Full-length sequences of 11 hepatitis C virus genotype 2 isolates representing five subtypes and six unclassified lineages with unique geographical distributions and genetic variation patterns

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In this study, we characterized full-length hepatitis C virus (HCV) genome sequences for 11 genotype 2 isolates. They were isolated from the sera of 11 patients residing in Canada, of whom four had an African origin. Full-length genomes, each with 18–25 overlapping fragments, were obtained by PCR amplification. Five isolates represent the first complete genomes of subtypes 2d, 2e, 2j, 2m and 2r, while the other six correspond to variants that do not group within any assigned subtypes. These sequences had lengths of 9508–9825 nt and each contained a single ORF encoding 3012–3106 aa. Predicted amino acids were carefully inspected and unique variation patterns were recognized, especially for a 2e isolate, QC64. Phylogenetic analysis of complete genome sequences provides evidence that there are a total of 16 subtypes, of which 11 have been described here. Co-analysis with 68 partial NS5B sequences also differentiated 18 assigned subtypes, 2a–2r, and eight additional lineages within genotype 2, which is consistent with the analysis of complete genome sequences. The data from this study will now allow 10 assigned subtypes and six additional lineages of HCV genotype 2 to have their full-length genomes defined. Further analysis with 2021 genotype 2 sequences available in the HCV database indicated that the geographical distribution of these subtypes is consistent with an African origin, with particular subtypes having spread to Asia and the Americas.

INTRODUCTION

Determination of hepatitis C virus (HCV) genotype is of clinical relevance (Simmonds et al., 2005), which is shown by differences in the sensitivity to alpha interferon and ribavirin combination therapy among patients. Those infected by genotypes 2 and 3 have a higher rate of sustained virological response than those infected by genotypes 1 and 4 (Feld & Hoofnagle, 2005; Manns et al., 2001, 2006; Pearlman, 2004). In addition, genotype 3 has been linked to the development of more pronounced hepatocellular steatosis (Abid et al., 2005; Hourioux et al., 2007; Jackel-Cram et al., 2007; Negro, 2006; Rubbia-Brandt et al., 2000). Although the mechanisms involved are currently not understood, there is probably an association with the genetic variation of the virus. Understanding the origin and nature of HCV genetic diversity is critical for defining preventive strategies and for the development of therapies and a vaccine. Such diversity is the source of information about the virus’s epidemic history prior to its discovery, which has not been well-documented in non-Western countries. Canada is a country encompassing many immigrants from non-Western countries, among which a variety of HCV genotypes and subtypes have been detected (Murphy et al., 2007). These include a number of genotype 2 isolates isolated in the Canadian province of Quebec. In addition to the globally prevalent 2a, 2b and 2c subtypes, other isolates representing less common subtypes (2d, 2e, 2i, 2j, 2k, 2m, 2r) and many novel variants were identified. This study focused on defining the full-length...
genome sequences for 11 of these strains, which have a limited spread, with the purpose of analysing their genetic diversity and organization. Our findings should add insights to HCV genomic studies and to clinical applications.

Although Africa is commonly thought to be the region where genotype 2 has originated, a whole spectrum of genotype 2 subtypes and unclassified variants have been identified outside this continent (Simmonds et al., 2005; Pouillot et al., 2008; Markov et al., 2009). This raises the question, could HCV genotype 2 have originated in another region such as Europe? In the current study, we addressed this question in association with the geographical distribution and genetic variation of all genotype 2 sequences available in the Los Alamos HCV database (http://hcv.lanl.gov/).

