Human parechoviruses (HPeVs) often cause severe illness among young children. National surveillance with routine testing of all cerebrospinal fluid, fecal, and tissue samples was conducted during January 2009–December 2012 in all counties in Denmark (6,817 samples from 4,804 children were screened for HPeV). We detected HPeV RNA in 202 (3.0%) specimens from 149 persons. Young infants were at highest risk for HPeV, and 9 (6%) of the HPeV-infected children died, probably of their HPeV illness. HPeV3 was the most common genotype identified, and 5 closely related clades of HPeV3 circulated in Denmark throughout the study period. Our study adds perspective on the prevalence and clinical and molecular virologic characteristics of HPeV infection.

Human parechoviruses (HPeVs) have recently been recognized to cause a variety of symptoms ranging from mild diarrhea to sepsis and meningitis, particularly among young children. HPeVs belong to a large family of nonenveloped, positive-sense, single-stranded RNA viruses, the Picornaviridae, which comprises 12 genera (and 5 proposed genera). Six genera are associated with human infections: cardiovirus (saffold virus), cosavirus, enterovirus (EV), hepatovirus (hepatitis A), kobuvirus (Aichi virus) and HPeV. HPeV1 and HPeV2, originally known as echovirus 22 and 23 of the EV genus, respectively, were reclassified in 1999 as a separate genus (Parechovirus) on the basis of genetic and biologic differences (1). Since the reclassification, the number of known HPeVs has increased and now totals 16 genotypes (www.picornaviridae.com/parechovirus/hpev/hpev.htm).

HPeV1 is known to be associated with asymptomatic infection. HPeV3 seems to be more or less well established (2–4). The remaining known HPeVs are clinically unexplored.

Whereas HPeV3 has been reported to be associated with sepsis-like syndrome, meningitis, encephalitis, and hepatitis in neonates and young infants (2), most HPeV infections are asymptomatic or associated with mild respiratory and/or gastrointestinal symptoms (4). HPeV incidence has been reported to show a seasonal pattern in temperate climates, with different types cocirculating simultaneously (5). Although HPeV infections are relatively common in most settings, experience from long-term population surveillance is somewhat sparse. We used 4 years of national laboratory surveillance to describe the molecular epidemiology of HPeV, including phylogenetic characteristics of the emerging HPeV epidemic, in Denmark.

Materials and Methods

Study Design

During January 2009–December 2012, a total of 6,817 specimens were collected from 4,808 children from all regions of Denmark: 2,006 cerebrospinal fluid (CSF) samples from 1,952 children; 1,963 fecal samples from 1,608 children; 1,057 blood samples from 1,025 children; 682 respiratory samples from 610 children; 571 autopsy samples from 228 children; 302 swab specimens from 291 children; and 236 other samples. In general, samples were collected from patients with symptoms compatible with EV infection, particularly meningitis, sepsis-like illness, respiratory symptoms, and/or diarrhea. The CSF samples were collected from patients who typically had fever and/or clinical signs compatible with meningitis or encephalitis. The fecal samples were collected from a mixture of patients with community-acquired infection and hospitalized patients who typically had fever and/or diarrhea, and/or meningitis/sepsis-like illness. Autopsy material was analyzed for EV and HPeV as part of standard procedure irrespective of symptoms. Samples were sent by mail to the Department of Microbiological Diagnostics and Virology at the Statens Serum Institut (Copenhagen, Denmark) to undergo viral diagnostic and isolation and real-time PCR for EVs and HPeVs. HPeV infections detected in the same person within a 3-week period were considered to be the same infection. During the study period, the Department of Microbiological Diagnostics and Virology was the only laboratory conducting HPeV diagnostics in Denmark; therefore, the data reflect all HPeV infections detected in Denmark.
Laboratory Analyses

Sample Preparation

Only fecal samples and biopsy samples needed special preparation before use of the general nucleic acid extraction protocol. Fecal samples were prepared as a 10% wt/vol suspension in minimal essential medium, followed by centrifugation at 3,500 × g for 30 min to remove inhibitors. Biopsy samples were suspended in Lysis/Binding Buffer from the MagNa Pure LC Total Nucleic Acid Isolation Kit (Roche Diagnostics, Mannheim, Germany), followed by homogenization.

