Hepatitis A virus (HAV) is classified as *Hepatovirus* within the *Picornaviridae*. In virology, HAV is a positively stranded ribonucleic acid (RNA) virus of about 7,500 nucleotides. The viral capsid is composed of three exposed polypeptides VP1, VP2, VP3, and a putative VP4, with a highly conserved antigenic structure. HAV is transmitted through fecal to oral route and is diagnosed by positive serum IgM anti-HAV antibody test. Isolates of HAV are of a single serotype, but human isolates can be categorized into three genotypes (I, II and III) with 2 subgenotypes (A and B for I and III). Genotype I is the most abundant type worldwide, and particularly genotype IA is found in North America, Europe, China, Japan, the former USSR and Thailand. Strains of subgenotype IIIA have been collected from humans infected with HAV in India, Sri Lanka, Nepal, Malaysia and the USA. For the past several years, studies on HAV have not been performed because most people in Korea carry the HAV antibody by natural infection. Once HAV antibodies are acquired, they give life-long immunity. Moreover, hepatitis A infection is usually mild in children, and rarely progresses into chronicity and fulminant hepatitis. However, the high risk for HAV infection in Korea has been focused on young adults and adolescents who, did not get infected in childhood due to improved hygiene measures, and are more prone to infection later in life.
with more serious disease.1,8 In 2001, sporadic acute hepatitis A was reported in Korea, being subgenotype IA.7 In 2007, an outbreak of acute hepatitis A in a Korean hospital was reported, and the subgenotypes of this hepatitis A virus were found to be IA and IB.10 HAV phylogenetic studies can provide important information for the design of appropriate public health cares and HAV genotypic changes. Therefore, we investigated the genotypes of recently isolated HAV cases in the south-east area of Gyeonggi-do in Korea.

**MATERIALS AND METHODS**

**Patients and collection of blood samples**

From June 2004 to June 2006, 46 patients from the Bundang CHA Hospital were identified prospectively. All patients, showing clinical and biochemical signs of acute hepatitis A and serological evidence of acute hepatitis A, were classified as acute hepatitis A patients. The serological evidence was obtained with IgM anti-HAV antibody tests (EIA, VIDAS, BioMérieux, Marcy-l’Etoile, France). Patients lived in the south-east area of Gyeonggi-do (Seongnam-si, Icheon-si, Gwangju-si) in Korea and all were Koreans.

During the two study years, there was no reported outbreak of acute hepatitis A in the study area. Routine contact tracing was done. All patients visited a hospital as an outpatient once a week for two or three weeks. All cases were sporadic acute hepatitis A.

Ten mL of whole blood was collected from the veins of each patient in order to analyse the genotype of HAV. Each sample was numbered in order of date. The whole blood was centrifuged to separate the serum. The serum was preserved at -20°C. For the study, sera of the enrolled patients were numbered from 1 to 46. The study was approved by the local ethical committee and conform to the manufacturer’s instructions. The cloned plasmids were sequenced in ABI PRISM® 3100 Genetic Analyzer (Applied Biosystems, Foster City, CA, USA).

**Direct sequencing**

The target PCR products were re-amplified and extracted from the gel using the GENCLEAN II Kit (Q-BIO gene, Carlsbad, CA, USA) and directly sequenced with ABI PRISM®3100-Avant Genetic Analyzer (Applied Biosystems, Foster City, CA, USA) using a HAV specific primer (5’-GTCTACCAGGCAATTGCATTCTCAT-3’).

**Phylogenetic analysis and genotype determination**

The CLUSTAL W (version 1.83) program was used to generate the phylogenetic tree. The putative VP1/2A junction sequences were amplified and sequenced from clinical samples and compared with the corresponding Gene Bank reference sequences for genotypes. IA : AF234864 (KU97-1), AF234876 (KU98-42), AB205676, L07717 (PRC37). IB : L07700 (Ag11). II : L07693 (CF-53). IIIA : AJ299467 (NOR-24), L07668 (GA76), L07688 (H-122), L07725 (India90), L20530 (A-177). IIIB : L07729 (SLF88). The amino acid alignments of the putative VP1/P2A junction sequences were amplified and sequenced from clinical samples and compared with the corresponding Gene Bank reference sequences for genotypes. IA : AF234864 (KU97-1), AF234876 (KU98-42), AB205676, L07717 (PRC37). IB : L07700 (Ag11). II : L07693 (CF-53). IIIA : AJ299467 (NOR-24), L07668 (GA76), L07688 (H-122), L07725 (India90), L20530 (A-177). IIIB : L07729 (SLF88). The amino acid alignments of the putative VP1/P2A junction in this study submitted to Genebank were numbered: FJ372910 - FJ372970.

