Hepatitis C Virus of Subtype 2l in Marseille, Southeastern France

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**Key Words**
Hepatitis C virus · Genotype · Subtype 2l · Treatment · Polymerase

**Abstract**
The rate of eradication of chronic hepatitis C considerably increases with direct-acting antiviral agents, particularly hepatitis C virus (HCV) polymerase inhibitors. While implementing full-length HCV NS5B polymerase sequencing in our clinical microbiology laboratory, we identified atypical HCV sequences, classified as subtype 2l, from 2 patients. HCV-2l NS5B polymerase sequences were detected from 5 and 14 additional patients by screening our laboratory hepatitis virus sequence database and the NCBI GenBank sequence database. Phylogenetic analyses show unambiguously that all HCV-2l sequences are clustered apart from HCV 2 non-l sequences, which compose a second cluster. Mean (±SD) nucleotide identity between near full-length NS5B fragments of subtype 2l was 93.4 ± 0.8% (range: 92.4–95.1). Of note, all HCV-2l sequences obtained in our laboratory and in other centers were from serum samples collected in France. Analysis of the HCV-2l NS5B polymerase amino acid sequences at 30 positions critical for interaction with or resistance to HCV polymerase inhibitors showed specific patterns.

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**Introduction**

Hepatitis C virus (HCV) infection entered a new era with the availability of direct-acting antiviral agents to treat chronic infections in combination with pegylated interferon and ribavirin [1, 2]. Thus, clinical trials have shown considerable increases in the rates of sustained virological response. Direct-acting antiviral agents encompass HCV NS3 protease inhibitors, approved for the treatment of chronic HCV genotype 1 infections, and pangenotypic inhibitors of HCV NS5B RNA-dependent RNA polymerase and HCV NS5A.
HCV is characterized by a high level of genetic diversity [3]. Seven HCV genotypes and more than 100 subtypes, including 69 confirmed, were described. Genomes of the same subtype differ by <13% over their complete coding region sequences, and those that differ by >15% belong to different genotypes and subtypes. While implementing full-length HCV NS5B polymerase gene sequencing in our laboratory, we identified atypical HCV sequences related to recently proposed subtype 2l [3, 4], which were obtained from the serum of 2 patients. Then, we screened our laboratory hepatitis virus sequence database [5] and detected that HCV sequences of subtype 2l had been obtained from serum samples of 5 additional patients, whereas HCV-2l sequences collected from only 14 patients were detected in the NCBI GenBank nucleotide sequence database.

Materials and Methods

Conserved regions in HCV RNA-dependent RNA polymerase gene (NS5B) were identified and used to design PCR primers using the SVARAP tool [6] based on a set of aligned HCV reference genomes obtained from the Los Alamos HCV sequence database [7] (online suppl. table S1; for all online suppl. material, see www.karger.com/10.1159/000369015). HCV RNA was extracted from 200 μl of serum using the EZ1 viral nucleic acid extraction kit (Virus Mini kit version 2.0, Qiagen, Courtaboeuf, France) according to the manufacturer’s recommendations. Aliquots (10 μl) of extracted RNA were amplified by reverse transcription PCR using 0.4 μM of each dNTP, 0.5 μl of SuperScript III/Platinum Taq polymerase (Invitrogen) for initial denaturation, then 40 cycles including 30 s at 95°, 45 s at 56° and 2 min at 72°, and a final elongation step at 72° for 10 min. Thereafter, 1,570 nucleotide-long HCV genome fragments that cover the near full-length NS5B-encoding gene were obtained using the ABI Prism 3100xl Genetic Analyzer (Applied Biosystems, Branchburg, N.J., USA) under conditions previously described [5] and were analyzed using the SeqScape v2.5 software (Applied Biosystems). Primers used for PCR amplification and sequencing are indicated in online suppl. table S1. In addition, a full-length HCV NS3 protease encoding gene and a fragment of 5′-untranslated region (5′-UTR) were obtained as previously described [5]. These sequences were aligned using Muscle software [8] with an HCV sequence reference panel [7]. Pairwise p-distances were computed using the BioEdit software (http://www.mbio.ncsu.edu/bioedit/bioedit.html). Phylogenetic trees were built using MEGA v5.1 software [9] with the maximum-likelihood method based on the most appropriate substitution model, as determined by MEGA v5.1 for the sequence set, or the neighbor-joining method based on the Kimura 2-parameter model; tests of phylogeny used the bootstrap method with 1,000 resamplings of the data. BLAST searches were done for HCV sequences obtained in the present study against the NCBI GenBank nucleotide sequence database (http://blast.ncbi.nlm.nih.gov/Blast.cgi?PROGRAM=blastn&PAGE_TYPE=BlastSearch&LINK_LOC=blasthome) and our clinical microbiology laboratory sequence database [5]. Intergenotypic or intersubtypic recombination was excluded using the SimPlot tool (sray.med.ohiohealth.edu/SCRoftware/simplot/) through comparison with a reference set of HCV genomes, including those classified as 2l (KC197235 and KC197240).