RESULTS

Genome sequence and organization

Full-length genomic sequences were successfully obtained for 11 HCV genotype 2 isolates each with 18–25 overlapping fragments (see Fig. S1, available in JGV Online). These sequences had the lengths of 9508–9825 nt, starting from the extreme 5' UTR termini through to the 3' poly(U) tracts or the highly conserved 3' X tails (Table 1). Four isolates QC64/2e, QC178/2m, QC302/2 and QC331/2 had their sequences determined up to the 3' X tail while all the other isolates only had their sequences determined to the poly(U) tract. The genomes each contained a single ORF of 9036–9318 nt, with 10 protein-coding regions of the following sizes: core (573 nt/191 aa), E1 (576 nt/192 aa), E2 (1095–1101 nt/365–367 aa), p7 (189–390 nt/63–130 aa), NS2 (651 nt/217 aa), NS3 (1893 nt/631 aa), NS4A (162 nt/54 aa), NS4B (783 nt/261 aa), NS5A (1398–1413 nt/466–471 aa) and NS5B (1776 nt/591 aa). Among these regions, E2, p7 and NS5A were variable in length (Table 1).

Genotype 2 classification

Fig. 1(a) shows the maximum-likelihood tree reconstructed with full-length genome sequences for all HCV genotypes. The tree displays seven major branches/clusters, each representing one genotype. Each major branch showed a bootstrap support of 100% when a cluster was formed (i.e. genotypes with more than one subtype). Within genotype 2, the 11 isolates were distinct from subtypes 2a, 2b, 2c, 2i and 2k, for which full-length sequences were previously determined. Prior to our report, subtypes 2d, 2e, 2j, 2m and 2r were provisionally assigned only based on partial sequences. In the present study, these five subtypes were confirmed by their full-length genomes being determined. In addition, six isolates QC114, QC182, QC289, QC297, QC302 and QC331 showed evidence that they represent six new lineages. Fig. 1(b) shows the maximum-likelihood tree based on the amino acid sequences deduced for these isolates. A topology consistent with that in Fig. 1(a) was observed.

Fig. 2 shows a neighbour-joining tree reconstructed with full-length sequences for genotype 2. Besides the 11 isolates from the current study, the tree also included 47 sequences retrieved from the HCV database. Among them, 19 belonged to subtype 2a (12 from Japan, seven with unknown origin), 25 belonged to subtype 2b (23 from Japan and two from the USA), and one belonged to each of subtype 2c (from an Italian patient) (Nakao et al., 1996), subtype 2k (Samokhvalov et al., 2000) and subtype 2i (Noppornpanth et al., 2006). The tree showed that the 11 isolates were distinct from each other and from subtypes 2a, 2b, 2c, 2k and 2i. This provides evidence that the 11 isolates represent 11 distinct genotype 2 subtypes/lineages.

A segment of the NS5B region, corresponding to nt 8276–8615 in the numbering of the H77 genome (GenBank accession no. NC_004102), is reliable for differentiating HCV genotypes and subtypes. This region was further analysed using sequences from 68 genotype 2 isolates in addition to our 11 genotype 2 isolates. The tree (Fig. 3) shows 26 branches or clusters corresponding to assigned subtypes and unclassified lineages, congruent with those obtained for the complete genome sequences. Among them 18 (yellow-coloured branches) have been assigned subtypes 2a–2r and eight (green-coloured branches) have not been assigned to a subtype. QC259, QC64, QC232, QC178 and QC283 represent five of the 18 assigned while QC114, QC182, QC289, QC297, QC302 and QC331 represent six of the eight unassigned. QC259/2d grouped with five isolates, two from the Netherlands (Stuyver et al., 1994) and three from Africa (Jannet et al., 1998); QC64/2e grouped with 10 isolates all from Indonesia (Tokita et al., 1996; Utama et al., 2010); QC232/2j grouped with nine isolates, one from Spain (BA047) (Holland et al., 1996) and eight from France (Thomas et al., 2007; Cantaloube et al., 2008; Le Pogam et al., 1998); QC178/2m and QC283/2r each grouped with three isolates, all from Canada; QC297 grouped with two isolates, one from the French Caribbean island of Martinique (Martial et al., 2004) and one from France (Thomas et al., 2007); QC331 grouped with three isolates all from Cameroon (Njouom et al., 2003a, 2005; Ndjomou et al., 2003a); QC182 grouped with eight isolates, two from Canada and six from Cameroon (Ndjomou et al., 2003; Pasquier et al., 2005; Njouom et al., 2003b). In contrast, QC302 and QC289 each represented an orphan isolate while QC114 grouped most closely with MRS50, a variant reported in France (Cantaloube et al., 2008).