Nucleic Acid Isolation

Nucleic acids were extracted from 200 mL sample material. All sample types, except CSF, were processed by using the MagNa Pure 96 DNA and Viral NA Small Volume Kit on the MagNa Pure 96 instrument (Roche Diagnostics) according to the manufacturer’s specifications. Nucleic acids from CSF were isolated by using the QIAamp DNA Blood Mini Kit on the QIAcube instrument (QIAGEN, Hilden, Germany) following the manufacturer’s specifications.

Amplification and Detection

For amplification, 5 mL of extracted nucleic acids were used per reverse transcription PCR (RT-PCR) reaction (total volume 25 mL) by using the OneStep RT-PCR Kit (QIAGEN). The reaction mixtures contained 1 µmol/L of each primer and 0.2 µmol/L probe. The primers and probe used have been published (6), and the reaction mixture also contained an assay for EV. The HPeV-specific probe was labeled with a Hex dye. The Mx3005P real-time thermocycler (Agilent Technologies A/S, Hoersholm, Denmark) was used for amplification and detection with the following settings: 50°C for 20 min, 95°C for 15 min, followed by 45 cycles of 95°C for 15 s and 55°C for 1 min.

Genotyping

We amplified 256–259 bp of the viral protein (VP) 3/VP1 region (from position 2159–2458 in relation to L02971) in a nested PCR. The first-round RT-PCR was conducted by using primers Harv1-F and Harv1-R (7) with a OneStep RT-PCR kit (QIAGEN) and the following thermocycler conditions: 50°C for 30 min, 95°C for 15 min, followed by 40 cycles of 95°C for 30 s, 42°C for 30 s, and 60°C for 45 s. The second-round PCR was conducted by using primers Harv2-F and Harv2-R (8) and the following thermocycler conditions: 95°C for 6 min, followed by 40 cycles of 95°C for 30 s, 60°C for 30 s, and 72°C for 45 s. This protocol was followed by a final extension at 72°C for 10 min. PCR amplification was followed by sequencing.

Before sequencing, PCR products were treated with exo-SAP IT (GE Healthcare, Buckinghamshire, UK). PCR products were sequenced by using the dideoxynucleotide chain termination method with the ABI Prism BigDye Terminator Cycle Sequencing Reaction kit on an ABI Prism 3100 automated sequencer (Applied Biosystems, Naerum, Denmark). Sequencing was performed with the forward and reverse primers from the second-round PCR. Sequences were assembled in BioNumerics 6.5 (Applied Maths, Kortrijk, Belgium). The sequences have been submitted to GenBank under accession nos. KF300772–KF300885.

Phylogenetic Analysis

Assembled sequences were aligned with reference sequences by using the Simmonic sequence editor (www.virus-evolution.org). Phylogenetic and molecular evolutionary analyses were conducted by using MEGA 5.0 software (www.megasoftware.net). Genetic distances were calculated by using the Jukes-Cantor parameter at the nucleotide level, and the phylogenetic trees were constructed by using the maximum-likelihood method with 500 bootstrap replications (8).

Results

Sample Material

Study Population

During January 2009–December 2012, from 6,817 samples from 4,804 children, we detected HPeV RNA in 202 (3%) specimens from 149 (3%) children from all counties of Denmark. Of the 149 individual HPeV cases, 25 (17%) were detected in CSF samples, 105 (70%) in fecal samples, and 19 (13%) in a variety of clinical specimens (8 pharynx/tonsil swabs, 2 bronchio-lavage fluids, and 9 biopsy specimens from children who died unexpectedly [5 from abdominal lymph nodes, 1 from the small intestine, 2 from the large intestine, and 1 from pulmonary tissue]). Of the 149 individual HPeV cases, 125 had sufficient material or RNA for further VP3/VP1 regional subtyping.

Frequency of HPeV-infected Patients

HPeV was detected in 52 (3%) of 1,744 patients in 2009, in 31 (2%) of 1,729 patients in 2010, in 33 (2%) of 1,939 patients in 2011, and in 33 (2%) of 1,405 patients in 2012. Of the 149 HPeV-positive patients, 52 (35%) tested positive in 2009, 31 (21%) in 2010, 33 (22%) in 2011, and 33 (22%) in 2012.

HPeV3 infections occurred in all months of the year, albeit with a clear seasonal pattern; only few cases occurred during the winter and spring months of November–May, with a marked increase during June and July, peaking during the fall months of September and October (Figure 1).
HPeV1 infections show a seasonality pattern very similar to that of the HPeV3 infections, although no cases were observed during March–June.

