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<td>Zheng, Wei-Yun</td>
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GENOTYPE IDENTIFICATION OF HEPATITIS C VIRUS (HCV) ISOLATED FROM A SINGLE JAPANESE CARRIER IN NAGASAKI PREFECTURE AND GENOME ANALYSIS OF E1 AND E2/NS1 ENVELOPE GLYCOPROTEIN REGIONS

WEI-YUN ZHENG

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Abstract: The nucleotide (nt) sequence of a Hepatitis C virus genome (HCV-N) which was derived from a single Japanese patient's serum in Nagasaki Prefecture has been determined by multiple clones covering 22 overlapping regions of the HCV genome. The sequenced region consisted of 9295 nt, including 248 nt of 5'-untranslated region (UTR), a single large open reading frame (ORF) encoding a polyprotein of 3010 amino acids (aa) and a 17 nt of 3'-UTR. Phylogenetic analysis indicated that HCV-N belongs to II/1b genotype of group 1. Two other Nagasaki HCV strains (HCV-N1 and HCV-N2) were also sequenced in the E1 and N-terminus of the E2/NS1 regions. Two hypervariable regions (HVR1 and HVR2) were found in the N-terminus of E2/NS1 region among 3 Nagasaki strains and 7 other HCV strains with published sequences. Two well-conserved aa sequences were also identified among 10 HCV strains in the E1 and N-terminus of the E2/NS1 regions. The results will be useful for future understanding on the pathogenesis, virological diagnosis and development of vaccine for HCV.

INTRODUCTION

Hepatitis C virus (HCV) is the major cause of nonA-nonB (NANB) hepatitis, and chronic infection with HCV has been linked to the development of liver cirrhosis and hepatocellular carcinoma (Plagemann, 1991). Many entire and partial HCV sequence informations published so far have shown that HCV genome consisted of about 10 Kb single-stranded positive-sense RNA with 5'UTR, a single long ORF followed by 3'UTR. The viral genome organization resembles that of the flaviviruses and pestiviruses (Kato et al., 1990; Plagemann, 1991).

Since Choo et al. (1989) cloned the genome of an RNA virus from the plasma of a chimpanzee inoculated with plasma from patient with NANB hepatitis and designated it as HCV, entire sequences have been reported for at least 15 HCV strains. There are: HCV-J (Kato et al., 1990), HCV-1 (Choo et al., 1991), HCV-H (Inchauspe et al., 1991), HCV-BK (Takamizawa et al., 1991), HC-J6 (Okamoto et al., 1991), HC-J1 (Okamoto et al., 1992a), HC-J8 (Okamoto et al., 1992b), HC-J4/83, HC-J4/91 (Okamoto et al., 1992c), HCV-JT (Tanaka et al., 1992), HCV-T (Chen et al., 1992), HCV-JK1 (Honda et al., 1993), HC-C2 (Wang et al., 1993), HC-G9 (Okamoto et al., 1994) and NZL1 (Sakamoto et al., 1994). Depending on the HCV sequence similarity, HCV genome can be classified into 6 genotypes: I, II, III, IV, V and VI as reviewed by Sakamoto et al. (1994), or into 6 major genotypes: 1 (a, b, c), 2 (a, b, c), 3 (a, b), 4, 5 and 6 (Simmonds et al., 1993b).

Recently, the newest classification has been summarized by Sakamoto et al. (1994) who classified HCV genome into 3 major groups with entire published sequences. The entire HCV sequence of Japanese strains have been assigned either to group 1 (I/1a genotype: HC-J1; II/1b genotype: HCV-J, HCV-BK, HC-J4/83, HC-J4/91, HCV-JT and HCV-JK1) or group 2 (III/2a genotype: HC-J6; IV/2b genotype: HCV-J8) according to Sakamoto et al. (1994). In order to know whether there is particular HCV in the local area of Nagasaki, Japan, HCV patient serum which came from Nagasaki Prefecture was obtained and the genome sequence of the HCV-N in this serum was determined by multiple overlapping clones.