Geographical distribution of HCV genotype 2 isolates

In the Los Alamos HCV database (end of year 2010), a total of 4044 sequences have been classified in genotype 2. These sequences were filtered out and a total of 201 individual isolates were distinguished that had their sampling countries recognized. The geographical distribution of these isolates is summarized in Tables 2 and 3.
Subtypes 2a, 2b and 2c were ubiquitous; they were identified in 31, 21 and 19 countries/regions, respectively, accounting for 21.6, 24.7 and 17.8% of the total 2021 isolates. This was followed by subtype 2k, which was found in nine countries in Europe, America, Africa, Asia and the Middle East, accounting for 3.27% of isolates. In contrast, subtype 2i can be found only in four countries but in different continents; 2d, 2e, 2f and 2j each in three countries; 2h and 2l each in two countries; and the remaining seven subtypes (2g, 2m, 2n, 2o, 2p, 2q, 2r) each in one country. Diversity of reported genotype 2 isolates was the largest in Europe, especially in France, where a total of 17 subtypes were identified. However, 85.6% (107 out of 125) of the African isolates were of unknown subtype, and 14.4% (18 out of 125) belonged to six different subtypes (Table 3). The area of greatest diversity is presumably the region of origin, which therefore suggests Africa.

Pairwise comparison

Pairwise comparisons were performed between the 11 isolates and other isolates within genotype 2. The differences ranged from 16.0% (QC82 vs QC31) to 23.6% (2e_QC64 vs 2b_JPUT971017) at the nucleotide level and from 10.5% (QC259 vs QC297) to 16.5% (2e_QC64 vs 2b_JPUT971017) at the amino acid level. These results further support Fig. 1 and distinguish each isolate; they each belong to a different subtype.

To exclude the possibility of viral recombination, pairwise nucleotide similarity curves were plotted along HCV genomes. Comparing the 11 isolates against each other and against the 19 reference sequences representing different genotypes and subtypes shown in Fig. 1(a), no such evidence was revealed (data not shown).

Table 1. Patient information for the 11 genotype 2 isolates and the number of nucleotides/amino acids in each genomic region

<table>
<thead>
<tr>
<th>ID</th>
<th>Age</th>
<th>Sex</th>
<th>Origin</th>
<th>Full ORF</th>
<th>5' UTR</th>
<th>Core</th>
<th>E1</th>
<th>E2</th>
<th>p7</th>
<th>NS2</th>
<th>NS3</th>
<th>NS4A</th>
<th>NS4B</th>
<th>NS5A</th>
<th>NS5B</th>
<th>3' UTR</th>
</tr>
</thead>
<tbody>
<tr>
<td>H77+</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>9646</td>
<td>9036/3011</td>
<td>341</td>
<td>573/191</td>
<td>576/192</td>
<td>1089/363</td>
<td>189/63</td>
<td>651/217</td>
<td>1893/631</td>
<td>162/54</td>
<td>783/261</td>
<td>1344/448</td>
<td>1776/591</td>
</tr>
<tr>
<td>QC288</td>
<td>53</td>
<td>F</td>
<td>Haiti</td>
<td>9512</td>
<td>9102/3033</td>
<td>340</td>
<td>573/191</td>
<td>576/192</td>
<td>1101/367</td>
<td>189/63</td>
<td>651/217</td>
<td>1893/631</td>
<td>162/54</td>
<td>783/261</td>
<td>1398/466</td>
<td>1776/591</td>
</tr>
<tr>
<td>QC297</td>
<td>68</td>
<td>F</td>
<td>Italy</td>
<td>9510</td>
<td>9096/3031</td>
<td>340</td>
<td>573/191</td>
<td>576/192</td>
<td>1101/367</td>
<td>189/63</td>
<td>651/217</td>
<td>1893/631</td>
<td>162/54</td>
<td>783/261</td>
<td>1398/466</td>
<td>1776/591</td>
</tr>
</tbody>
</table>