**RESULTS**

Forty six patients were enrolled in this study. Among the total 46 patients, 21 were male and 25 were female. The mean age was 31 ± 6 years. The mean period of hospitalization was 11 ± 6 days. The baseline clinical and laboratory results are summarized in Table 1. Most clinical and laboratory results of each genotype did not show differences between the genotypes. Case of co-infection with HBV or HCV was not found. The yearly distribution according to genotype is shown in Table 2.
We found that sera from 41 out of 46 patients were positive for HAV-RNA. As can be seen in the phylogenetic tree, the HAV sequences from 25 patients (60%) out of 41 ones were found to be of subgenotype IIIA, and HAV sequences from 16 patients (40%) were confirmed as subgenotype IA (Fig. 1). Compared to the formerly reported subgenotype IA in Korea,7 the subgenotype IA in our investigation showed > 95% sequence homology.

Several amino acid substitutions were found in subgenotype IA and IIIA. The 2A-19 substitution was found in ten samples of subgenotype IA. The substitution of 2A-19 is identical to fulminant hepatitis of AB020567 in Japan. On the serum No. 37, glutamic acid of 2A-11, on L07725 found in China was substituted by aspartic acid. On the serum No. 26, arginine of 2A-6, on L20530 found in Japan was substituted by glycine.

On serum No. 18 of subgenotype IIIA, valine of VP1-288 was substituted by arginine. On serum No. 45, glutamine of 2A-25 was substituted by arginine. On serum No. 7 and 36, methionine of VP1-276 was substituted by arginine. On serum No. 9, methionine of VP1-276 was substituted by leucine, and aspartatic acid of VP1-283 was substituted by glycine. On serum No. 10, lysine of 2A-18 was substituted by glutamic acid.

**Table 1. Baseline Clinical and Laboratory Results of 46 Hospitalized Patients with Positive IgM Anti HAV Test**

| Variable                      | Total |
|-------------------------------|-------|
| Number                        | 46    |
| Age (yrs)                     | 31 ± 6|
| Period of hospitalization (days) | 11 ± 6|
| AST on admission (IU/L)       | 2383 ± 2523 |
| Peak AST (IU/L)               | 2589 ± 2669 |
| ALT on admission (IU/L)       | 2587 ± 1415 |
| Peak ALT (IU/L)               | 2809 ± 1544 |
| TB on admission (mg/dL)       | 5.1 ± 1.9 |
| Peak TB (mg/dL)               | 6.6 ± 3.3 |
| Platelet (cells/mm³)          | 168000 ± 73000 |
| PT (%)                        | 75 ± 25 |
| Albumin (g/dL)                | 3.7 ± 0.4 |

AST, aspartate aminotransferase; ALT, alanine aminotransferase; HAV, Hepatitis A virus; IgM, immunoglobulin M; PT, prothrombin time; TB, total bilirubin. Values represent mean ± standard deviation.

HAV-RNA, Hepatitis A virus-ribonucleic acid.

**Table 2. Yearly Distribution of HAV-RNA Detected Acute Hepatitis A Patients**

| Genotype | Genotype | Total |
|----------|----------|-------|
| IA       | IIIA     |       |
| June 2004 to June 2005 | 4       | 10    | 14 |
| July 2005 to June 2006 | 12      | 15    | 27 |
| Total    | 16       | 25    | 41 |

HAV-RNA, Hepatitis A virus-ribonucleic acid.