**Epidemiology of HPeV-infected Persons**

Age was inversely associated with risk for HPeV infection among children <5 years of age: the overall median age of HPeV-infected children was 39 days (interquartile range [IQR] 22–71 days), and the median age of HPeV-negative children was 16.3 years (IQR 318 days–41.8 years). Children with HPeV3 infection (median age 37 days [IQR 19–59 days]) were significantly younger than children with HPeV1 (median age 199 days [IQR 84–303 days]).

**Clinical Features of HPeV Infection**

Information about symptoms and the source of the sample was available for 89 (60%) of children with HPeV. A wide range of symptoms were reported: slightly more cases were associated with meningitis (36 cases) than diarrhea (29) and fever (19), but sepsis-like symptoms, convulsions, apathy, and general discomfort also were reported (Table).

**Virologic and Molecular Findings**

Our sequencing of 1 of the least conservative regions of the capsid (VP3/VP1) resulted in the following: 90 HPeV3, 21 HPeV1, 8 HPeV6, 4 HPeV5, and 2 HPeV4. The genotype was assigned by BLAST analysis of the sequence against all published sequences in GenBank (http://blast.ncbi.nlm.nih.gov/).

HPeV3 was more frequently associated with disease in neonates than was any other HPeV genotype. The mean age of HPeV3-infected children was 1.6 months, compared with 8.9 months for HPeV1-infected children (n = 20), and 8.1 months for HPeV6-infected children (n = 6) (t test for mean age difference, p = 0.01). Sequences were available from 8 of the 9 children who died unexpectedly; of these, 4 sequences were HPeV1, 2 were HPeV3, 1 was HPeV5, and 1 was HPeV6. Of the 36 patients with reported meningitis, HPeV types were available for 32; of these, 30 (94%) types were HPeV3, and 2 (6%) were HPeV1. Of the 4 patients with sepsis-like syndrome, sample material for typing was available for 3; all 3 were identified as HPeV3. HPeV types were available for 25 of the 29 patients with reported diarrhea; of these, 19 (76%) types were HPeV3, 4 (16%) were HPeV1, and 2 (8%) were HPeV6. All 25 CSF samples were HPeV3. HPeV1 and HPeV3 were detected throughout the study period; HPeV6 was detected in all years except 2010; and HPeV5 emerged only in 2012.

Sequence data from 124 of these 125 samples were of sufficient quality and were used to create phylogenetic trees (Figure 2, Appendix, wwwnc.cdc.gov/EID/article/19/12/13-0569-F2.htm), which also included reference sequences obtained from GenBank for each genotype identified in this study. The phylogenetic analyses revealed the existence of 5 closely related clades of HPeV3 circulating in Denmark throughout the study period, with most clades found all years of the study, implying cocirculation of these clades without genetic selection of either clade in the sequenced area. In addition, there was no particular geographic distribution of the individual clades. The other genotypes appear to follow the same pattern of circulation and of little evolutionary change within each genotype over time. However, data for the genotypes other than HPeV3 were insufficient to substantiate a division into individual clades.

We created a phylogenetic tree (Figure 3) including a representative strain from each of the HPeV3 clades detected in our study in Denmark, combined with matching HPeV sequence from Europe available in GenBank. This tree showed that clades 1–4 were most closely related to strains from Spain (9) and Italy (10) during 2006–2009, whereas clade 5 was identical (in the sequenced area) to a simultaneous strain from Germany (11). We found an intragenotype variation of 3.9%, within HPeV1 of 5.6%, within HPeV4 of 11.6%, within HPeV5 of 1.6%, and within HPeV6 of 3.5%.

