Full-length genome of wild-type hepatitis A virus (DL3) isolated in China

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METHODS: A stool specimen was collected from hepatitis A patient from Dalian, China. HAV (DL3) was isolated and viral RNA was extracted. The genome of DL3 was amplified by reverse transcription and polymerase chain reaction (RT-PCR), followed by cloning into pGEM-T vector. The positive colonies were selected and sequenced. The full-length genome of DL3 was analyzed and compared with other wild-type HAV isolates.

RESULTS: The genome of DL3 was 7476 nucleotides (nt) in size, containing 732-nt 5' untranslated region (UTR), 6681-nt open reading frame (ORF) which encoded a polyprotein of 2227 amino acids (aa), and 63-nt 3' UTR. The base composition of 2227 amino acids, a short 3' UTR with a polyA tail. The HAV polyprotein is co- and posttranslationally cleaved into smaller structural proteins (VP1, VP2, VP3 and a putative VP4) and nonstructural proteins (2A, 2B, 2C, 3A, 3B, 3C and 3D) by the virus-encoding proteinase.

Human isolates of HAV possess a single serotype, and monoclonal antibodies raised to various isolates of human HAV have failed to distinguish between individual isolates. However, the nucleotide sequencing of selected genome regions that encode the putative VP1/2A junction region of wild-type HAV isolates present in human specimens has demonstrated significant sequence heterogeneity. Using this approach, HAV isolates could be differentiated into seven unique genotypes based on the sequence of the VP1/2A junction region. A genotype is defined as a group of viruses that differed from each other by no more than 15%, and a subgenotype as a group of viruses with >92.5% nucleotide sequence identity.

Until now, complete nucleotide sequences of eleven different human wild-type HAV isolates and partial nucleotide sequences of wild-type isolates or cell-adaptive variants of HAV have been reported. These isolates were isolated from hepatitis A patients in diverse geographic origins, including partial sequence of the isolates from fulminant hepatitis A in Shanghai in 1988. But the complete nucleotide sequence of genome of the isolate from China remains unknown. In order to elucidate the genetic characteristics and molecular epidemiology and evolution of wild-type HAV in China, we determined the complete nucleotide sequence of genome of an acute HAV isolate (DL3) from Dalian, and compared the complete genome sequences and deduced amino acid sequence of isolate DL3 with those of other eleven wild-type HAV isolates. There are significant differences and identities among the twelve independent isolates.

INTRODUCTION

Hepatitis A virus (HAV) is an important human pathogen causing hepatitis, with a higher incidence in developing countries than in developed countries. Direct person-to-person spread by the fecal/oral route is the most important means of transmission of hepatitis A, and infection with HAV can cause sporadic and epidemic acute hepatitis in humans. Although improvements in sanitation have led to a significant reduction in the endemicity of hepatitis A virus infection, hepatitis A is still the most common viral hepatitis infection and a cause of substantial morbidity in China.

HAV is classified as one of two members of the genus Hepatovirus within the family Picornaviridae. HAV virion is a naked, spherical particle with a diameter of 27-32 nm. The virion consists of a genome of a linear, single-stranded, 7.5-kb positive-sense RNA and of a protein shell made up of three major proteins, VP1-VP3. The genome can be divided into a long 5' terminal untranslated region (5'UTR) of about 735 nucleotides, a large open reading frame encoding a polyprotein of 2227 amino acids, a short 3' UTR with a polyA tail. The HAV polyprotein is co- and posttranslationally cleaved into smaller structural proteins (VP1, VP2, VP3 and a putative VP4) and nonstructural proteins (2A, 2B, 2C, 3A, 3B, 3C and 3D) by the virus-encoding proteinase.

Human isolates of HAV possess a single serotype, and monoclonal antibodies raised to various isolates of human HAV have failed to distinguish between individual isolates. However, the nucleotide sequencing of selected genome regions that encode the putative VP1/2A junction region of wild-type HAV isolates present in human specimens has demonstrated significant sequence heterogeneity. Using this approach, HAV isolates could be differentiated into seven unique genotypes based on the sequence of the VP1/2A junction region. A genotype is defined as a group of viruses that differed from each other by no more than 15%, and a subgenotype as a group of viruses with >92.5% nucleotide sequence identity.