The founding works on HAV genetic variability made use of discrete, selected partial HAV genomic regions, such as the C terminus of VP3 to the N terminus of VP1 or the putative VP1/P2A junction region. HAV genotype is defined as a group of viruses with > 85% nucleotide sequence identity within one of these regions. Genotypes I and III are further subdivided into two distinct subgroups, which differ in sequence in no more than 7.5% of base positions.1,4,11 The subgenotypes, IA and IB have been reported in Korea.3,10 In conclusion, the three genotypes (IA, IB, IIIA) are co-circulating in Korea. It has been reported before
that genetic changes are rare in the HAV genome, which is unlike other RNA viruses.12 Also, nationwide research of HAV genotypic distribution in Korea has not yet been performed. It can be hypothesized that the influx from the other regions of Korea or predominant countries is one of the causes in the subgenotype IIIA circulating in Korea. It was reported that the dominant HAV genotype in India, Sri Lanka, Nepal, and Malaysia is IIIA, even though the research was insufficient.13 Recent reports showed that the HAV subgenotype IIIA in those countries has continuously been the circulating type.14,15 The acute hepatitis A samples gathered from this research area and subgenotype IIIA dominant country need to be compared by further molecular analysis. To verify possible infection from travel, 41 patients, detected with HAV-RNA, were questioned about their travel history. One patient (No. 22, subgenotype IA) had a travel history to Philippines 15 days before hospitalization. Tjon et al.17 reported that HAV subgenotype IIIA was found in homeless, bad hygiene patients and intravenous drug users. However, these trends were not found in this research.

The outbreak of HAV subgenotype IIIA has been reported among intravenous drug users in Europe.16-18 Up to date, HAV has been known to be transmitted through fecal to oral route. However, by the outbreak of the subgenotype IIIA of intravenous drug users and infection of HAV in hemophilia patients by clotting factor,19 it can be possible that HAV is also transmitted by the parenteral route. This change of mode of transmission is likely due to nucleotide variability that reach 4% of the subgenotype IIIA, and it is also possible that HAV has high selection pressure through fecal to oral route and infection through blood has low selection pressure.20,21 The nucleotide variability of the HAV subgenotype IIIA needs to be studied through continuous molecular study in Korea. Through the research, future outbreaks of the subgenotype IIIA in intravenous drug users and blood products can be predicted.

Little information is available on the possible relation between the severity of hepatitis A and the infecting genotype.22,23 It was previously considered that HAV disease severity is linked to individual host factors such as age and underlying liver disease. However, through continuous investigation of HAV, many factors that affect the severity of acute hepatitis A have been proven. First, it has been reported that nucleotide variations in the central portion of the 5' NTR of HAV could affect the severity of acute hepatitis A.24 Low or undetectable HAV viral load and a high bilirubin level could independently affect the progression to fulminant hepatitis. It has also been reported that HAV genotype did not affect the progress to fulminant hepatitis.2,3 The present study found also similar number of subgenotypes IA and IIIA in the same research area, thus making it possible to directly compare their clinical and laboratory results. Most clinical and laboratory results of each genotype did not reveal any differences between the genotypes, but it showed that mean peak total bilirubin (TB) level in subgenotype IIIA is significantly higher (data not shown). Also, the mean TB level on admission showed a high level of tendency on subgenotype IIIA, even though it is not statistically significant. However, difference has limitations, mostly stemming from this study’s small sample size, small area of study and selection of patients. Future work is required to ascertain the difference of total bilirubin according to genotype.

The nucleotide sequence heterogeneity found in our study of the HAV genome of these 41 virus strains results in only limited differences in the amino acid sequence. Among all of the subgenotype IA and IIIA, amino acid sequence differences showed to be maximum 3% (2/56). On the former prevalent subgenotype IA of Korea, it was reported that lysine 2A-10 was substituted by arginine, and glutamine 2A-19 by serine.7 However, our study showed that substitution of 2A-10 did not occur in every sample, and substitution of 2A-19 occurred only in 10 of 16 patients who showed subgenotype IA.

In conclusion, this study examined HAV subgenotype IIIA in Korea, and the results showed that three HAV subgenotypes (IA, IB, IIIA) co-circulate in Korea. The genetic relatedness of HAV from Korea provides valuable new data on the distribution of subgenotype IIIA.

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