

Novel Hepatitis E Virus in Ferrets, the Netherlands

To the Editor: Hepatitis E virus (HEV), a member of the family *Hepeviridae* and the genus *Hepevirus*, is transmitted by the fecal–oral route and causes liver inflammation, which leads to mortality rates of $\approx 20\%$ in pregnant woman (1,2). Human hepatitis E is a major disease not only in developing countries but also in industrialized countries, and identification of animal strains of HEV in pigs and deer and its zoonotic potential has raised considerable public health concerns (1,3). Recent reports suggest that other animals such as rats, mongooses, chickens, rabbits, and trout also may harbor HEVs (1–5). The genomes of these viruses are ≈ 6.6 kb–7.2 kb and encode 3 open reading frames (ORFs) flanked by a capped 5' end and a poly A tail at the 3' end (1,3). We used random PCR amplification and high-throughput sequencing technology to investigate HEV sequences in ferrets (*Mustela putorius*) from the Netherlands.

In 2010, fecal samples were collected from ferrets in the Netherlands and stored at -80°C . Samples that were negative for ferret coronavirus (6) were further characterized for other pathogens. Viral RNA was isolated and viral metagenomic libraries were constructed for 454 pyrosequencing as described (7,8), and 248,840 sequence reads were generated from 7 fecal samples. Using Blastn and Blastx (www.ncbi.nlm.nih.gov/BLAST), we identified 289 sequence reads in 1 sample that were related to rat HEV and that could be assembled into 6 contigs covering $\approx 50\%$ of the ferret HEV (FRHEV) genome.

We then developed a set of nested PCR primers on the basis of obtained sequences to detect viral RNA (online Technical Appendix Table 1,

www.cdc.gov/EID/pdfs/11-1659-Techapp.pdf). Total RNA extracted from 43 ferret fecal samples collected from 19 locations in the Netherlands was used to perform reverse transcription PCR amplification. Using this PCR, we detected viral RNA in 4 (9.3%) fecal samples tested from 4 locations (distance between each sampling location ranged from 25 km to 127 km). All amplicons were confirmed by nucleotide sequencing. We have limited information regarding the clinical disease this virus may cause because these samples were obtained from household pet ferrets that did not show overt clinical signs. In addition, 4/16 animals from a single farm were IgG positive when tested for IgG against HEV by using recombinant human HEV protein (Wantai, Beijing, China).

To characterize the complete genome of this virus, we selected 2 PCR-positive samples (FRHEV4 and FRHEV20), developed different sets of specific primers on the basis of sequence fragments obtained by 454 pyrosequencing, directly sequenced

amplicons by Sanger sequencing, and used a rapid amplification of cDNA ends PCR to obtain 5' and 3' frame end sequences. Using overlapping fragments we assembled 2 complete FRHEV genome sequences that contained 6,854 nt, including a 13-nt 3' poly A tail and a 12-nt 5' end. FRHEV full-genome sequences FRHEV4 and FRHEV20 showed 98.6% sequence identity and were deposited into GenBank under accession nos. JN998606 and JN998607, respectively.

The FRHEV genome contains a complete ORF1 gene that encodes a nonstructural protein of 1,596 aa, an ORF2 gene that encodes a capsid protein of 654 aa, an ORF3 gene that encodes a phosphoprotein of 108 aa, and a 3' noncoding region of 78 nt. Sequence analyses indicated that the FRHEV genome shared the highest identity (72.3%) with rat HEV. Sequence identity with HEV genotypes 1–4 and rabbit and avian HEVs ranged from 54.5% to 60.5% (online Technical Appendix Table 2). The FRHEV genome organization

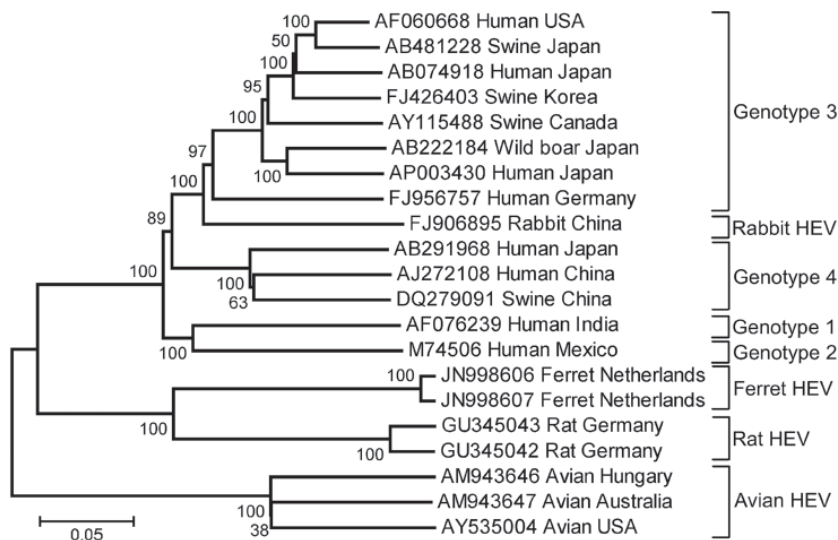


Figure. Phylogenetic tree based on the complete genomic sequences of ferret hepatitis E viruses (HEVs) and human, rabbit, swine, avian, and rat HEV strains. Names of HEV strains follow GenBank accession numbers. Sequence alignment was performed by using ClustalW in the MEGA5.0 software package (www.megasoftware.net), and the trees were constructed by using the neighbor-joining method with p-distance (gap/missing data treatment; complete deletion) and 1,000 bootstrap replicates as in MEGA version 5.0. Scale bar indicates nucleotide substitutions per site.