*The H77 genome (GenBank accession no. NC_004102), HCV genotype 1, is included for comparison.
insertion of 201 nt (67 aa) located directly after the p7 gene. Since the insertion appeared to be related to the p7 gene itself, a phylogeny was constructed for p7 sequences of all genotype 2 subtypes and representative sequences of the six other genotypes. For QC64, the 189 nt corresponding to the standard p7 sequence and the 201 nt corresponding to the insertion sequence were individually included in the analysis. Interestingly, this QC64 insertion was shown to cluster with the p7 sequence from QC64, and they both shared a nucleotide similarity of 78%. The phylogenetic tree and the alignment of amino acid sequences are shown in Fig. 4.

**NS5A.** Genotype 2 isolates show the longest NS5A region. For the 11 isolates, this region ranged between 1398 and 1413 nt (466–471 aa). Various insertions were observed in NS5A. To begin with, the interferon-sensitivity determining region (ISDR) of QC64 showed an insertion of 4 aa at amino acid numbering 2303–2306 (aa 261–264 in NS5A). Such an insertion is also observed in the BEBE1/2c isolate and in genotypes 1, 3, 5, 6 and 7 isolates, but not in genotype 4 isolates. An insertion was also observed in both QC64 and QC178 directly at the end of protein kinase R-binding domain (PKR-BD) region following the ISDR. They each had an insertion of 1 aa, corresponding to amino acid numbering 2349 of QC64 and 2278 of QC178 (aa 306 and 302 in NS5A, respectively). In the V3 region, QC182 and QC331 each displayed an insertion of 4 aa at amino acid numbering 2376–2379 and 2375–2378. Finally, when genotype 2 sequences as a whole were compared to the other HCV genotypes, they were found to contain an insertion in domain III of NS5A. The insertion occurred following the sequence SMPPLEGEPGDPDL (amino acid numbering 415–428 in the NS5A of H77) that is highly
conserved in all HCV genotypes. At this position, genotype 2 isolates showed an insertion of 20 aa that was highly variable between subtypes with the median p-distance being 0.65 (0.30–0.90) (data not shown).

**DISCUSSION**

In this study, full-length genome sequences were characterized for 11 genotype 2 isolates. Among them, QC259, QC64,
QC232, QC178 and QC283 were previously assigned to subtypes 2d, 2e, 2j, 2m and 2r, respectively, based on partial C/E1 and NS5B sequences (Murphy et al., 2007). In the consensus paper, these five subtypes have been classified as 'provisionally assigned subtypes' (Simmonds et al., 2005) due to the absence of available full-length genome sequences. In this study, the full-length sequences of the five isolates all grouped independently from each other and from other genotype 2 isolates for which complete sequences are available. This now allows for the confirmed designations of subtypes 2d, 2e, 2j, 2m and 2r.

Full-length sequences were also obtained for six isolates designated QC114, QC182, QC289, QC297, QC302 and QC331. They also grouped independently from each other and from other genotype 2 complete sequences. However, we prefer not to assign them to new subtypes at the present moment. QC114, QC289 and QC302 do not group with a sufficient number of closely related isolates; they are not considered for a new subtype designation. Analysis of partial sequences in the NS5B region showed QC182, QC331 and QC297 each clustered with three or more isolates from the same geographical regions (Fig. 3). However, only a fraction of unclassified variants were included in Fig. 3. Inclusion of additional variants in the analysis did not allow for clear subtype separations (data not shown). They may represent continual variants resulting from endemic evolution. We would suggest that new subtype designations be given for these isolates only after their epidemiological significance is approved, the possibility of viral recombination is excluded, and if possible, more related candidates are entirely sequenced (Simmonds et al., 2005).