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In order to see the genetic variability of HCV in the Nagasaki area, HCV genome sequence in 2 more HCV patient sera in Nagasaki Prefecture (HCV-N1 and HCV-N2) were also analyzed similarly in the E1 and N-terminus of the E2/NS1 envelope glycoprotein regions. Two hypervariable regions have been found among these 3 Nagasaki HCV strains using the sequence diversity comparison with published sequence data of 7 HCV patient sera in Nagasaki Prefecture.

**Materials and Methods**

**RNA extraction from HCV patient serum**

Serum samples were obtained from NANB patients in Nagasaki Prefecture which were kindly provided by the Second Department of Internal Medicine of Nagasaki University Hospital. These patients were confirmed to be infected with HCV by anti-C100 HCV ELISA Kit (Ortho Diagnostic Systems, Tokyo, Japan) and HCV reverse-transcription polymerase chain reaction (RT-PCR) as published by Kurihara (1992). One hundred microliters of HCV patient serum was mixed with 20 μl of 10% sodium dodecyl sulfate (SDS) and 80 μl of sterile distilled H2O at room temperature for 5 min. HCV RNA was extracted with phenol / chloroform and precipitated with 3 volumes of ice-cold absolute ethanol. After storage at -80°C for 1hr, HCV RNA was pelleted in an Eppendorf centrifuge with 15,000xg at 4°C for 30 min. The pellet was washed once in 70% ethanol, vacuum dried and dissolved in 100 μl sterile distilled water. RNA solution was stored at -80°C.

**Selection and synthesis of oligonucleotide primers**

Oligonucleotide primer sequences were selected based on the published sequence data (Choo et al., 1991, Takamizawa et al., 1991) and gene walking method. Oligonucleotide primers were synthesized by Applied Biosystems Model 392 DNA / RNA Synthesizer and confirmed for their purity by ion-exchange gel chromatography (Gen-pack; Waters).

**RT-PCR**

Ten microliters of RNA solution were added with 90 μl of RT-PCR mixture [100 pmol of each primers, 0.2 mM deoxynucleoside triphosphate, 10 mM Tris (pH 8.9), 1.5 mM MgCl2, 80 mM KCl, 0.5 mg of bovine serum albumin per ml, 0.1% sodium cholate, 0.1% Triton X-100, 10 U of reverse transcriptase (Life Science Inc.) and 2U of Tth DNA polymerase, a thermostable DNA polymerase (Toyobo Co.).] The reaction mixture was covered by 2 drops of mineral oil and incubated for 10 min at 53°C for RT. PCR amplification (94°C for 60 sec, 53°C for 90 sec and 72°C for 120 sec by thermal cycler; Iwaki Co.) was started immediately after RT and repeated 35 times. cDNA product was subjected to agarose gel electrophoresis and visualized by ethidium bromide staining.

**Cloning and sequencing of HCV cDNA product**

The amplified HCV cDNA was excised from agarose gel, phosphorylated with T4 polynucleotide kinase (Nippon Gene Co.) and blunted with T4 DNA polymerase (Takara Co.). The modified cDNA fragment was ligated into Smal site of pUC19 and transformed into *Escherichia coli* JM 109 strain. The recombinant pUC19 carrying inserted cDNA fragment was purified with Wizard™ Minipreps DNA Purification System (U.S. A.). The cDNA fragment sequence was determined in both directions with sense and antisense primers by dideoxy chain termination method using both 35S radi-isotope-labeling (DNA Sequencing Kit Version 2.0, U.S. A.) and fluorescent dye-labeled DNA sequencing system (373A DNA Sequencer, Applied Biosystems). To avoid sequence variability, 3 colonies from each RT-PCR product were isolated independently for nucleotide sequence determination.

Analysis and homology comparison in nucleotide and deduced amino acid sequences were carried out for HCV-N and 7 other HCV strains with published entire sequence, using a computer system with DNAISIS Mac Version 2.2, NEW CD2 system (Hitachi Software Engineering Co., Ltd, 1992).