Results

HCV sequences corresponding to the near full-length HCV NS5B polymerase gene, most similar to genotype 2l apart from other genotype 2 HCV, were first obtained from serum samples of 2 case-patients (fig. 1). These sequences exhibited 93% nucleotide identity between each other. BLAST searches against our laboratory sequence database identified 5 other sequences that belong to the same phylogenetic cluster. In addition, these 2 sequences...
showed 93–94 and 92–95% identity, respectively, with their 3 top BLAST hits in the NCBI GenBank nucleotide sequence database, which were genomes KC197235 and KC197240 that were released in GenBank in September 2013 and classified as subtype 2l [3, 4] and sequence FJ872270 [10]. These 3 latter sequences were obtained in France. Apart from these 3 sequences, the 10 best BLAST hits in GenBank included HCV sequences of subtypes 2c, 2q or 2b, with 79–80, 79 and 79% nucleotide identity, respectively, for the first NS5B sequence recovered here, and HCV sequences of subtypes 2c or 2b with 79% nucleotide identity for the second NS5B sequence recovered here. Overall, mean (±SD) nucleotide identity between NS5B long fragments of subtype 2l was 93.4 ± 0.85% (range: 92.4–95.1), whereas the mean nucleotide identity between sequences of subtype 2l and 2 non-l was 77.9 ± 0.80% (75.7–79.9). Therefore, sequences represented in the present work can be classified as of subtype 2l because, in addition to their phylogenetic clustering with the 2 recently described HCV-2l genomes, they exhibit far lower nucleotide differences with HCV-2l sequences than those defined as threshold for subtype classification for genotype 2 (13.1–17.6%) in the recent article on expanded classification of HCV genotypes [3]. Intergenotypic or intersubtypic recombination was excluded by using the SimPlot tool (sray.med.som.jhmi.edu/SCSoftware/simplot/) with HCV reference genomes (including HCV-2l genomes).

BLAST searches using a shorter HCV NS5B fragment (nt 8,342–8,650 in reference to HCV 2a genome accession No. AF177036) classically used for genotype identification and used in routine clinical microbiology practice in our institution detected 5 additional HCV-2l sequences in our laboratory sequence database and 9 additional HCV-2l sequences in the GenBank sequence database, 7 being labeled as 2l (fig. 2). All of these additional HCV sequences were obtained in France. The same results were obtained when incorporating into the analysis sequences from the recent article by Smith et al. [3] on expanded classification of HCV genotypes and other HCV-2 non-l sequences recovered from the GenBank sequence database (online suppl. fig. S1). Two of these sequences were branched with HCV subtype 2l sequences, apart from HCV-2 non-l subtypes, but they exhibited a mean nucleotide identity of 84.3 ± 1.4 (81.3–86.6) with HCV-2l sequences. In congruence with the HCV NS5B encoding gene, the best BLAST hits for HCV NS3 protease encoding sequences (543 nucleotides) obtained from the 2 case-patients were sequences identified either as belonging to subtype 2l (KC197235 and KC197240 [4]; with 91–92% nucleotide identity for the first sequence and 87–88% nucleotide identity for the second sequence) or as belonging to genotype 2 but unclassified at the subtype level (HQ623293 and HQ623295 [10]; with 91% nucleotide identity for the first sequence and 88–91% nucleotide identity for the second sequence; fig. 3). All of these HCV NS3 sequences found in GenBank had been recovered from serum samples collected in France. Finally, the best matches for 5′-UTR nucleotide sequences obtained from the 2 case-patients were the 2 HCV-2l genomes (KC19740 and KC197235, with 99% nucleotide identity; online suppl. fig. S2).

Overall, HCV-2l sequences represented 0.25, 0.15 and 1% of 2,788, 1,333 and 688 HCV NS5B polymerase, NS3 protease and 5′-UTR sequences, respectively, available in our laboratory. For all HCV sequences analyzed here, phylogeny reconstructions delineated with strong confidence of phylogeny used the bootstrap method with 1,000 resamplings of the data. This analysis involved a set of 58 nucleotide sequences with a total of 250 positions (nucleotides 8,344–8,591 in reference to GenBank accession No. NC004102) in the final dataset. A tree built by using the maximum likelihood method with an expanded set of HCV genotype 2 sequences is shown in online supplementary figure S1.