The Los Alamos HCV database contained 53 complete sequences of genotype 2. Among them, 38 had their geographical origins recognized. This included nine subtype 2a isolates from Asia, 23 subtype 2b isolates from both Asia and North America, one each of subtypes 2c, 2k and 1b/2k from Europe, and one each of subtypes 2i, 2i/6p and 2b/1b from Asia. Albeit 18 subtypes have been designated, the 53 full-length sequences only represent five of the genotype 2 subtypes (excluding recombinants), most of them belonging

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**Table 2. Geographical distribution of HCV genotype 2 subtypes**

<table>
<thead>
<tr>
<th>Subtype</th>
<th>No. of countries in which subtype is detected</th>
<th>Total (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2a</td>
<td>31</td>
<td>437 (21.6)</td>
</tr>
<tr>
<td>2b</td>
<td>21</td>
<td>499 (24.7)</td>
</tr>
<tr>
<td>2c</td>
<td>19</td>
<td>360 (17.8)</td>
</tr>
<tr>
<td>2k</td>
<td>9 (AZ, CA, FR, MD, MG, MQ, RU, UZ, VN)*</td>
<td>66 (3.27)</td>
</tr>
<tr>
<td>2i</td>
<td>4 (CA, DE, FR, VN)</td>
<td>36 (1.78)</td>
</tr>
<tr>
<td>2d</td>
<td>3 (BJ, NL, CA)</td>
<td>6 (0.300)</td>
</tr>
<tr>
<td>2e</td>
<td>3 (ID, CA, BN)</td>
<td>22 (1.09)</td>
</tr>
<tr>
<td>2f</td>
<td>3 (DE, ID, CN)</td>
<td>5 (0.247)</td>
</tr>
<tr>
<td>2j</td>
<td>3 (FR, CA, BA)</td>
<td>9 (0.445)</td>
</tr>
<tr>
<td>2h</td>
<td>2 (DE, GN)</td>
<td>2 (0.099)</td>
</tr>
<tr>
<td>2l</td>
<td>2 (FR, MQ)</td>
<td>8 (0.396)</td>
</tr>
<tr>
<td>2g</td>
<td>1 (GN)</td>
<td>10 (0.495)</td>
</tr>
<tr>
<td>2m</td>
<td>1 (CA)</td>
<td>4 (0.198)</td>
</tr>
<tr>
<td>2n</td>
<td>1 (NL)</td>
<td>1 (0.045)</td>
</tr>
<tr>
<td>2o</td>
<td>1 (FR)</td>
<td>2 (0.099)</td>
</tr>
<tr>
<td>2p</td>
<td>1 (NL)</td>
<td>1 (0.045)</td>
</tr>
<tr>
<td>2q</td>
<td>1 (BA)</td>
<td>1 (0.045)</td>
</tr>
<tr>
<td>2r</td>
<td>1 (CA)</td>
<td>4 (0.198)</td>
</tr>
<tr>
<td>Recomb.</td>
<td>6 (IE, VN, PH, AR, RU, UZ)</td>
<td>48 (2.38)</td>
</tr>
<tr>
<td>2?</td>
<td>31</td>
<td>500 (24.7)</td>
</tr>
<tr>
<td>Total</td>
<td>56</td>
<td>2021 (100.0)</td>
</tr>
</tbody>
</table>