**Results**

**The genome characterization of HCV-N**

Multiple cDNA clones were isolated from altogether 22 overlapping regions which covered almost entire genome of HCV-N, except extreme 5' and 3' terminals (Fig. 1). For each of the overlapping region, 3 independent cDNA clones were isolated from the RT-PCR product and sequenced. The cleavage site of the polyprotein coded by the ORF of the HCV-N genome was assigned according to the publications by Okamoto et al. (1992b) and Honda et al. (1993). Total length of the sequenced region of HCV-N consisted of 9295 nt and 3010 aa, respectively (Fig. 2). The sequenced region can be divided into 248 nt in the 5'UTR, 9030 nt in a single ORF and 17 nt in the 3'UTR, respectively. The ORF was considered to be translated into C (191 aa), E1 (192 aa), E2/NS1 (346 aa), NS2 (277 aa), NS3 (609 aa), NS4 (398 aa) and NS5 (997 aa) proteins.

**Genotype classification and genome homology comparison among HCV-N and 7 other HCV strains with published sequences**
Entire sequence informations have been published for the following 7 HCV strains: HCV-1 (Choo et al., 1991), HCV-BK (Takamizawa et al., 1991), HCV-J (Kato et al., 1990), HC-J6 (Okamoto et al., 1991), HC-J8 (Okamoto et al., 1992b), HCV-JT (Tanaka et al., 1992) and HCV-T (Chen et al., 1992). Homology comparison between HCV-N and these 7 HCV strains was shown in Table 1. The most conserved region is the 5'UTR which showed nt homology more than 91.5% among 8 HCV isolates. The sequence homology of the C protein region was also highly conserved (nt > 81.1%, aa > 88.5%). In the putative E1 and E2/NS1 regions, the HCV-N showed high homology with HCV-BK, HCV-J, HCV-JT and HCV-T (nt > 84.5%, aa > 85.8%). Whereas HCV-N showed low homology with HC-J6 and HC-J8 (nt < 68.8%, aa < 72.3%). Nt sequence homology between HCV-N and HCV-1 in the E1 and E2/NS1 regions was 73.7%, 74.5% and aa sequence homology was 76.0%, 81.2%, respectively. From NS2 to NS5 regions, it was evident that HCV-N is closer to HCV-BK, HCV-J, HCV-JT and HCV-T and remote from HC-J6 and HC-J8, while HCV-1 seems to be located intermediate. The total sequence homology comparison among 8 HCV isolates gave us a conclusion that HCV-N belongs to HCV-BK/HCV-J/HCV-JT/HCV-T genotype. In contrast, HCV-N is remote from J6 and J8 genotypes.

Table 1 Homology comparison of nucleotide and deduced amino acid sequence among 8 HCV strains (HCV-N, HCV-1, HCV-BK, HCV-J, HCV-JT, HCV-T, HC-J6 and HC-J8). The homology is indicated by %. The nucleotide sequence homology is shown in upper and amino acid sequence homology is shown in lower with parenthesis, respectively.

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<td>91.5 (93.7)</td>
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<tr>
<td>C</td>
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<td>59.5 (71.8)</td>
<td>59.2 (71.5)</td>
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Figure 2 Nucleotide and deduced amino acid sequence of HCV-N strain. The nucleotide sequence of HCV-N is shown in the upper line and deduced amino acid sequence of HCV-N is shown in the lower line, respectively.
respectively. The sequence homology of HCV-N also showed its closer relationship to HCV-1 than to HC-J6 and HC-J8 isolates (Table 1). HC-J8 possessed 23% divergence with HC-J6 and has been clearly identified as a separate genotype from HC-J6 (Okamoto et al., 1992b). In order to confirm this conclusion, phylogenetic trees were constructed base on nucleotide

Figure 2 Phylogenetic trees are based on nucleotide divergence (%). Sequence of all 8 HCV strains were compared and analyzed by the nearest neighbor method.