Until now, complete nucleotide sequences of eleven different human wild-type HAV isolates and partial nucleotide sequences of wild-type isolates or cell-adaptive variants of HAV have been reported. These isolates were isolated from hepatitis A patients in diverse geographic origins, including partial sequence of the isolates from fulminant hepatitis A in Shanghai in 1988. But the complete nucleotide sequence of genome of the isolate from China remains unknown. In order to elucidate the genetic characteristics and molecular epidemiology and evolution of wild-type HAV in China, we determined the complete nucleotide sequence of genome of an acute HAV isolate (DL3) from Dalian, and compared the complete genome sequences and deduced amino acid sequence of isolate DL3 with those of other eleven wild-type HAV isolates. There are significant differences and identities among the twelve independent isolates.

MATERIALS AND METHODS

Virus sample

Wild-type human HAV isolate DL3 was recovered from stool specimens stored at -70°C from an patient with acute hepatitis A, infected during an outbreak at Dalian, Liaoning Province, and was named as DL3. HAV contained in stool were identified by ELISA, immune precipitation and immune electron microscopy. The concentrated and purified HAV for RT-PCR were prepared by chloroform extraction for three times from 10% of stool supernatant, followed by discontinuous sucrose/glycerol density gradient ultracentrifugation.
Table 1. The primers used for amplification of DL3 genomic RNA

| Clones | Primers       | Sequences                          |
|--------|---------------|------------------------------------|
| A (0.5kb) | 5'RACE RT-primer | 5'-PGCAGAATGAATC-3                 |
|        | 5'RACE A1 | 5'-CTTCGACAAGCACTCCAAGG-3          |
|        | 5'RACE S1 | 5'-TCTTCCTCTCAATATCTGCC-3          |
|        | 5'RACE A2 | 5'-TGTTCTTCCTCAATATCTGCC-3         |
|        | 5'RACE S2 | 5'-CTGGAATAAATTCCTCCTGGCC-3        |
| B (0.5kb) | B1 | 5'-GCTTGAAGCTCTCAAGGGCC-3          |
|        | B2 | 5'-AGATTAAACAGTCAGAGGTC-3          |
| C (1.1kb) | C1 | 5'-ACATCATCCAAGGTCAGGAGG-3         |
|        | C2 | 5'-ATCCATACGTGATAAGAGG-3           |
| D (1.1kb) | D1 | 5'-CTCGAGTTTCGAGACCTCC-3           |
|        | D2 | 5'-CATGGATAACATGGCATTTG-3          |
| E (1.1kb) | E1 | 5'-TCTAGCAGCAAGCAATCATGAG-3        |
|        | E2 | 5'-AAATCCTCTGAAACTTAGCAGTC-3       |
| F (1.2kb) | F1 | 5'-TCCCCAGATGAGCTAGGAG-3           |
|        | F2 | 5'-GCTTGAAGCTCTCAAGGGCC-3          |
| G (1.2kb) | G1 | 5'-AGTCTCTCTAGTAAGCAGTTCG-3        |
|        | G2 | 5'-GCTCGAGTTTCGAGACCTCC-3          |
| H (1.1kb) | H1 | 5'-AATCATCAAGGTCAGGAGG-3           |
|        | H2 | 5'-ATCCATACGTGATAAGAGG-3           |
| I (0.5kb) | I1 | 5'-GATTCTCTTGGTGGAGTGC-3           |
|        | I2 | 5'-CTTCTCTTGGTGGAGTGC-3            |

**DNA sequencing and analysis**

Sequencing strategy of genome of DL3 was showed in Figure 1. Oligonucleotide primers specific for HAV and primers corresponding to the T7/SP6 promoter region of pGEM®-T Vector were used to sequence the inserted and identified HAV fragment. A Taq DyeDexy Terminator Cycle sequencing kit and a 377 DNA sequencer (Perkin Elmer) were used to determine nucleotide sequences. To eliminate the possibility of errors in the sequence due to Taq polymerase for PCR, at least three clones of each amplified fragment, derived from two individual PCR products, were sequenced. Also to correctly determine the sequence of extreme 5’ terminus of HAV genome, a 5’ RACE reaction[22] was used to obtain a cDNA fragment from 5’ UTR of genome with 5’-Full RACE Core Set (TaKaRa). Analysis, alignment and translation in the amino acids of the obtained nucleotide sequences were done using the sequence analysis program OMEGA2.0 (Oxford Molecular).