was found to be slightly different from other HEVs and included a putative ORF (ORF4) of 552 nt that overlapped with ORF1 (online Technical Appendix Figure). A similar pattern of genome organization was observed for both FRHEVs.

Phylogenetic analysis of the complete genomes clearly showed that FRHEV was separated from genotype 1–4 HEVs and clustered with rat HEV (Figure). Similar phylogenetic clustering was observed when nucleotide and deduced amino acid sequences of ORF1, ORF2, and ORF3 were analyzed separately. The phylogenetic distance between rat HEV and FRHEV is larger than the distance between genotype 1 and genotype 2 HEV.

In recent years, an increasing number of sporadic cases of hepatitis E have been reported (1,9). Several observations suggest that autochthonous cases are caused by zoonotic spread of infection from wild or domestic animals (1,3,9). In addition, IgG anti-HEV seropositivity in the United States has been associated with several factors, including having a pet at home (10). Further studies are needed to identify the zoonotic potential of FRHEV.

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Epidemic *Clostridium* *difficile* Ribotype 027 in Chile

To the Editor: The increased severity of *Clostridium difficile* infection is primarily attributed to the appearance of an epidemic strain characterized as PCR ribotype 027 (1). The only report that identified epidemic *C. difficile* ribotype 027 in an American country outside of North America comes from Costa Rica, raising the possibility that strains 027 might also be present in other countries of Latin America (2). Several studies between 2001 and 2009 have been conducted in South American countries to detect the incidence of *C. difficile* infection in hospitalized patients, but they did not identify which *C. difficile* strains were causing these infections (3).

During an epidemiologic screening of patients with *C. difficile* infection in a university hospital in Chile, we analyzed all stool samples of patients with suspected *C. difficile* infection during a 5-month period (June–November 2011). Two cases of *C. difficile* infection were associated with ribotype 027.

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Technical Appendix

Table 1. Primers used for amplification and sequencing of the ferret HEV genome*

Primer	Sequence, 5' → 3'	Nucleotide position†
FRHEV-F1	GGCTGGCGTTTGCTTGAGG	256–275
FRHEV-R1	TTCGAATCCAACGCTGGTGAC	1158–1178
FRHEV-F2	GTAATATCACGGCCAATGAG	1077–1096
FRHEV-R2	CAGCCTATAGGGCATAGTAAG	1681–1701
FRHEV-F3	GCCCTGACCTTGGAGCTGAC	1592–1611
FRHEV-R3	CTATTGGCGGCGTTAACTAG	2078–2097
FRHEV-F4	GAGCTTTTGCCGGATGGGTC	2015–2034
FRHEV-R4	CACTAGCCCCTTATGGTCGA	2922–2941
FRHEV-F5	CTGTTGCCAAATCGGTCGTA	2865–2884
FRHEV-R5	GCATCAGACACGCCAGCTC	3308–3327
FRHEV-F6	TGCCCTTGACGCGCCCACTG	3250–369
FRHEV-PCR-InR	CTACCKGAATGCTTCTTCC	4272–4291
FRHEV-PCR-ExF	TCCAGAAGGACTGCAACAAG	3876–3895
FRHEV-R7	CTCACCACGTCAGGAACAACC	4525–4545
FRHEV-F8	AGTTGAAGGTGGCGTTCCAC	4461–4480
FRHEV-R8	GCGCCTCGAGAATCGTAATC	5183–5202
FRHEV-F9	CGTGTTCCCGATGCCGGGCA	5114–5133
FRHEV-R9	CAGGCGAAGGCGTCGGCC	6164–6181
FRHEV3'RACE-ExF	TATACCTCTGTTGAGGCTGC	6049–6068
FRHEV3'RACE-InF	CCTTGGACTTTTCTGTTAC	6108–6128
FRHEV-5' end-F‡	GACCATCTAGCGACCTCCACTTTTTTTTTTTTTTTTTTTTTT	NA
FRHEV-5' end-R	TGCATATCTCGACCGTCAGG	338–357
FRHEV-R10	TATTTGCTGGACCCCTTC	380–397
FRHEV-PCR-ExR	CAAGATCACCGCCATGTTCCA	4319–4339
FRHEV-PCR-InF	GTCAGGGCATWTCRCSTGG	3933–3952

*HEV, hepatitis E virus; FRHEV, ferret HEV; F, forward; R, reverse; RACE, rapid amplification of cDNA ends; NA, not applicable.

†Positions of primers located in the complete genome are shown according to ferret HEV strains.

‡Nonviral primer.

