

Molecular Detection of Hepatitis E Virus in Sewage Samples[∇]

Giuseppina La Rosa,* Manoochehr Pourshaban, Marcello Iaconelli,
 Valentina Spuri Vennarucci, and Michele Muscillo

Istituto Superiore di Sanità, Department of Environment and Primary Prevention, Rome, Italy

Received 8 February 2010/Accepted 24 June 2010

Human hepatitis E virus (HEV) is considered an emerging pathogen in industrialized countries. In Italy, the true burden of HEV infection is unknown. Molecular HEV screening of raw sewage samples from 11 wastewater treatment plants yielded 19 positives (16%; 18 genotype I, 1 genotype III) evenly distributed throughout Italy. Evidence that HEV could be establishing itself in our region is accumulating and may justify more active surveillance to monitor its spread.

Hepatitis E is a self-limited, enterically transmitted acute viral hepatitis that occurs most frequently in epidemic outbreaks and often spreads by way of fecally contaminated drinking water (5, 20). Hepatitis E virus (HEV) infections are caused by a positive-sense, nonenveloped RNA virus of the *Hepevirus* genus. The four major genotypes (GI to GIV), all belonging to a single serotype, are known to infect humans. While GI and GII are restricted to humans, GIII and GIV are zoonotic and may infect animals (swine, chickens, deer, mon-gooses, and rabbits), as well as humans, in both industrialized and nonindustrialized countries (18, 19). GI consists of epidemic strains circulating in Africa and Asia. GII is found in Mexico and Africa. GIII is widely distributed, mainly—but not exclusively—in the United States, Europe, and Japan. GIV is present in Asia (16). An HEV strain belonging to a fifth genotype has been identified in birds (12).

HEV is transmitted by the fecal-oral route. Large water-borne outbreaks with high attack rates among young adults have been described in regions characterized by poor sanitary conditions (22). Hepatitis E is responsible for over 50% of cases of acute viral hepatitis in countries where the disease is endemic (Central and Southeast Asia, North and West Africa, and Mexico), where seroprevalence rates range from 15% to 60% (8). North America and Europe have traditionally been considered areas where HEV is not endemic, with acute infection diagnosed rarely and largely confined to travelers returning from areas where the disease is endemic. The high rates of HEV IgG positivity reported in different studies, however, suggest that unrecognized or subclinical infection is common (8). In Europe, increasing numbers of HEV infections not associated with travel have been recently reported (15).

HEV infection may vary in severity from asymptomatic to fulminant. Case fatality rates range between 0.5% and 4% overall but may reach 25% among pregnant women (1). In industrialized countries, the case fatality rate seems to be higher than in areas where the disease is endemic, since infec-

tion occurs more frequently in elderly people with chronic liver disease, a subgroup of patients with a case fatality rate approaching 70% (26).

HEV, which is shed in the feces of infected individuals, has been detected in sewage samples, suggesting that HEV contamination of aquatic environments may also be present (2, 6, 7, 23). In Italy, the true burden of HEV infection is still unknown and there are no available studies on the presence of this virus in sewage. The prevalence of anti-HEV antibodies among healthy individuals has been found to be approximately 1% in the northern regions and up to 5% in the southern regions, including Sicily and Sardinia. Higher prevalence rates have been found among drug users (especially HIV-infected individuals), hemodialysis patients, and patients with chronic hepatitis C, suggesting that HEV may be transmitted not only by the fecal-oral route (the main mode of transmission) but also parenterally (27).

The objective of the present study was to investigate the occurrence of HEV through the molecular screening of raw sewage samples collected from urban wastewater treatment plants (WTPs) in different regions of Italy.

MATERIALS AND METHODS

Samples (118 inflow grab samples) were collected on a monthly basis from April 2008 to March 2009 at 11 WTPs located in the following regions throughout Italy: Campania, Umbria, Tuscany, Piedmont, Friuli-Venezia Giulia, Basilicata, Lombardy, Emilia Romagna, Veneto, Latium, and Sardinia (this region was enrolled later in the project) (Table 1). Due to incomplete compliance and to the fact that one of the regions, Sardinia, was enrolled only in December 2008, 118 samples were collected rather than the expected 132.