**Figure 1** Sequencing strategy of genome of DL3.

**Phylogenetic analysis**

Multiple alignments of genome sequences of twelve HAV isolates were done using the Clustal W Program[21]. Phylogenetic tree was calculated from genomes to determine the association of DL3 with other eleven wild-type HAV isolates with Vector NTI Suite6.0 software using the Neighbor Joining method[24].

**Reference isolates**

The accession numbers of the sequences included in the analysis were as follows: LA, K02990; HM175, M14707; MBM, M20273; GBM, X75215; AH1: AB020564; AH2: AB020564 AB020565; FH1: AB020564 AB020566; FH1: AB020567; FH2: AB020568; FH3: AB020569; SLF88, AY302861.

**RESULTS**

**Complete nucleotide sequence of DL3 genome and deduced amino acids**

The Complete nucleotide sequence of DL3 has been deposited into GenBank under the accession no. AF512536.

The genome was 7 476-nucleotide long and encoded a polyprotein of 2 227 amino acids. The genome contained a 5’ UTR of 732 nucleotides that was two nucleotides shorter than that of HAV HM175/WT. The long open reading frame of genome with 5'-Full RACE Core Set (TaKaRa). Analysis, alignment and translation in the amino acids of the obtained nucleotide sequences were done using the sequence analysis program OMEGA2.0 (Oxford Molecular).

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The 5’ UTR contained three main pyrimidine-rich tracts. The first region, near the 5’ terminus (nucleotide 99–138), had...
a 92.3\% pyrimidine content. The second region (nucleotide 204-250) had a 83\% pyrimidine content. The third tract (nucleotide 711-725) with a 93.9\% pyrimidine content lay immediately before the initiation codon. The 5' UTR had a G+C content of 46.29\% which is higher than that of the genome.

**Comparisons with other isolates**

The complete nucleotide sequence and deduced amino acid sequences of HAV DL3 have been compared with those of reported wild-type isolates (Table 2, Table 3). DL3 had the highest identity of 97.5\% for nt (185 differences) with AH1, but the highest identity of 99.2\% for amino acid (18 differences) with AH2 and FH3. DL3 shared the lowest identities of 85.7\% for nucleotide (1,066 differences) and 96.8\% for amino acid (72 differences) with SLF88. Most of the changes were nucleotide transitions.

**Table 2** Nucleotide identities of full-length genomes between DL3 and reference isolates

|       | AH1 | AH2 | AH3 | FH1 | FH2 | FH3 | GBM | LA  | HM175 | MBB  | SLF88 |
|-------|-----|-----|-----|-----|-----|-----|-----|-----|-------|------|-------|
| Full  | 185/97.5 | 263/96.5 | 307/95.9 | 282/96.2 | 200/97.3 | 199/97.3 | 323/95.7 | 344/95.4 | 642/91.4 | 659/91.2 | 1066/85.7 |
| 5'NTR | 15/98.0 | 29/96.0 | 25/96.6 | 16/97.8 | 13/98.2 | 13/98.2 | 18/97.4 | 14/98.1 | 38/94.8 | 50/93.1 | 84/88.5 |
| VP4   | 5/98.6 | 2/97.1 | 4/94.2 | 2/97.1 | 2/97.1 | 2/97.1 | 3/95.7 | 2/97.1 | 7/89.9 | 7/89.9 | 8/88.4 |
| VP2   | 15/97.7 | 25/96.2 | 28/96.4 | 18/97.3 | 24/96.4 | 39/94.1 | 28/95.8 | 58/91.3 | 58/91.3 | 98/85.3 |
| VP3   | 15/98.0 | 25/96.6 | 28/96.2 | 13/98.2 | 23/96.9 | 35/95.3 | 40/94.6 | 67/90.9 | 63/91.5 | 114/84.6 |
| VP1   | 23/97.2 | 27/96.7 | 25/97.0 | 23/97.2 | 13/98.4 | 30/96.4 | 44/94.6 | 76/90.8 | 78/90.5 | 124/84.9 |
| 2A    | 5/97.7 | 10/95.3 | 11/94.8 | 9/95.8 | 3/98.6 | 9/95.8 | 9/95.8 | 12/94.4 | 17/92.0 | 14/93.4 | 27/87.3 |
| 2B    | 13/98.3 | 26/96.5 | 29/96.1 | 46/93.9 | 18/97.6 | 28/96.3 | 33/95.6 | 37/95.1 | 75/90.0 | 73/90.3 | 99/86.9 |
| 2C    | 30/97.0 | 42/95.8 | 42/95.8 | 35/95.6 | 35/96.5 | 52/94.8 | 55/94.5 | 114/88.7 | 120/88.1 | 152/84.9 |
| 3A    | 1/99.5 | 4/98.2 | 4/98.2 | 9/95.9 | 4/98.2 | 3/98.6 | 13/94.1 | 8/96.4 | 14/93.7 | 18/91.9 | 23/89.6 |
| 3B    | 1/98.6 | 1/98.6 | 2/97.1 | 3/95.7 | 3/95.7 | 0 | 2/97.1 | 3/95.7 | 6/91.3 | 7/89.9 | 12/82.6 |
| 3C    | 14/97.9 | 20/97.0 | 24/96.3 | 25/96.2 | 16/97.6 | 21/96.8 | 30/95.4 | 27/95.9 | 56/91.5 | 50/92.4 | 98/85.1 |
| 3D    | 51/96.5 | 51/97.0 | 51/96.5 | 53/96.4 | 52/96.5 | 27/98.2 | 57/96.1 | 64/95.6 | 113/92.3 | 114/92.2 | 209/85.8 |
| 3'NTR | 1/98.4 | 1/98.4 | 1/98.4 | 1/98.4 | 1/98.4 | 1/98.4 | 2/96.8 | 10/84.4 | 1/98.4 | 7/88.9 | 18/71.4 |