**Fig. 2.** Phylogenetic reconstruction based on a short fragment of the HCV NS5B RNA polymerase encoding gene recovered in the present study, their best BLAST hits in our laboratory [5] and the NCBI nucleotide sequence databases (http://blast.ncbi.nlm.nih.gov/Blast.cgi?PROGRAM=blastn&PAGE_TYPE=BlastSearch&LINK_LOC=blasthome) and a set of HCV reference genomes [7]. The legend is the same as for fig. 1, except that the tree was built using the neighbor-joining method with the Kimura 2-parameter model; test bootstrap method with 1,000 resamplings of the data. This analysis involved a set of 58 nucleotide sequences with a total of 250 positions (nucleotides 8,344–8,591 in reference to GenBank accession No. NC004102) in the final dataset. A tree built by using the maximum likelihood method with an expanded set of HCV genotype 2 sequences is shown in online supplementary figure S1.

**Fig. 3.** Phylogenetic reconstruction based on HCV NS3 protease encoding gene recovered in the present study, their best BLAST hits in our laboratory [5] and the NCBI nucleotide sequence databases (http://blast.ncbi.nlm.nih.gov/Blast.cgi?PROGRAM=blastn&PAGE_TYPE=BlastSearch&LINK_LOC=blasthome) and a set of HCV reference genomes [7]. The legend is the same as for fig. 1, except that the tree was built using the neighbor-joining method with the Kimura 2-parameter model; test of phylogeny used the bootstrap method with 1,000 resamplings of the data. The analysis involved a set of 39 nucleotide sequences with a total of 542 positions (nucleotides 3,420–3,962 in reference to GenBank accession No. NC004102) in the final dataset.

(For figures 2 and 3 see next pages.)
a cluster encompassing HCV-2l sequences apart from other genotype 2 subtypes (fig. 1–3; online suppl. fig. S1).

The same phylogenetic tree topology as the near full-length NS5B nucleotide sequences was obtained with the near full-length NS5B amino acid sequences (online suppl. fig. S3). Analysis of the NS5B RNA-dependent RNA polymerase amino acid sequence at 30 positions critical for interaction with or resistance to HCV polymerase inhibitors [11–14] showed presence of amino acid substitution C451T in all HCV-2l sequences, as in all HCV genotype 4, 6 and 7 sequences and most of the genotype 3 HCV, whereas subtypes 2 non-l HCV harbored the C451V substitution. The M414L substitution that confers resistance to HCV nonnucleoside inhibitors [15, 16], already described in most of the genotype 4 HCV [17], was found in all HCV-2l sequences, whereas HCV of subtypes 2 non-l harbor amino acid substitution M414Q that was also associated with drug resistance in replicon assays [18]. In contrast, among the amino acid substitutions identified at 7 positions in the NS3 protease as associated with decreased susceptibility to HCV protease inhibitors, only the Q80G substitution was found in HCV-2l, as in other HCV genotype 2 subtypes [13].

Clinical and epidemiological data were available for 4 of the 7 HCV-2l-infected patients followed up at our institution and are summarized in table 1.

**Discussion**

We describe here 7 HCV NS5B RNA polymerase sequences of subtype 2l, representing more than half the number of sequences (n = 12) previously available worldwide. When considering any HCV genome region, only 19 HCV-2l sequences were available from GenBank, corresponding to 14 HCV strains, including only 2 full-length genomes released in 2013, and some of these sequences were un- or misclassified at the subtype level. Phylogenetic analyses based on HCV NS5B and NS3 genes show in congruence with a previous report [4] that these HCV sequences are clustered together apart from HCV 2 non-l sequences that compose a second cluster. In addition, HCV NS5b sequences recovered here can be classified as 2l based on their level of nucleotide identity with other HCV-2l genomes and with sequences classified as 2 non-l based on the recent recommendation for the classification within HCV genotype 2 subtypes [3]. It is noteworthy that all currently available HCV-2l sequences and those recovered in the present study were obtained from serum samples collected in France (in metropolitan France in all but 1 case) and the reason for this geographical distribution is unresolved. In addition, HCV-2l NS5b polymerases were found to specifically harbor 2 amino acids that may influence susceptibility to currently available HCV polymerase inhibitors, albeit the clinical relevance of these observations is unclear, particularly with respect to the tremendous potency of these drugs. Taken together, previous findings indicate that HCV-2l has been rarely detected worldwide and only from serum samples collected in France, and they highlight that HCV diversity could still be incompletely explored.

**Disclosure Statement**

No potential conflict of interest or financial disclosure for all authors.
References