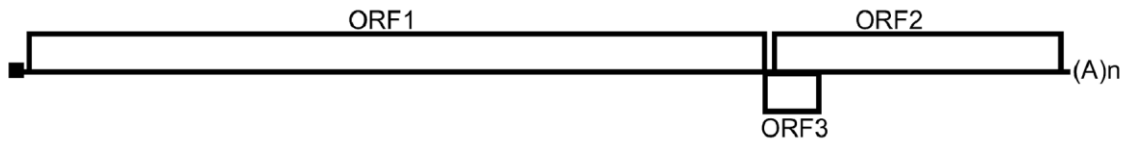
Table 2. Nucleotide and amino acid sequence identities between rat, human, rabbit, and avian HEV strains compared with ferret HEV*

Strain, GenBank accession no.	% Identity with ferret HEV							
	FRHEV 4, NJ998606				FRHEV 20, NJ998607			
	Genome, nt	ORF, aa	ORF2, aa	ORF3, aa	Genome, nt	ORF1, aa	ORF 2, aa	ORF 3, aa
RatHEV, GU345042	72.2	77.1	81.4	44.4	72.3	77.2	82.3	45.7
Genotype 1, † AF076239	59.2	53.9	60.1	19.8	59.4	54.0	60.6	21.0
Genotype 2, † M745056	58.7	54.0	60.6	19.8	58.8	61.7	61.0	21.0
Genotype 3, † AF060668	60.6	55.7	61.0	18.5	60.5	55.8	61.7	19.8
Genotype 4, † AJ272108	58.8	54.7	60.5	17.3	58.9	54.9	61.2	17.3
Rabbit HEV, FJ906895	59.8	54.9	60.1	17.3	59.8	54.9	60.8	18.5
Avian HEV, AM943646	54.5	46.9	46.0	21.0	54.5	46.9	46.5	19.8

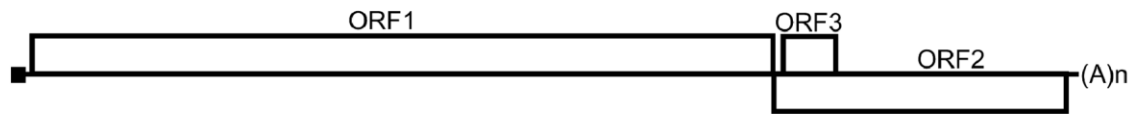
*HEV, hepatitis E virus; FRHEV, ferret HEV; ORF, open reading frame.

†Human strains.

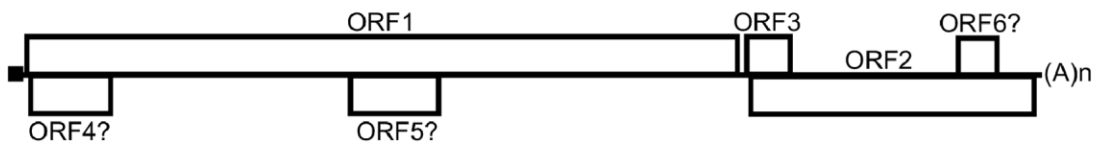
HEV genotypes 1–3



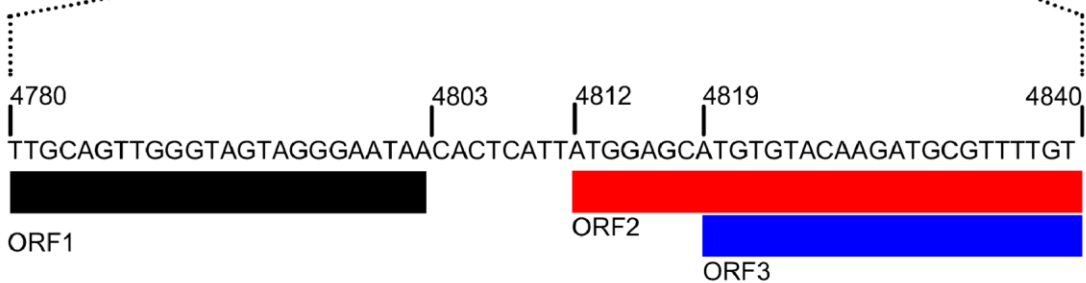
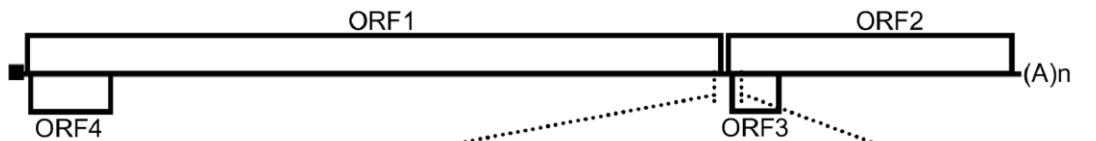
HEV genotype 4



Rat HEV



Ferret HEV



Technical Appendix Figure. Genome organization of hepatitis E viruses (HEVs) and initiation of translation of open reading frame 1 (ORF1) (black bar), ORF2 (red bar), and ORF3 (blue bar) of ferret HEV.