Figure 3
divergence throughout entire sequence using nearest neighbor method (Williams and Lance 1977) (Fig. 3). This result indicated that 8 HCV strains can be classified into 2 groups: the group 1 contains genotype II/1b (HCV-N, HCV-BK, HCV-J, HCV-JT, HCV-T) and genotype I/1a (HCV-1); while group 2 contains genotype III/2a (HC-J6) and genotype IV/2b (HC-J8).

Sequence variation in the E1 and N-terminus of E2/NS1 envelope glycoprotein regions among 3 HCV Nagasaki strains and 7 other HCV strains with published sequences

Recently, many papers reported that hypervariable regions existed in the N-terminus of E2/NS1 envelope glycoprotein region of HCV genome (Hijioka et al., 1991; Weiner et al., 1991; Honda et al., 1993). In order to know the HCV sequence diversity of the envelope glycoprotein in local area of Nagasaki, 2 other HCV Nagasaki strains (HCV-N1 and HCV-N2) were also cloned and sequenced for the region of nt 623-1988, aa 126-580 (nt and aa base on the HCV-N number).

Alignment comparison of deduced aa sequence among 3 Nagasaki strains and other 7 HCV strains with published sequences were shown in Fig. 4. Two hypervariable regions (HVR) were discovered in the N-terminus of E2/NS1 region. HVR 1 (aa 384-411) was located directly downstream at the beginning of the E2/NS1, whereas HVR 2 (aa 475-480) was observed 64 aa downstream from the HVR 1. In the HVR 1 consisting of 28 aa, 10 aa residues were well-conserved among 3 Nagasaki strains. Whereas only 3 aa were conserved in this

[Figure 4 Alignment of amino acid sequence among 10 HCV strains (HCV-N, HCV-N1, HCV-N2, HCV-1, HCV-BK, HCV-J, HCV-JT, HCV-T, HC-J6 and HC-J8) in the E1 and N-terminus of E2/NS1 regions (aa 126-580). Two hypervariable regions (HVR 1 and HVR 2) are shown by boxes. Potential N-glycosylation sites (N-X-S/T) are underlined.]


Table 2  HCV genotype classification including HCV-N from this study and 15 entire HCV strains which have been reported by Sakamoto et al. (1994).

<table>
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<th>Group</th>
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<td>Genotype</td>
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<td>II/1b</td>
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<td>HCV-N</td>
<td>HCV-J</td>
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region when a total 10 HCV strains were compared. In the HVR 2 consisting of 6 aa, none of the aa were conserved among 3 Nagasaki strains. This hypervariability was also observed when a total 10 HCV strains were compared simultaneously. In the E1 and N-terminal of the E2/NS1 regions, 2 well conserved aa sequences were observed among 10 HCV strains: G-H-R-M-A-W-D-M-M (aa 315-323) and W-F-G-C-T-W-M-N (aa 549-556). Although 14 N-glycosylation sites (N-X-S/T) were identified in HCV-N, one of them (NNS at aa No. 475-477) was unique to this strain and not present in 2 other Nagasaki strains and 7 other HCV strains with published sequences. Instead, HCV-N1 and HCV-N2 possessed another N-glycosylation site (NFS or NSS at aa No. 250-252). This glycosylation site is found also in HCV-BK, HCV-J, HCV-JT, and HCV-T (Fig. 4). Altogether, only 9 N-glycosylation sites were conserved in the E1 and N-terminal of the E2/NS1 regions among 10 HCV strains shown in Fig. 4.