RNA was extracted from 10 ml of sewage using the NucliSens miniMAG (bioMérieux Italia S.p.A., Rome, Italy) nucleic acid isolation kit. RNAs were then eluted in 100 μ l elution buffer and stored in aliquots at -80°C until use, as previously described (13). A feline calicivirus (FCV; strain CVF9) was used as an internal control for some of the samples. A known amount of CVF9 (10⁶ 50% cell culture infective doses) was added to the samples prior to processing. Average recovery, used as a measure of extraction efficiency, was calculated as the mean ratio of genome copies (GCs) detected after and before concentration (GCs after/GCs before) by using a specific real-time PCR for CVF9 (PCR 652). Both the primers and the TaqMan probe were designed using Primer Express Software v3.0 (Applied Biosystems) (Table 2). The calibration curve used to calculate the number of FCV GCs was based on a linearized plasmid prepared from a recombinant pCR4-TOPO vector containing 630 bp of the polymerase region of strain CVF9 (PCR 650).

The above PCR, using the reagents and enzymes employed for HEV detection, was also used on HEV-negative samples in order to exclude the presence of inhibitors.

* Corresponding author. Mailing address: Istituto Superiore di Sanità, Department of Environment and Primary Prevention, Viale Regina Elena 299, 00161 Rome, Italy. Phone and fax: 39-06-49902718. E-mail: giuseppina.larosa@iss.it.

[∇] Published ahead of print on 2 July 2010.

TABLE 1. Environmental samples used in this study^a

Sample	City	WTP	Collection date
1179	Trieste	Servola	7 May 2008
1183	Bologna	Corticella	21 May 2008
1185	Venice	Fusina	21 May 2008
1191	Brescia	Verziano	10 June 2008
1249	Rome	Roma Sud	30 September 2008
1307	Venice	Fusina	15 December 2008
1309	Perugia	Pian della Genna	16 December 2008
1315	Turin	Castiglione T.se	17 December 2008
1317	Turin	Castiglione T.se	13 January 2009
1321	Brescia	Verziano	19 January 2009
1325	Bologna	Corticella	21 January 2009
1327	Venice	Fusina	19 January 2009
1329	Perugia	Pian della Genna	20 January 2009
1341	Brescia	Verziano	9 February 2009
1345	Naples	Cuma	12 February 2009
1349	Rome	Roma Sud	25 February 2009
1357	Venice	Fusina	11 March 2009
1359	Brescia	Verziano	11 March 2009
1363	Potenza	Tiera di Vaglio	11 March 2009

^a GenBank database accession no. FN796464 to FN796481.

Samples were analyzed by reverse transcriptase PCR (RT-PCR), followed by nested PCR, using both published and newly designed primers targeting portions of the RdRp gene (Table 2). Among the primers used to detect GI (RT-PCR 653, primers 1661 and 1662; nested PCR 654, primers 1663 and 1664), two (1662 and 1663) were originally described by McCaustland et al. (17). In the present study, however, several degeneracies, based on information drawn from public sequence databases, were introduced into the above primers to obtain a broader range of detection. Primers for GIII (RT-PCR 660,

primers 1669 and 1670; nested PCR 661, primers 1671 and 1672) were designed by our group to target the same fragment as that targeted by the GI assay. This was done through multiple alignment of GenBank HEV GIII sequences.