*Country codes officially assigned in ISO 3166-1: AR, Argentina; AZ, Azerbaijan; BA, Bosnia; BJ, Benin; BN, Brunei Darussalam; CA, Canada; CM, Cameroon; CN, China; DE, Germany; FR, France; GN, Guinea; ID, Indonesia; IE, Ireland; MD, Moldova, Republic of; MG, Madagascar; MQ, Martinique; NL, the Netherlands; PH, the Philippines; RU, Russian Federation; SP, Spain; UZ, Uzbekistan; VN, Vietnam (http://en.wikipedia.org/wiki/ISO_3166-1_alpha-2).
Molecular epidemiology studies have revealed much greater diversity of HCV in certain regions of sub-Saharan Africa and in South and South-East Asia (Simmonds et al., 2005). Africa is where the diversity of genotype 2 isolates is thought to be the greatest. Table 3 showed that there are six subtypes found in Africa but more than 80% genotype 2 sequences have not been classified into any subtypes. Although their origin in West Africa has been validated (Markov et al., 2005), genotype 2 viruses are widely distributed. Not only have they been found in other continents, particular subtypes have also shown unusual patterns of prevalence. The identification of 2e isolates all with an Indonesian origin is such an example. Another example is observed for 2m isolates, which are exclusively detected in Vietnam or in immigrants from Vietnam (unpublished data). Similarly, 2r isolates have been identified only in patients originating from the island of Hispaniola. Our speculation is that all genotype 2 subtypes had an ancestral origin from Africa but many had been later brought to Europe and European dominions by immigrants. This would be supported by the finding of a high number of distinct lineages in Africa. However, the number of lineages identified in Africa has been lower than that on other continents (Table 3). This could be due to two reasons. First, there has been limited sampling in Africa compared with that in the developed countries in Europe and in the other parts of the world. For example, 2i which is found in Europe, Vietnam and Canada has recently been found in Morocco in Africa (based on the updated database on March, 2011) (Qu et al., 1994; Cantaloube et al., 2003; Noppornpanth et al., 2006; Le Pogam et al., 1998; Murphy et al., 2007; Thomas et al., 2007). Secondly, for many of the isolates in Africa only short segments of sequences are available (Markov et al., 2009). This has limited their analysis and hampered their comparison to other isolates. Greater efforts are needed for more extensive surveys in

Table 3. Genotype 2 subtypes in different countries

<table>
<thead>
<tr>
<th>Location</th>
<th>No. (subtypes)*</th>
<th>No. of isolates</th>
<th>Subtotal (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Europe</td>
<td>17 (a, b, c, d, f, g, h, i, j, k, l, m, n, o, p, q, r) (2?)</td>
<td>724</td>
<td>1020 (52.5)</td>
</tr>
<tr>
<td>North America</td>
<td>11 (a, b, c, d, e, i, j, k, l, m, r) (2?)</td>
<td>207</td>
<td>254 (12.6)</td>
</tr>
<tr>
<td>Asia</td>
<td>9 (a, b, c, f, i, k, 1b/2k, 2i/6p) (2?)</td>
<td>510</td>
<td>540 (26.7)</td>
</tr>
<tr>
<td>Africa</td>
<td>6 (b, c, d, g, h, k) (2?)</td>
<td>18</td>
<td>125 (6.19)</td>
</tr>
<tr>
<td>South America</td>
<td>4 (a, b, c, a/c) (2?)</td>
<td>53</td>
<td>69 (3.41)</td>
</tr>
<tr>
<td>Middle East</td>
<td>3 (a, b, k) (2?)</td>
<td>5</td>
<td>9 (0.445)</td>
</tr>
<tr>
<td>Australia–Oceania</td>
<td>2 (a, b)</td>
<td>4</td>
<td>4 (0.198)</td>
</tr>
<tr>
<td>Total</td>
<td>2021</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*The first row for each location shows the number of total assigned subtypes found in the location; the second row (2?) for each location represents isolates that are classified into genotype 2, but the subtype has not yet been assigned.
Fig. 4. Analysis of p7 sequences. (a) Phylogeny. p7 sequences from the 11 isolates in this study are marked ($\bullet$). The insertion of isolate QC64 is indicated ($\bigcirc$). Subtypes 1a, 1b, 1c, 2a, 2b, 2c, 2i, 2k, 3a, 3b, 3k, 4a, 4f, 5a, 6a, 6i and 7a were included as references. Bar, 0.1 nucleotide substitutions per site. (b) Multiple sequence alignment. The consensus sequence and distinct amino acids are shown in standard International Union of Pure and Applied Chemistry (IUPAC) codes. Dashes indicate gaps relative to the 2e QC64 insertion sequence. The isolate names are listed to the right of the alignment and the QC64 insertion is indicated in grey.
these African countries. Optimally, migration analyses will supply more accurate estimates.