**Table 3** Amino acid identities of full-length genomes between DL3 and reference isolates

|       | AH1 | AH2 | AH3 | FH1 | FH2 | FH3 | GBM | LA  | HM175 | MBB  | SLF88 |
|-------|-----|-----|-----|-----|-----|-----|-----|-----|-------|------|-------|
| Full  | 21/99.1 | 18/99.2 | 36/98.4 | 26/98.7 | 21/99.1 | 18/99.2 | 36/98.4 | 34/98.5 | 36/98.4 | 49/97.8 | 72/96.8 |
| VP4   | 1/95.7 | 1/95.7 | 1/95.7 | 1/95.7 | 1/95.7 | 1/95.7 | 1/95.7 | 1/95.7 | 1/95.7 | 1/95.7 | 1/95.7 |
| VP2   | 0 | 0 | 0 | 1/99.5 | 0 | 0 | 0 | 1/99.5 | 0 | 2/99.1 | 2/99.1 |
| VP3   | 1/99.6 | 1/99.6 | 1/99.6 | 2/99.2 | 1/99.6 | 1/99.6 | 1/99.6 | 1/99.6 | 2/99.2 | 2/99.2 | 4/98.4 |
| VP1   | 0 | 0 | 0 | 1/99.6 | 0 | 0 | 1/99.6 | 2/99.3 | 13/95.3 | 1/99.6 | 2/99.3 | 2/99.3 |
| 2A    | 1/98.6 | 1/98.6 | 1/98.6 | 1/98.6 | 1/98.6 | 0 | 0 | 1/98.6 | 0 | 1/98.6 | 2/97.2 |
| 2B    | 2/99.2 | 2/99.2 | 3/98.8 | 5/98.0 | 5/98.0 | 5/98.0 | 3/98.8 | 4/98.4 | 4/98.4 | 6/97.6 | 6/97.6 |
| 2C    | 6/98.2 | 5/98.5 | 4/98.8 | 6/98.2 | 5/98.5 | 4/98.8 | 7/97.9 | 5/98.5 | 11/96.7 | 17/94.9 | 14/95.8 |
| 3A    | 0 | 1/98.6 | 1/98.6 | 1/98.6 | 0 | 0 | 5/93.2 | 1/98.6 | 1/98.6 | 3/95.9 | 1/98.6 |
| 3B    | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1/95.7 | 0 | 0 | 0 |
| 3C    | 2/99.1 | 2/99.1 | 3/98.6 | 2/99.1 | 2/99.1 | 2/99.1 | 3/98.6 | 2/99.1 | 2/99.1 | 2/99.1 | 8/86.3 |
| 3D    | 8/98.4 | 5/99.0 | 22/95.5 | 8/98.4 | 6/98.8 | 4/99.2 | 11/97.8 | 7/98.6 | 10/98.0 | 12/97.5 | 33/93.3 |
5' Untranslated region
The 5’ UTR of HAV forms conserved and highly ordered secondary structure and plays an important role in controlling viral translation[25, 26]. Comparisons with eleven other isolates in 5’ UTR showed that DL3 shared the lowest identity of 88.5 % (84 nucleotide changes) with SLF88, the highest identity of 98.2 % (13 nucleotide changes) with FH2 and FH3. There were a G deletion at position 28 and a T deletion at position 102 compared with HM175 and MBB. In contrast to the comparison with complete nucleotide sequence, identities of 5’ UTR for DL3 with other isolates were higher than that of complete genomes except the identity with AH2, which suggested that the 5’ UTR of HAV was conserved. In contrast to ribosomes scanning from the capped 5’ ends of the majority of cellular mRNA, the long and highly structured 5’ UTR of HAV mediated the binding of ribosomal subunits internally at internal ribosome entry site (IRES) of nucleotide 324 to nucleotide 692 (base position was corresponding to HM175/WT) that directed cap-independent initiation of viral translation at correct AUG codon[26]. In this region DL3 shared higher identities of 97.9 %, 98.4 %, 99.2 %, 98.6 %, 98.6 %, 95.9 %, 95.4 % and 94.9 % with AH2, FH1, FH2, FH3, LA, HM175, MBB and SLF88 than that in whole 5’ UTR. But DL3 was identical to SLF88 between nucleotide 223 to nucleotide 371 (base position is corresponding to HM175/WT). The result suggested higher conservative and importance of common secondary structure of IRES in initiation of viral translation.