Two microliters of the extracted RNA and 22 pmol of each primer were used in a final mixture of 25 μ l using GoTaq Green Master Mix 2x, RNase inhibitor, and Moloney murine leukemia virus RT by Promega as previously described (14). GIII amplifications were carried out in a GeneAmp PCR System 9700 thermocycler (Applied Biosystems) under the following conditions: reverse transcription at 42°C for 45 min, 1 cycle of template denaturation at 94°C for 5 min, and 35 cycles at 94°C for 30 s, 50°C for 30 s, and 72°C for 30 s, followed by 1 cycle of elongation for 10 min at 72°C. Following the reaction, 1 μ l of this mixture was subjected to a second round of PCR involving 35 cycles of amplification under the same conditions. For GI, the annealing temperature was raised to 54°C. The second round is able to amplify a 175-bp fragment of HEV GI (PCR 654) and a 163-bp fragment of HEV GIII (PCR 661) (Table 2). Standard precautions were taken to prevent PCR contamination.

PCR amplicons were directly sequenced with a capillary automatic sequencer (ABI PRISM 310 Genetic Analyzer; Applied Biosystems). Some products were also cloned into the pCR4-TOPO vector (Invitrogen), following the manufacturer's protocol, and sequenced using vector-specific primers.

Bioinformatic analysis included all of the nucleotide sequences obtained here, as well as 27 prototype sequences obtained from GenBank (GI to GIV sequences and an avian HEV used as an outgroup). The analysis was performed as follows. The raw forward and reverse ABI files were aligned and assembled into a single consensus sequence using GAP4 of the Staden package. All sequences were submitted to BLAST analysis for genotyping at <http://blast.ncbi.nlm.nih.gov/Blast.cgi>. The nucleotide alignment was tested using Modeltest v3.1 (25), coupled with PAUP* v4.0 software to find the best-fit model of nucleotide substitution. The model selected (general time reversible) was used for the Bayesian analysis as implemented in the BEAST (Bayesian evolutionary analysis sampling trees) program available at <http://beast.bio.ed.ac.uk>.

Nucleotide sequence accession numbers. All of the sequences determined in this study were deposited in the GenBank database under accession no. FN796464 to FN796481.

TABLE 2. PCRs and primers used in this study

PCR no. and primer	Sequence (5' → 3')	Product length (bp)	Primer position (5' → 3') ^b	Virus
653				
1661	TTAYGGKATGCCTTTGATGACACC	302	4329–4353	HEV, GI
1662	TRATAACGGCCATRTTCCAGACAGTATTCC		4630–4601	
654				
1663	TGTTTGAGAATGACTTTTCTGAGTTTGAYT	175	4394–4423	HEV, GI
1664	TTCCAAAACCTCGCAGAGAC		4568–4548	
660				
1669	GGYGACGCYATGAGGAGT	299	4362–4380	HEV, GIII
1670	GCTATRATYGCCATRTTCCA		4660–4641	
661				
1671	TGGTRTTTGARAATGAYTT	163	4420–4438	HEV, GIII
1672	AGAGACTCCTTCGGCGCCTG		4582–4563	
650				
1654	CCCCTGGAGAAGGTTAGTGA	630	4256–4272	FCV
1655	CTTGTCACCCGAGTTGGTT		4885–4866	
651				
1656	CAAGGTATTTGCGGTCGATT	340	4453–4472	FCV
1657	ACCATCATCCCCGTAAGTCA		4792–4773	
652				
1658	GGCCTATACAGGTTGGTATCAACA	100	4380–4403	FCV
1659	TTGGAATAATCGACCGCAAAT		4479–4459	
1660 ^a	ATAGCCCCAGCGTCAAGCGC		4407–4427	

^a Primer 5' labeled with 6-carboxyfluorescein and 3' labeled with 6-carboxytetramethylrhodamine.

^b For PCRs 653 and 654, the primer positions are based on GenBank sequence accession no. M73218; for PCRs 660 and 661, the primer positions are based on GenBank sequence accession no. AB291960. For PCRs 650, 651, and 652 (FCV) the primer positions are based on GenBank sequence accession no. M86379.

