Comparison of epidemiology and clinical characteristics of infections by human parechovirus vs. those by enterovirus during the first month of life

María Cabrerizo¹ · Gloria Trallero¹ · María José Pena² · Amaia Cilla⁴ · Gregoria Megias⁴ · Carmen Muñoz-Almagro⁵ · Eva Del Amo⁵ · Diana Roda⁵ · Ana Isabel Mensalvas⁶ · Antonio Moreno-Docón⁶ · Juan García-Costa⁷ · Nuria Rabella⁸ · Manuel Omeñaca⁹ · María Pilar Romero¹⁰ · Sara Sanbonmatsu-Gámez¹¹ · Mercedes Pérez-Ruiz¹¹ · María José Santos-Muñoz³ · Cristina Calvo³ · And the study group of “Enterovirus and parechovirus infections in children under 3 years-old, Spain” PI12-00904

Received: 5 March 2015 / Revised: 6 May 2015 / Accepted: 8 May 2015 / Published online: 16 May 2015

Abstract Human parechoviruses (HPeV) have been recently recognized as important viral agents in paediatric infections. The aims of this study were to investigate the HPeV infection prevalence in infants <1 month in Spain and, secondly, to analyse the clinical and epidemiological characteristics of the infected patients compared with those infected by enterovirus (EV). Infants ≤1 month with neurological or systemic symptoms were included in a multicentre prospective study. EV and HPeV detection by RT-PCR and genotyping were performed in cerebrospinal fluids (CSF), sera or throat swabs. Out of the total of 84 infants studied during 2013, 32 were EV positive (38 %) and 9 HPeV positive (11 %). HPeV-3

Communicated by David Nadal

Cristina Calvo
calvorey@ono.com
María Cabrerizo
cabrerizo@isciii.es
Gloria Trallero
trallero@isciii.es
María José Pena
mpenalopd@gobiernodecanarias.org
Amaia Cilla
cillaamaia@gmail.com
Gregoria Megias
megias@saludcastillayleon.es
Carmen Muñoz-Almagro
mu@hsjdbcn.org
Eva Del Amo
delamo@fsjd.org
Diana Roda
roda@hsjdbcn.org
Ana Isabel Mensalvas
amenasalvas@hotmail.com
Antonio Moreno-Docón
antonio.moreno13@carm.es
Juan García-Costa
garcia@sergas.es
Nuria Rabella
rabella@sanpau.cat
Manuel Omeñaca
omeñaca@aragon.es
María Pilar Romero
promero.hulp@salud.madrid.org
Sara Sanbonmatsu-Gámez
saral.sanbonmatsu.sspa@juntadeandalucia.es
Mercedes Pérez-Ruiz
mercedes.perez.ruiz.sspa@juntadeandalucia.es
María José Santos-Muñoz
mjsanmu@hotmail.com

¹ Enterovirus Unit, National Centre for Microbiology, Health Institute “Carlos III”, Madrid, Spain
² Hospital Negrín de Gran Canaria, Las Palmas, Spain
³ Pediatrics Department, Hospital Severo Ochoa, Avda. Orellana, s/n., Leganés 28911, Madrid, Spain
⁴ Hospital de Burgos, Burgos, Spain
was identified in eight cases and HPeV-5 in one. Mean age of HPeV-positive patients was 18 days. Diagnoses were fever without source (FWS) (67%), clinical sepsis (22%) and encephalitis (11%). Leukocytes in blood and CSF were normal. Pleocytosis (p = 0.03) and meningitis (p = 0.001) were significantly more frequent in patients with EV infections than with HPeV.

Conclusions: Although HPeV-3 infections were detected less frequently than EV, they still account for approximately 10% of the cases analysed in infants younger than 1 month. HPeV-3 was mainly associated with FWS and without leukocytosis and pleocytosis in CSF. In these cases, HPeV screening is desirable to identify the aetiologic agent and prevent unnecessary treatment and prolonged hospitalization.

What is Known:
• Human parechovirus may be a cause of fever and clinical sepsis in the neonatal period.
• HPeV-3 might be one of the main agents causing severe neonatal neurological infections.

What is New:
• This is the first multicenter prospective study focused on newborns and contributes to a better knowledge of these viral infections. Clinical characteristics of enterovirus and parechovirus infections are compared specifically in the neonatal period.
• Knowledge of HPeV infections by paediatricians and neonatologists can guide the diagnosis of these patients and avoid unnecessary treatment and prolonged hospitalization.

Keywords Enterovirus · Human parechovirus · Sepsis · Fever · Neonate

Abbreviations
CSF Cerbroespinal fluid
EV Enterovirus
HPeV Human parechovirus
FWS Fever without source
NICU Neonatal intensive care unit

Introduction

Human parechoviruses (HPeV) are RNA viruses belonging to the family of Picornaviridae. Formerly described as echovirus

22 and 23 in the Enterovirus genus, HPeV were reclassified into their own genus, Parechovirus, in the 1990s based on genetic differences and biological properties. Then, they were renamed as HPeV-1 and HPeV-2, respectively [13, 25]. Recently, additional types of HPeV have been reported, and a total of 16 different types have been recognized to date (HPeV-1 to 16). The most common genotype detected worldwide is HPeV-1 followed by HPeV-3. Other types such as HPeV-2 and HPeV-4 are less common.

Infections with HPeV are prevalent in young children and have been associated with mild diseases of the respiratory and gastrointestinal tract but also with serious diseases such as meningitis, encephalitis and sepsis in young infants [3, 12, 13]. According to recent studies, HPeV-3 might be one of the main agents causing severe neonatal neurological infections in Europe [5, 14, 17], although its real incidence is unknown since HPeV detection is not routinely performed in the majority of clinical microbiology laboratories. Clinical data specific for neonates are also scarce.