Coding region
The open reading frame of HAV RNA encoding a large polyprotein with 2,227 amino acids was divided into P1, P2 and P3 region. The P1 region encodes structural or capsid proteins, VP4, VP2, VP3 and VP1. The P2 and P3 region encoded nonstructural proteins, 2A, 2B, and 2C, and 3A, 3B, 3C and 3D, respectively. In coding region, the amino acid differences for the structural and nonstructural proteins of twelve isolates are showed in Table 3. In contrast to the high heterogeneous of nucleotide sequences, the amino acid sequences were highly conserved among twelve wild-type isolates. Comparisons within the coding region (6,681 nucleotides) of DL3 and SLF88 yielded 964 nucleotide changes (14.4 % difference), resulting in only 72 amino acid substitutions (3.2 % difference). In comparison with eleven isolates, DL3 had the highest identity (99.2 %) of amino acids with AH2 and FH3, lowest identity (96.8 %) with SLF88.

P1 region, the structural protein region of HAV, consisted of 2,295 nucleotides and encodes 765-amino acid polypeptide which was processed into four structural proteins, VP4, VP2, VP3 and VP1. The capsid protein of HAV consisted of VP1, VP2 and VP3. The presence of a fourth protein VP4 has been described repeatedly, but the reported apparent molecular weights (7-14KD) contrasted sharply with those predicted from nucleic acid sequence data (1.5 or 2.3KD). Conclusive physical identification of VP4 was still unavailable. The alignment of amino acid sequence indicated that no amino acid mutation of polyprotein of DL3 occurred at cleavage site of VP4/VP2, VP2/VP3, VP3/VP1, implying the importance of conserved cleavage sites in entire viral structure. In VP4 DL3 shared lowest amino acid sequence identities with other isolates in structural region, 91.3 % identity with SLF88, 95.7 % identity with other ten wild-type isolates. DL3 had a unique amino acid mutation (S to A) at position 4 different from all the other isolates. In VP2 region amino acid sequences were highly conserved. There was 100 % identity between DL3 and AH1, AH2, AH3, FH2 and FH3 and LA. And even for DL3 and SLF88, the identity was also up to 99.5 %. In VP3 region DL3 had another unique amino acid mutation (K to R) at position 47 different from all the other isolates. It had been reported that VP1 had the most amino acid diversity of the capsid proteins[16]. But it was not the case for DL3. In VP1 region there was 100 % amino acid sequence identity between DL3 and AH1, AH2, AH3 and FH2. The highest difference was between DL3 and LA (13 amino acid differences, 95.3 % identity), just because one amino acid was inserted into the LA between amino acids 540-548, at which a cluster of 8 amino acids was altered due to 3 changes in the open reading frame resulting from the insertion of 3 nucleotides.