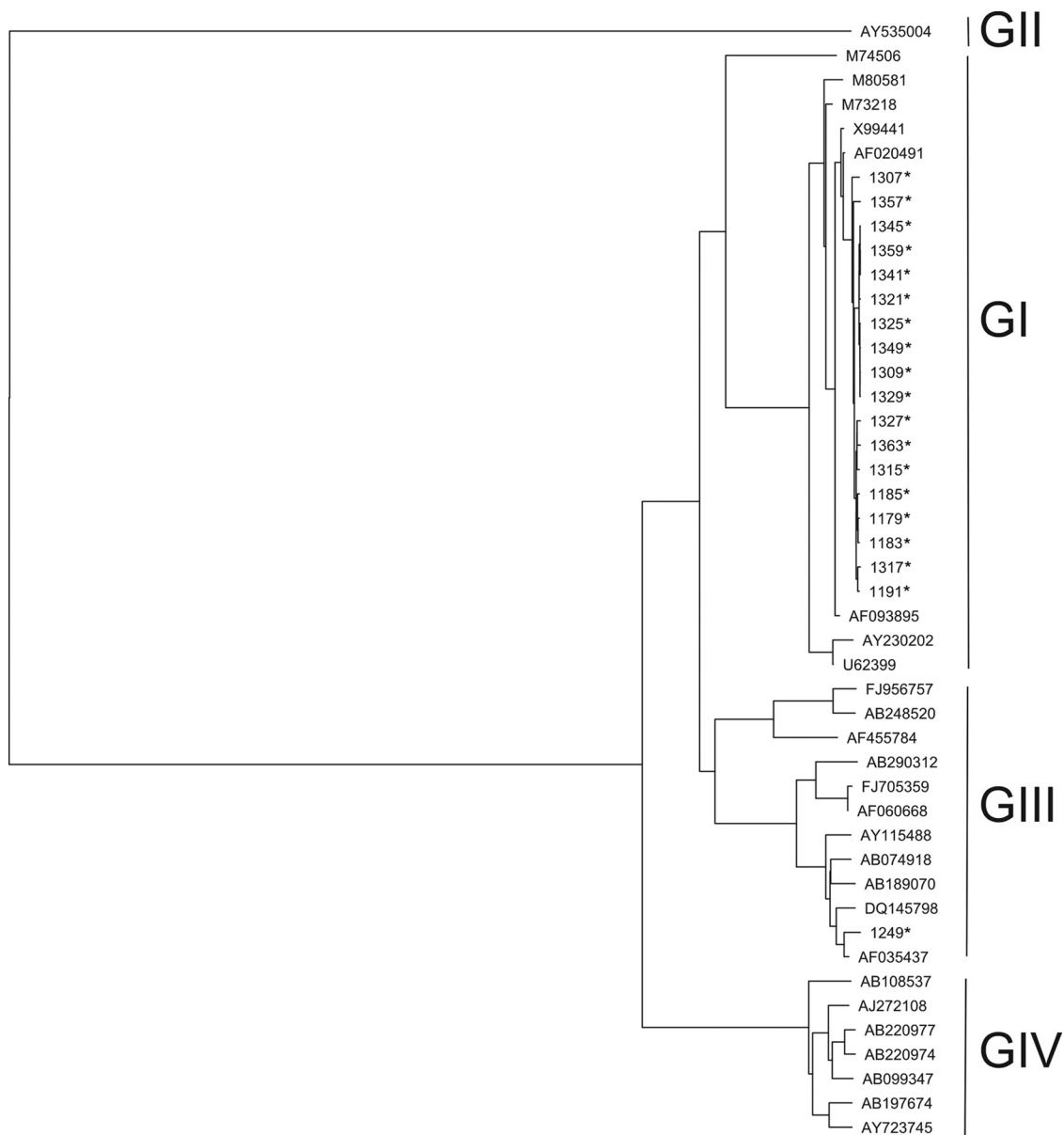


FIG. 1. Phylogenetic analysis of 19 HEV strains isolated in the present study and 27 prototype HEV strains obtained from GenBank. *, HEV-positive sewage sample.

