Molecular epidemiology of hepatitis E virus infections in Shanghai, China

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Abstract

Background: Hepatitis E virus (HEV) causes acute or fulminant hepatitis in humans and is an important public health concern in many developing countries. China has a high incidence of HEV epidemics, with at least three genotypes (1, 3 and 4) and nine subtypes (1b, 1c, 3b, 4a, 4b, 4d, 4g, 4h and 4i) so far identified. Since genotype 3 and the newly identified subtype 4i have been exclusively limited geographically to Shanghai and its neighboring provinces, the epidemiology of HEV infections within the municipality, a major industrial and commercial center, deserves closer attention.

Findings: A total of 65 sequences, 60 located within the HEV SH-SW-zs1 genome [GenBank:EF570133], together with five full-length swine and human HEV genomic sequences, all emanating from Shanghai, were retrieved from GenBank. Consistent with the primary role of genotype 4 in China overall, analysis of the sequences revealed this to have been the dominant genotype (58/65) in Shanghai. Six HEV subtypes (3b, 4a, 4b, 4d, 4h and 4i) were also represented. However, although subtype 4a is the dominant subtype throughout China, subtype 4i (29/65) was the most prevalent subtype among the Shanghai sequences, followed by subtypes 4d (10/65) and 4h (9/65). Subtypes 4h, 4i and 4d were found in both swine and humans, whereas 4b was found only in swine and subtype 4a only in humans.

Conclusions: Six different swine and human HEV subtypes have so far been documented in Shanghai. More molecular epidemiological investigations of HEV in swine, and particularly among the human population, should be undertaken.

Keywords: Hepatitis E virus, Epidemiology, Shanghai municipality, Virus genotypes, Virus subtypes

Findings

Hepatitis E virus (HEV), the causative agent of acute or fulminant hepatitis in humans, is an important public health concern in many developing countries. It is estimated that about two billion people, or one-third of the world population, live in areas where HEV is endemic and are therefore at risk of infection [1]. The disease is thought to be transmitted by the fecal-oral route, usually through contaminated drinking water.

HEV is a non-enveloped, single stranded, positive-sense RNA virus belonging to the family *Hepeviridae*. At least four mammalian HEV genotypes have been recognized [2]. Genotypes 1 and 2 are primarily associated with fecal-oral transmission among humans and, in developing countries, can lead to waterborne jaundice epidemics. Genotypes 3 and 4 circulate in humans and several animal species, and are associated with sporadic infections among humans in industrialized countries [3]. In addition, two putative HEV genotypes, one from the Norway rat (*Rattus norvegicus*) [4] and the other from wild boar [5], were recently reported.

A recent study in China, where there is a high frequency of HEV epidemics, has shown that HEV seroprevalence among the general population was almost 40% and increased with age at a rate of about 1% per year [6]. Furthermore, the number of fecal samples taken from young swine that tested positive for HEV RNA ranged between 20-48% [7,8]. According to Zhu *et al.*
and the sequence percent identity was calculated using Clustal W (version 1.8), together with five full-length swine and human HEV genomic sequences, all derived from Shanghai, were retrieved from GenBank as of June, 2011. All the sequences were aligned with Clustal W (version 1.8), and the sequence percent identity was calculated using Lasergene (version 7.10; DNASTar). Phylogenetic trees were constructed by the neighbor-joining method [10], based on the partial nucleotide sequences of the ORF2 region. Bootstrap values were determined on 3,000 resamplings of the data sets [11]. The criteria used to define HEV genotypes or subtypes were adopted from Lu et al. [12]. These authors demonstrated that an assemblage of 300-450 nucleotides at the 5’ end of the HEV ORF2 region was highly conserved and that phylogenetic analysis based on this region provided accurate information about the genetic relationships between the HEV isolates and their evolutionary state. The accession numbers of HEV reference sequences are shown in Figure 1.