One of the features observed among all genotype 2 sequences is the unique insertion of 20 aa near the end of the NS5A region. In this study, certain isolates also showed insertions in the ISDR and the PKR-BD. Although controversial, mutations in these latter regions have been associated with sensitivity to interferon treatment (Enomoto & Sato, 1995; Gale & Katze, 1998; Gale et al., 1998; Taylor et al., 1999). Another unique and striking insertion was found in the p7 region of the QC64 isolate, which contained 67 aa directly following the end of the p7 protein. Interestingly, the insertion and the p7 protein shared 78% amino acid similarity and both were very different from all other isolates in the Los Alamos HCV database. A survey of additional isolates is required to see whether the insertion is common among other 2e viruses. The p7 protein is found to be critical for HCV infectivity and contains functionally important genotype-specific elements (Sakai et al., 2003). Repetition of this element may have the effect to increase viral infectivity and could be tested using the JFH1 cell culture model.

METHODS

Samples. Serum samples were obtained from 11 patients (four male and seven female) living in Quebec, Canada, between November 2000 and June 2006 as previously described (Murphy et al., 2007). Of these patients, four were Caucasians, four were African immigrants, two were Asians, and one was Haitian (Table 1). They were aged between 36 and 75 years old when sampled. HCV infection was confirmed by routine virological assays for clinical diagnosis.

PCR strategy. Starting with 100 ng of serum, HCV genomic fragments were amplified using methods described previously (Li et al., 2006, 2009). In brief, RNA was extracted using TriPure (Roche), cDNA converted using random hexamers (Promega) and SuperScript III (Invitrogen) reverse transcriptase, and HCV fragments amplified using conventional PCR.

Using the BioEdit software (Hall, 1999), an alignment of full-length genomic sequences available for HCV genotype 2 subtypes 2a, 2b, 2e, 2k and 2i was made and degenerate primers were chosen according to the conserved regions in the alignment. A first set of fragments were amplified and sequenced using these degenerate primers. Specific primers were then designed based on the sequences of the fragments. These primers in combination with additional primers, degenerate or specific, were used to amplify the missing regions. Such a procedure continued until fragments covering the full HCV genome length were obtained. This strategy has been referred as ‘DNA walking through bridges on islands’ and has been successfully used to characterize many genotype 4 and 6 variants (Lu et al., 2007; Li et al., 2006, 2009). The degenerate primers used in this study are listed in Table S1 while the specific primers are not shown.

5’ and 3’ ends. Because the extreme 5’ end of the HCV genome is highly conserved, the 25 nt at the beginning of the 5’ UTR of strain JFH1 were used as a primer in conventional PCR for amplifying the 5’ ends of all the 11 genotype 2 isolates. However, amplification of the 3’ ends employed a nested RACE PCR. The upstream primers were specific to the 3’ ends of the respective NS5B regions while the downstream primer (9587R, TTTACAGCTAGCCGTCGACATG) was based on the highly conserved 98 nt 3’ X tail, corresponding to nt 9587–9565 in the H77 genome. For isolates failing in this amplification, poly A and nested universal primer (NUP) adapter primers were used instead (Li et al., 2009). PCR was conducted for 35 cycles each consisting of 94 °C for 30 s, 55 °C for 30 s, and 72 °C for 30 s except for the first cycle for which the denaturation step was conducted at 94 °C for 3 min. This was followed by a final cycle of extension at 72 °C for 7 min. FastStart Taq DNA Polymerase (Roche) was used as the reagent and standard procedures were taken to prevent false RT-PCR positives (Kwok & Higuchi, 1989).

Sequence purification. Amplicons were purified with the QIAquick PCR purification kit (Qiagen) according to the manufacturer’s protocol and directly sequenced in both directions to confirm the genetic heterogeneity observed at numerous positions throughout the genome. Amplicons obtained with degenerate primers were cloned with pGEM-T Easy Vector (Promega) and at least four clones were sequenced. The sequencing jobs were done using the ABI Prism BigDye 3.0 terminators and an appropriate primer on an ABI Prism 3137 genetic analyser (Applied Biosystems). All observed ambiguous nucleotides were recorded using standard IUPAC codes. ABI raw files were transferred onto a work station where chromatograms were visually inspected, manually edited and automatically assembled using the Seqman program of the LaserGene software (DNASTar, Inc.). After the chromatograms of all amplicons successfully overlapped the whole HCV genome, the sequence job was cleared and the full-length consensus sequence was compiled.