RESULTS

Extraction efficiency was good, with an average FCV recovery exceeding 50%. Of 118 sewage samples, 19 (16%) were positive for HEV RNA. PCR inhibitors were not detected in the negative samples.

Figure 1 shows the genetic relationships between the HEV sequences obtained in this study and prototype sequences obtained from GenBank (multialignment of 127-bp fragments). The 19 HEV isolates formed two clusters, one comprising GI strains (18 isolates) and the other comprising a single GIII strain. The mean pairwise distance between the GI sequences detected in this study (18 taxa) was 0.7%. The HEV GIII strain showed sequence similarity to a strain isolated in the United States from a non-travel-related case

(AF035437) and with strain CAM-3F13 (DQ145798), detected in swine feces in Cambodia in 2006.

The vast majority of positive samples were collected in winter or spring (no positive samples were found in July, August, October, or November; only one positive in September, peak of five positives in January).

Geographically, HEV sequences were detected in 9 out of the 11 regions analyzed, distributed throughout the country.

DISCUSSION

The purpose of the present study was to provide preliminary information regarding the occurrence of HEV in the population of Italy. This was done through molecular analysis of urban sew-

age samples. Similar studies have been conducted in other European countries (Spain and France), as well as in the United States. In these studies, HEV strains belonging to GIII and also, sporadically, to GI were found in urban sewage and biosolids (2, 6, 7, 23). While the detection of virions by molecular methods does not necessarily entail the presence of infectious particles, infectious HEV particles have been reported to occur in sewage (24), indicating the existence of a possible public health risk from the contamination of surface water with HEV.

Despite the scarcity of information on the occurrence of HEV in Italy, different authors have demonstrated a sustained circulation of the virus in this country. The virus is generally diagnosed in travelers returning from areas where the disease is endemic, but non-travel-related cases have also been documented, indicating that a reservoir of HEV may exist (3, 11, 30).

Our results revealed a high proportion of HEV-positive sewage samples in Italy (16%), evenly distributed across the entire country. GI was found in all but one of our positive samples. This high proportion of GI in Italy has yet to be explained and its public health implications studied, seeing that this genotype has been associated with significant epidemics in nonindustrialized countries and has thus far been identified in Western Europe relatively rarely. Increased migration, tourism, and international trade, particularly of food products, may all contribute to the spread of HEV into new regions. GIII, found in only one sample in our study, comprises both human and animal (mainly swine and wild boar) strains and includes strains responsible for sporadic cases of hepatitis in industrialized countries. The GIII strain detected in this work (sample 1249) showed high sequence identity (95%) with a strain isolated from a pig in 2006. In Italy, HEV strains have often been detected in swine populations (4, 9, 10). These data are in line with recent reports from Spain and Netherlands (28, 29) suggesting that HEV infection is widespread in European pig farms.

Different studies have demonstrated that human and swine HEV sequences, especially when obtained in the same geographical area, are highly homologous or even identical (16, 21), suggesting not only the possibility of zoonotic transmission but also that of food-borne transmission.

Results from this study indicate that HEV strains could be more widespread than previously thought. More studies are necessary to gather information on the occurrence and diversity of the strains circulating in humans and animals in order to gain a better understanding of the epidemiology of HEV and plan adequate preventive measures.

ACKNOWLEDGMENTS

This study was funded by the project Rapid Diagnosis of Viruses in Sewage (CCM no. 4393, ISS prot. 8M09) of the Italian Center for Disease Control and Prevention (CCM—Centro Nazionale per la Prevenzione e il Controllo delle Malattie) and by the Italian Ministry of Labor, Health, and Social Policy.

We thank Emilia Luca and Marta Fratini for technical assistance and Simona Sermoneta for insightful linguistic and critical revision of the manuscript. We are also grateful to Vito Martella, University of Bari, Italy, for providing the calcivirus strain used as a control.

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