Phylogenetic analysis indicated that six different HEV subtypes, i.e. 3b, 4a, 4b, 4d, 4h and 4i (Figure 1) have so far been prevalent in Shanghai. Genotype 4 was the most highly represented genotype among the 65 Shanghai sequences, and subtype 4i (29/65), followed by 4d (10/65) and 4h (9/65), the most prevalent subtypes. Of the 29 sequences classified as subtype 4i, four were of human origin and 25 were from swine. HEV strain SH-SW-zs1 [GenBank: EF570133], originating from swine and regarded as the Shanghai prototype HEV strain, belonged to subtype 4i. Furthermore, all the subtype 4i HEV strains so far reported in China were isolated in the eastern part of the country, with 70% emanating from Shanghai, suggesting that subtype 4i strains are perhaps indigenous to the municipality. Also included among this subtype was the HEV strain, E067-SJ105C [GenBank: AB369690], collected from a patient suffering from acute hepatitis E in Japan who had traveled to Shanghai before the onset of the disease. Furthermore, the partial sequence [GenBank: EU034710] of another subtype 4i strain isolated from a human source in Shanghai was found to be virtually identical to a partial HEV sequence [GenBank: EU034714] obtained from a sample of swine serum in eastern China during the same period, suggesting that the isolates had a common origin [13]. To date, a total of nine subtype 4i HEV strains identified from full genomic sequences deposited in GenBank have emanated either from Japan or from Shanghai and neighboring Jiangsu Province. HEV strains of human origin emanating from Japan (JYN-Shiz08L and JKS-Shiz07L) and from Shanghai (E067-SJ105C) were closely related to HEV strains isolated from wild boar (wbJGF08-1) [14,15] and from swine (SAAS-FX17) [16], respectively. In addition, when phylogenetic clustering of subtype 4i HEV strains was examined (data not shown), sequences from Japan and Shanghai were positioned on different subtype 4i branches, suggesting that each of these two groups of strains derived from different origins. When compared with other HEV subtype 4 strains across the whole genome, only two specific amino acid substitutions were identified within the ORF1 of subtype 4i. One substitution was located in the methyltransferase motif and the other in the macro domain and might therefore influence HEV replication [17].

In this study, only subtypes 4h, 4i and 4d were found in both swine and humans, whereas subtypes 4a and 4b were confined to either human or swine populations, respectively. According to Zhu et al. [16], subtypes 4a and 4b HEV were capable of infecting both humans and swine. Subtype 4a HEV is widely distributed among both the swine and human populations in China as a whole but swine subtype 4a HEV has not been identified in Shanghai. A human subtype 4a strain isolated in Shanghai, JYI-ChiSai01C [GenBank: AB197674], showed the highest nucleotide similarity (94.0%) with a swine HEV strain Ch-S-1, from Jilin Province [GenBank: EF077630], suggesting that 4a subtypes in Shanghai were also zoonotic. Human subtype 4b HEV has a more limited distribution in China than the swine counterpart [16] and, so far, no human subtype 4b strain has been reported in Shanghai. However, not every sample from hepatitis E cases in Shanghai has been analyzed sufficiently and the possibility that human subtype 4b exists among the population of the municipality cannot be excluded.

Genotype 3 HEV strains were geographically limited to Shanghai and neighboring provinces, and seven sequences were classified into subtype 3b. No human genotype 3 HEV sequences were among those from Shanghai although one strain, EChN22 [GenBank: HM439285], was reported recently in neighboring Jiangsu Province [18]. The EChN22 isolate clustered closely with Shanghai swine isolate FJ527832 and shared 97.2% nucleotide and 99.6% amino acid homologies, which suggested that genotype 3 strains prevalent in Shanghai possibly participated in human-swine transmission.
Figure 1 Phylogenetic tree depicting genotypic/subgenotypic status of swine and human HEV strains isolated in Shanghai. A) Analyses based on 5 full-length swine and human HEV genomic sequences and 44 ORF2 sequences located between nt 6,104 and nt 6,256 within the HEV SH-SW-zs1 genome [GenBank:EF570133]. B) Analyses based on 16 sequences located between nt 6,360 and nt 6,509 within the HEV SH-SW-zs1 genome. Percent bootstrap support is indicated at each node. For each phylogeny, HEV subtypes were indicated on the outside of the square brackets that define the HEV subtypes. Each branch is labeled with the host of the HEV isolate, GenBank accession number and the year the strain was isolated. Reference sequences are labeled with the prefix RE, the subtype of the sequence, the GenBank accession number and the geographical source.
No genotype 1 HEV representatives were found in this study, possibly due to the limited number (10/65) of human HEV sequences available.

In summary, six different HEV subtypes, i.e. 3b, 4a, 4b, 4d, 4h and 4i, have so far been shown to be prevalent in Shanghai, with genotype 4 and subtype 4i the dominant forms. Only subtypes 4h, 4i and 4d were found in both swine and human hosts, whereas subtypes 4a and 4b were confined to either human or swine populations, respectively. A recent epidemiological investigation has shown that the average incidence of HEV RNA positives among the swine population of Shanghai was approximately 20.0%, and that serum IgG positives reached 72.18% [19]. Hepatitis infection rates (all types) ranged between 15.0–35.0% among the younger population and reached 47.0% in older people. Furthermore, 39.4–69.7% cases of sporadic hepatitis E were identified. Serological epidemiology investigations of HEV among swine, after hepatitis A and B [21]. Consequently, more molecular epidemiology investigations of HEV among swine, and particularly human, populations should be undertaken to assist in HEV control and prevention.

List of abbreviations
HEV: Hepatitis E virus; ORF: Open reading frame

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Authors’ contributions
YMZ, FYS, RSY, YSZ and ZL participated in the study design, YMZ, SJD and FSS performed the sequence analysis, and all authors participated in writing and revising the manuscript. All authors have read and approved the final version.

Competing interests
The authors declare that they have no competing interests.

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