The HAV P2 region encoded nonstructural proteins 2A, 2B and 2C. The proposed 2A region encoded for 71 amino acids, but the exact function of 2A was not yet determined. Recent study revealed that 2A protein might participate in virion morphogenesis[27]. 2B and 2C proteins were found to have 251 and 335 amino acids residues, respectively. Both proteins playing important roles in the replication of viral RNA were considered significant in host-dependent adaptation since many mutations had been detected in both regions of adapted variants[28]. HAV 2C protein was considered to have helicase and NTPase activities. With some variants, multiple mutations in P2 contributed to enhance viral replication with 5’ UTR and P5 proteins and to express the cytopathic phenotype[29]. In 2A, 2B and 2C proteins, 2C protein was the most variable between twelve isolates. DL3 shared the lowest identity of 94.9 % (17 amino acid differences) with MBB and the highest identity of 98.8 % with AH3 and FH3 (only 4 amino acid differences). In contrast, 2B was more conserved. The 99.2 % identities of DL3 with AH1 and AH2 and the 98.8 % identities of DL3 with AH3 and GBM revealed high conservation of 2B protein among wild-type viruses. It has been reported that a little more amino acid substitutions seemed to be found in FH than in AH in 2B region[18]. This research also showed that more amino acid substitutions were found in FH than that in AH when compared with DL3, suggesting the possible association between the severity of hepatitis A and amino acid substitutions in 2B region.

The HAV P3 region encoded 3A, 3B, 3C and 3D proteins with 74, 23, 219 and 489 amino acids, respectively. 3A functioned as pre-Vpg. 3B was a genome-linked viral protein (Vpg). 3C was the sole protease for HAV protein processing. The recombinant HAV 3C prepared in E. coli catalyzed the putative cleavage sites and released mature or intermediate proteins, VP0, VP3, VP1-2A, VP1, 2A, 2B, 2BC, 2C, 3ABC, 3AB, 3C and 3D[30-34]. 3D was an RNA-dependent RNA polymerase. By comparison, 3B protein showed highest amino acid sequence homology, with 100 % identities among eleven isolates except 95.7 % identity between DL3 and GBM. Compared with 3D proteins, 3C protein showed higher amino acid sequence identities in all twelve isolates. In this protein, DL3 shared the 96.3 % identity with SLF88, 98.6 % with AH3 and GBM, and the highest identity 99.1 % with the other eight isolates. The comparison indicated that high conservation of 3C protein in wild-type HAV played an important role in the replication of virus. 3D among isolates of genotype I showed no significant variation. But the 3D polymerase of SLF88 showed low identity with that of DL3, with only 93.3 % identity (33 amino acid differences). GBM showed a little higher variability among the eleven reference isolates in P3 region, especially for 3A and 3D. The identities for these two proteins between DL3 and GBM were 93.2 % (5 amino acid differences) and 97.8 % (11 amino acid differences), respectively. Even for 3B, GBM had 1 amino acid difference with DL3.

3' Untranslated region
The 3’ UTR of DL3 was 63-nucleotide long, the same as those of 9 isolates, 1 nucleotide less than that of LA, and 14 nucleotides more than that of SLF88. All twelve isolates had two stop codons, separated by 6 nucleotides. The 3’ UTR
exhibited the great identity in nucleotide sequence between DL3 and isolates from Japan, GBM and HM175, with only 1 or 2 nucleotide difference. But for DL3 and the other 3 isolates, identities were lower, especially 71.4% between DL3 and SLF88.

**Subgenotype classification**
Alignment analysis of 168 nucleotides in VP1/2A (nucleotide 3 024-3 191) of DL3 with other eleven wild-type isolates revealed that DL3 had >92.5% identities with six Japanese isolates, LA and GBM (IA subgenotype), <92.5% identity with HM175 (IB subgenotype), and only 83.9% identity with SLF88 (the newly determined isolate of genotype VII). According to these comparison, DL3 was classified as subgenotype IA. Something strange was that comparison of the same region between DL3 and MBB showed 94% identity, suggesting DL3 to be a subgenotype IB. To solve this paradox, the VP1 amino terminus (nucleotide 2 208-2 468) of DL3 was compared with isolate MBB. The result showed DL3 belonged to subgenotype IA.

