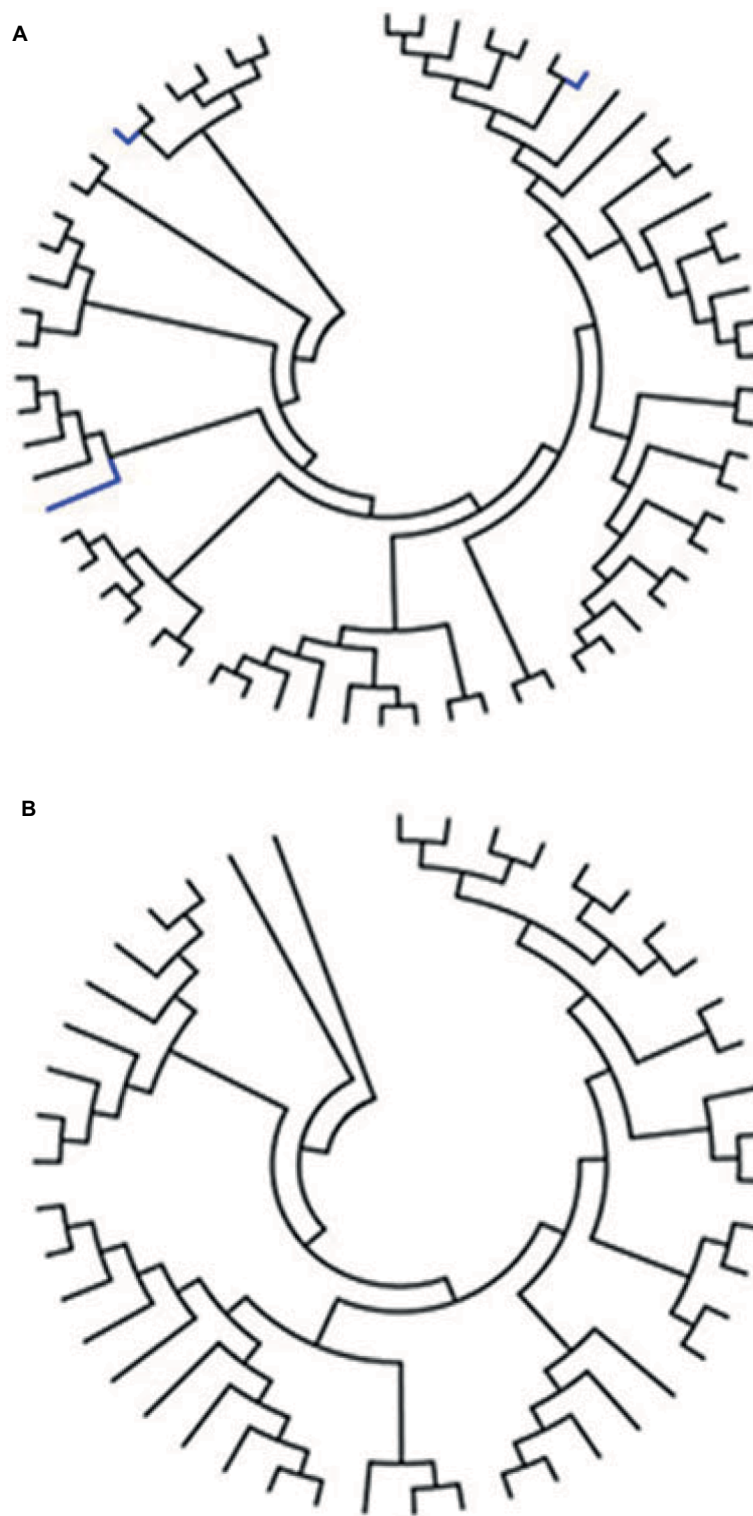


Figure S3 Phylogenetic trees for HCVGT1b NS3 and NS5B genes.

Notes: (A) Phylogenetic tree for HCV GT1b NS3 gene; (B) phylogenetic tree for HCV GT1b NS5B gene. Blue lines represent nonresistant lineages; black lines represent lineages carrying the resistance associated substitutions.

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