**DISCUSSION**

Relationship between genotype and geographic area of HCV strains including HCV-N was shown by their sequence comparison using the classification by Sakamoto et al. (1994) (Table 2). The sequence homology indicated that HCV-N strain did not possess its local genotypic character, and similar to HCV-BK, HCV-J and HCV-JT strain which were isolated from other areas of Japan. All these strains apparently belong to the same genotype II/1b of group 1. This genotype also included HC-J4/83; HC-J4/91; HCV-JK1 of Japanese HCV strains, HC-C2 of Chinese HCV strain and HCV-T of Taiwan strain. The genotype 1c of group 1 includes HC-G9 which was derived from Indonesian strain. While, the genotype I/1a of group 1 includes HCV-1 and HCV-H which were derived from American strains. Although another HC-J1 was Japanese strain, it was supposed to have originated from the US, because HC-J1 was derived from a Japanese haemophiliac who developed hepatitis C after receiving US-made factor VIII (Okamoto et al., 1992a). In group 2, HC-J6 and HC-J8 belong to the III/2a and the IV/2b genotype, respectively. Both of them were Japanese strains. In group 3, NZL1 that came from New Zealand belongs to V/3a genotype (Table 2). Simmonds et al. (1993b) have classified 6 major genotypes of HCV from 76 HCV isolates, which were almost worldwide collection, using phylogenetic analysis of the NS5 region. The 76 HCV isolates contained entire sequence of HCV-1, HCV-H, HCV-J, HCV-BK, HCV-T, HCV-JT, HC-J6 and HC-J8 strains and other partial sequence of HCV strains. From this classification, only genotype 4, 5 and 6 showed highly restricted geographical distributions, being apparently confined to Egypt, South Africa and Hong Kong respectively (Simmonds et al., 1993b).

Two hypervariable regions have already been observed in the N-terminal of the E2/NS1 region. The number of conserved aa in the HVR 1 among 3 Nagasaki strains were higher than among a total 10 strains including 3 Nagasaki strains and 7 other HCV strains with published sequences. The numbers of conserved aa (5 aa/28 aa) among HCV strains of genogroup 1 (HCV-N/HCV-1/HCV-BK/HCV-J/HCV-JT/HCV-T) were higher than those (3 aa/28 aa) among 10 HCV strains which included genogroup 2 (HC-J6 and HC-J8). This result may give us an idea that aa conservation in the HVR 1 of the same genotype or genogroup of HCV are
higher than among the different genotype or genogroup of HCV strains. Therefore, the 3 Nagasaki HCV strains may have been originated from the same ancestor.

In contrast, none of the aa was conserved among 3 Nagasaki strains nor among a total 10 HCV strains in the HVR 2 which showed higher variability than the HVR 1. The reason why mutations occur so frequently in only limited regions such as HVR is not known (Tanaka et al., 1992). The higher degree of divergence in HCV E2/NS1 region might reflect the immune selection and suggest that this region cannot probably be an ideal target for future vaccine development.

Regarding the N-glycosylation sites in the E1 and N-terminus of the E2/NS1 region, only 9 sites were conserved out of the 14 sites seen in HCV-N. Some of the glycosylation sites, therefore, would not be essential for the survival, transmission and maintenance of HCV in nature. On the other hand, 2 well-conserved aa sequences were found in the E1 and N-terminus of the E2/NS1 regions among the 10 HCV strains compared. These conserved sequences may be better targets of vaccine development if they were related with protective immunity.

The nt sequence homology of HCV-N in the 5'UTR shows high conservation comparing with other 7 HCV isolates (nt > 91.5%) and was used as an ideal target for PCR amplification to detect HCV RNA (Okamoto et al., 1990). But recently, some papers reported that several HCV strains have sequence variation in the 5' UTR (Lee et al., 1992; Bukh et al., 1992; Simmonds et al., 1993a). Therefore additional informations would be required to select optimal primer for PCR diagnosis on HCV. The functional motifs of the putative encoded proteins of HCV (Plagemann et al., 1991; Tanaka et al., 1992) have also been found in HCV-N. There are consensus sequences of RNA helicase in the NS3 region: Gly-Ser-Gly-Lys-Ser-Thr (aa 1233-1238) and Gln-Arg-Gly-Arg-Asp-Thr-Gly (aa 1486-1493), while NS5 region possessed consensus sequence of RNA-dependent RNA polymerase: Gly-Asp-Asp (aa 2736-2738). The sequence heterogeneity in putative structural proteins and nonstructural proteins of HCV provided significant evidence for genotypic classification. The genetic informations of HCV-N genome obtained in this study will be useful in future understanding on the pathogenesis, diagnosis and development of vaccine for HCV.

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REFERENCES