**Phylogenetic relationships**
Phylogenetic relations of various HAV wild-type isolates were analyzed on the genomes of twelve isolates (Figure 2). The result showed that twelve wild-type isolates formed four main subclusters, subcluster I contained DL3, AH1, FH2 and FH3, subcluster II contained AH2 and AH3, subcluster III contained LA and GBM, and subcluster IV contained MBB and HM175. Subcluster I, II, III and FH1 composed of subgenotype IA cluster. SLF88 located near this cluster. By this phylogenetic analysis DL3 isolate showed closer phylogenetic affiliation to AH1 and FH2. Besides, the comparison of full-length genome of nine isolates showed DL3 had higher identity with AH1 and FH2 than those with GBM, LA, HM175, MBB and SLF88. The analysis suggested that phylogenetic relations of various HAV wild-type isolates correlated with geographical region.

**Figure 2** Phylogenetic relations of DL3 full-length genome to those of other HAV.

**DISCUSSION**
Although hepatitis A remains a disease of major public health importance in China, the study on the genome of wild-type HAV isolate in China has not been reported. Here, we analyzed the full-length genome of a wild-type HAV isolate DL3 in China. The genome sequence of DL3 contained 7 476 nucleotides, dividing into three regions, 732-nucleotide 5′ UTR, 6681-nucleotide ORF coding for a polyprotein of 2 227 amino acid, 63-nucleotide 3′ UTR.

Sequence comparisons of genomes showed DL3 shared the highest nucleotide identity (97.5%) with AH1 and amino acid identity (99.2%) with AH2 and FH3, and lowest nucleotide identity (85.7%) and amino acid identity (96.8%) with SLF88. But the nucleotide sequence and amino acid sequence differences were not evenly distributed along the genome in all isolates. The 5′ UTR region showed high conservatism, and even SLF88 is 88.5% identical to DL3, an identity higher than that of full-length genome. This suggests the importance of this region for HAV. The 3′ UTR region showed high identities between DL3 and other isolates of genotype I. When compared with SLF88, DL3 had high divergence of 28.6% with SLF88, which lead to 18 nucleotide changes, including 14 deletions and 4 mutations. Although the exact function of 3′ UTR is unknown, several studies have revealed that sequence elements within the 3′ UTR are expected to play an important role in initiation and regulation of synthesis of HAV RNA[35, 36]. The high heterogeneity between DL3 and SLF88 in the 3′ UTR may suggest the difference of the replicating characteristics between DL3 and SLF88.

In structural region, the highest difference (13 amino acid differences, between DL3 and LA) existed in VP1. But this high divergence was due to 1 amino acid insertion into LA. For DL3 and other eleven isolates, high homology was observed, and the high variability of VP1 reported early was not observed. In nonstructural region the majority of amino acid identities between DL3 and other eleven isolates appeared in 3B and 3C proteins, implying that both proteins play important roles in the replication of virus and the processing of polyprotein. Differences between DL3 and other isolates of subgenotype IA in 2C region were a little more than those in 2B region. But differences between DL3 and other isolates of subgenotype IB or SLF88 in 2C region were much more than those in 2B region. SLF88 had 33 amino acid mutations with DL3 in 3D, suggesting the polymerase of SLF88 may have different efficiency compared with DL3.

Based on the higher nucleotide sequence identities of DL3 with AH1 and FH2 in the whole 7 476-nucleotide genome (97.5% and 97.2%, respectively), 732-nucleotide 5′ UTR region (98.0% and 98.2%, respectively), 63-nucleotide 3′ UTR region (98.4% for both), 168-nucleotide region of subgenotype classification (98.2%), 6681-nucleotide ORF (97.5% and 97.2%, respectively). DL3 is found to be in phylogenesis closest to AH1 and FH2. So we proposed that DL3, AH1 and FH2 probably represent different isolates of the same strain. Moreover, it is noteworthy that Japan where AH1 and FH2 were isolated geographically close to China, where DL3 was isolated. The phylogenetic affiliation and geographical closure suggested the epidemiological link of hepatitis A happened in Dalian and Japan.

By comparison of VP1/2A junction, genotypes were defined by at least 15%, and subgenotypes differed by at least 7.5%. According to this, we defined DL3 as subgenotype IA by comparisons with isolates HM175, AH1, AH2, AH3, FH1, FH2 and FH3. Interestingly, comparisons of this region with isolate MBB showed DL3 to be subgenotype IB. To solve this paradox, the VP1 amino terminus of DL3 was compared with isolate MBB. The comparison showed DL3 belonged to subgenotype IA. This suggests that it may not be so accurate to define a genotype only by comparison of VP1/2A junction. And the more accurate result may be achieved by comparison of more nucleic acid fragments.

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