**Discussion**

Our study establishes HPeV as important differential diagnosis in young and often severely ill children in
an endemic pattern of HPeV infection in Denmark. This
seasonality (i.e., occurrence of HPeV3 every year) indicates
12). The observed sea-
characteristics of HPeV and EV (12). In our selected material (selected
-ber, coinciding with the EV season. These findings under-
marked seasonality with peaks during July and Septem-
denmark with fever and/or central nervous system symp-
toms. The sample material received for diagnostic testing,
where a significant portion of the HPeV-positive samples
are CSF, illustrates the severity of these infections. With-
out routine testing for HPeV of infants suspected to have
EV infection (EV and HPeV3 infections are indistinguish-
able in infants [12]), these children’s illnesses would not
have been diagnosed, which would have resulted in unfo-
cused treatment, such as unnecessary antimicrobial drug
therapy and unnecessary long hospital stays, as suggested
by Wolthers et al. (13). In our selected material (selected
for suspected enteroviral disease), HPeV3 was by far the
most prevalent HPeV type and was the only genotype de-
tected in CSF. The high prevalence of HPeV3 contrasts
with findings from other studies (5,14), which have es-
hanced HPeV1 as the most common virus circulating
in the population. However, these studies did not select
the patients on the basis of symptoms because the chil-
dren were part of healthy birth cohorts from prospective
diabetes studies. In Denmark, although HPeV infections
were detected throughout the year, the illness showed a
marked seasonality with peaks during July and Septem-
ber, coinciding with the EV season. These findings under-
score the need for HPeV and EV differential diagnostics
in young children who have nonbacterial meningitis and
severe disease in general caused by the similar clinical
characteristics of HPeV and EV (12). The observed sea-
sonality (i.e., occurrence of HPeV3 every year) indicates
an endemic pattern of HPeV infection in Denmark. This
finding contrasts with other HPeV studies from the United
Kingdom and the Netherlands, where a biannual cycle of
HPeV3 infections has been observed; the reason for this
difference is unknown (15,16).
In our study, the children infected with HPeV (and in par-
ticular HPeV3) were very young (median age 39 days), which
is consistent with prior studies (17,18). However, a recent
study from Japan (5) has established that adults also can be
infected with HPeV3, possibly resulting in epidemic myalgia.
We also showed that HPeV3 in Denmark did not un-
dergo significant genetic diversification during 2009–2012,
at least not in the sequenced part of the capsid (VP3/VP1)
(Figure 2, Appendix). This observation was in line with
comparisons of the strains from Denmark with strains
from Europe in which only limited variation periodically
(1994–2012) and geographically was observed (Figure 3). The genetic distances in the trees are small (which further
substantiates the results of limited evolutionary change).
These indications of limited genetic evolution correspond
well with results from a study in the Netherlands that de-
scribed HPeV during 2000–2007 (19). The reason for the
lack of intratypic diversification of the otherwise intertypi-
cally heterogenic capsid area is unknown. Perhaps most
infections are subclinical and do not elicit an immunologic
response, which might otherwise cause the virus to evolve
to escape the immune response. Structural constraints
might also exist that limit the genetic variation to ensure
proper capsid formation.
Our data supported previous findings that HPeV3 is
commonly associated with meningitis/sepsis symptoms
among young infants, and we found that convulsions and
apathy were reported among children with such severe cas-
es. Because HPeV3 has been reported to cause white matter
injury in 9 of 10 infected neonates, resulting in a range of
complications from cerebral palsy in 1 child to epilepsy in
another child to learning disabilities in a third child (20),
focused longitudinal follow-up and cognitive evaluation of
these children during childhood might be indicated to bet-
ter understand the long-term consequences of this emerg-
ing viral disease.
In conclusion, HPeV was a clinically significant virus
rivaling EVs in young children in Denmark. In particular,
HPeV3 was circulating in our clinically selected material,
with 5 different clades cocirculating over most of the years
of the study. The patients were from all around Denmark
demonstrating the need for central or disseminated HPeV di-
agnostics in Denmark so that HPeV is tested for in all young
children suspected of infection with HPeV or EV. Testing
might possibly reduce the length of hospitalization and limit
the inappropriate use of antimicrobial drugs. Furthermore,
doctors and especially pediatricians should be aware of the
symptoms of HPeV-associated diseases and should be en-
couraged to collect a sample (primarily CSF and preferably

Figure 3. Phylogenetic analysis of strains of human parechovirus
type 3 from Europe compared with a representative strain of each
of the 5 clades found in Denmark (squares). Maximum-likelihood
analysis with bootstrap values >70% are shown. Scale bar indicates
nucleotide substitutions per site.
combined with a fecal sample) for diagnostics. Centralized surveillance of this virus could provide deeper insight into the behavior of HPeV and might shed light on the clinical significance of the HPeVs other than type 3.

Acknowledgments

We are grateful for the review expertise of Marian Jørgens and her assistance in preparing the manuscript and for the highly qualified assistance with laboratory analyses by Ann Berith Petersen, Shila Mortensen, and Jesper Roenn. We thank all the clinical internal medicine, pediatric, and microbiological departments of Denmark for submission of samples for analyses at the Department of Microbiological Diagnostics and Virology.

No external financial support was used in conducting this study.

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