Sequence analysis. The 11 full-length HCV genomes were denoted according to the numbering of the H77 isolate from the extreme 5’ end to the 3’ X tail. Their amino acid sequences were deduced with the EditSeq program of the LaserGene 8.1 software. To analyse these sequences, many reference sequences were retrieved. They were contained in three datasets. The complete sequence dataset included 19 full-length genome sequences, not only from genotype 2, but also from genotypes 1, 3, 4, 5, 6 and 7. The complete sequence dataset for genotype 2 included 58 full-length sequences all from genotype 2. The partial NS5B sequence dataset included 68 partial NS5B sequences, spanning 340 nt and corresponding to nt 8276–8615 in the H77 genome. The three datasets of sequences were aligned using BioEdit software and checked with MEGA5 editor (Tamura et al., 2011) to display the genomic organization and genetic differences, with a particular interest in the amino acids of the interferon-sensitivity determining region (ISDR) (Enomoto & Sato, 1995), the RNA (dsRNA)-activated protein kinase (PKR) region, and additional domains in the 5’ UTR, E2 and NS5A regions (Gale & Katze, 1998; Gale et al., 1998; Taylor et al., 1999).

Recently BEAST software, in which a large number of evolutionary models and tree-based models can be implemented, has been widely used for Bayesian analysis of molecular sequences related by a phylogenetic tree. These models are suitable for analysing both within- and between-species sequence data. Possibly the most distinguishing feature is explicitly modelling the rate of molecular evolution on each branch in the phylogenetic tree rooted by an incorporated timescale. However, an unavoidable burden of this analysis is that, if no obvious prior distribution of a parameter exists, the prior distribution selected will intentionally influence the posterior distribution of the parameters of interest (Drummond & Rambaut, 2007). While the majority of genotype 2 sequences analysed here do not have their sampling dates recorded, in BEAST analysis we would have to include an estimate of a prior distribution, which would create uncertainties and bias. Therefore, phylogenetic trees in this study are better estimated by the traditional method.

Evolutionary analyses of viral ancestral relationships were conducted with MEGA5 software (Tamura et al., 2007). The evolutionary history was inferred by using the maximum-likelihood (ML) method based on the Hasegawa–Kishino–Yano nucleotide-substitution model (Hasegawa et al., 1985) under six categories of the gamma-distribution among-site rate variation (HKY + I + F,0) (Posada & Crandall, 1998). The
transformation/transversion ratio, proportion of invariable sites and gamma distribution shape parameter were estimated from the real sequence datasets. Base frequencies were adjusted to maximize the likelihood. Bootstrap resampling tests were performed 500 times.

To exclude recent virus recombination events (Kalina et al., 2002, 2004; Colina et al., 2004; Legrand-Abraham et al., 2007; Noppornpanth et al., 2006; Lee et al., 2010), the RDP3 software (Martin et al., 2005) was run with default settings modified as follows: (i) window size was 40 nt; (ii) the option of linear sequences was chosen; (iii) six different methods (RDP, GENECOV, MaxChi, Bootscan, Chimaera and SiScan) were run simultaneously against the sequences listed; and (iv) only the events detected by more than two of the above methods were considered positive. This analysis was only performed for the complete genome sequence dataset.

Finally, to study the geographical distribution of genotype 2 viruses, a total of 4044 genotype 2 sequences were analysed. These sequences were retrieved from the Los Alamos HCV database (accessed at the end of 2010) and filtered with respect to their isolation time, sampling countries, genomic region and clonal status, which distinguished 2021 individual isolates. To clarify their diversity with regard to geographical distributions, we tabulated the information according to their subtypes, sampling countries and sampling continents.

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