This study describes the results obtained about the prevalence of HPeV infections in children up to 1 month of age in Spain and the clinical and epidemiological characteristics of the infected patients over a 1-year period. Furthermore, HPeV and enterovirus (EV) infections were compared.

Methods

Patients and samples In 2013, a prospective study was conducted in collaboration with 10 Spanish hospitals. The project was supported with a grant from the Health System (AES; PI12-00904). The personal data of the patients were protected, and the study was approved by the Ethics Committee of the Health Institute Carlos III. Criteria for inclusion were infants ≤1 month of age with fever without source (FWS), clinical sepsis disease or meningitis/encephalitis admitted to the participant centres, whose parents agreed to sign the informed consent. During the hospital stay, a physician filled out a study questionnaire with the clinical characteristics of the patients. FWS was defined as axillary temperature greater than 37.9 °C, which after an initial examination and laboratory evaluation has no apparent cause. Aseptic meningitis was defined by a neonate with fever, irritability, poor feeding, vomiting, or bulging fontanelle with pleocytosis >30 cells/mm³ in cerebrospinal fluid (CSF), with negative CSF culture for bacteria. Encephalitis was defined as a clinical diagnosis by the attending neurologist. It may be accompanied or not by CSF pleocytosis. Electroencephalogram and MRI compatible support the diagnosis but were not considered essential to it, and a clinical criterion of the attending neonatologist was considered enough for inclusion. Meningoencephalitis shares data from the two latter ones. Clinical sepsis is defined as an infant with an unwell appearance, with alteration of the paediatric
assessment triangle (appearance, respiratory and circulatory items). Exclusion criteria were absence of consent or insufficient clinical sample.

From January to December 2013, a total of 84 individual specimens were collected from 84 children up to the age of 1 month, who met the clinical inclusion criteria and were included in the study. Samples were 71 CSFs (85 %), 12 sera (14 %) and 1 throat swab (1 %). Mean age of the children was 13 ± 10 days and 46 of them were males (55 %).

**Viral detection and typing** Enterovirus (EV) and herpes simplex virus (HSV-1 and 2) and varicella zoster virus (VZV) screening was performed in the microbiology laboratory from the hospital where the patients were admitted or in the National Centre for Microbiology (CNM), Enterovirus Laboratory, using commercial molecular methods (Xpert EV, Cepheid, CA, USA) or “in-house” RT-PCRs as previously described [8, 21].

HPeV was tested at the CNM in those specimens negative for EV, HSV and VZV, using a real-time RT-PCR designed in the 5'-NCR of the genome [7]. Molecular typing of detected EV and HPeV was carried out by 3'-VP1 or VP3/VP1 amplification, respectively, and sequencing [6, 12].

**Statistical analysis** Clinical and laboratory characteristics of the HPeV-positive patients were compared with those of the children infected with EV. A descriptive analysis was performed, expressing qualitative variables as proportions and quantitative variables as mean and standard deviation or median and interquartile range when appropriate. The chi-square test was used to compare groups. A difference with p value <0.05 was considered to be significant.

**Results**

**Baseline data and frequency of EV and HPeV infections** Out of 84 infants included in the study, 32 were positive for EV (38 %). No HSV-1 and 2 and VZV were detected. The 52 negative specimens (47 CSFs, 4 sera and 1 throat swab) were tested for HPeV, with positive detection in 9 (17 %) of them. Overall, HPeV detection frequency was 11 %. During the clinical course, other diagnosis were made (urinary infection, bronchiolitis, gastroenteritis or non-infectious neurologic disease) for 13 of the infants, and those were finally excluded from the epidemiological and clinical data analysis; all were EV and HPeV negative (Fig. 1).

Viral detections were not found in any patient with suspected early-onset sepsis (11 newborns). In addition, four patients had bacteria positive blood culture, one *Staphylococcus epidermidis* (contaminated culture), two *Streptococcus agalactiae*, and one had a coinfection of *Pasteurella multocida* in blood and echovirus 9 in CSF. It was the only detected virus-bacteria coinfection [1].

**Epidemiological and clinical associations of HPeV infections** HPeV-positive samples were from four males and five females aged between 6 and 31 days (mean age, 17.6 days). Regarding monthly distribution, most of the HPeV infections (67 %) were detected between April and July, with another peak in October (22 %) (Fig. 2). All viruses were successfully characterized. Eight strains were HPeV-3 and the other was HPeV-5. All but one HPeV-positive specimens were found at CSFs. One of the HPeV-3 was detected in a serum sample. Clinically, six cases were diagnosed as FWS, two as clinical sepsis and one as encephalitis (Table 1). Fever was present in all cases (100 %), 38.6±0.4 °C and lasted a mean of 1.3±1.1 days. Although all except one of the HPeV were detected in CSF, none of the positive cases had cells in the range for pleocytosis in children <1 month of age (up to 30 cells/mm³). Protein and glucose were also normal in CSF. Leucocytes in blood were 8500±5696 cells/mm³. C-reactive protein and procalcitonine were normal in all patients. Four of the nine HPeV-infected patients (44 %) were admitted to the neonatal intensive care unit (NICU). One of them, diagnosed with encephalitis, had seizures. None of them needed mechanical ventilation, and all experienced total recovery except the infant with encephalitis who had persistent seizures. All patients received antibiotics until the bacterial aetiology could be ruled out.
Comparison between EV and HPeV infections

Similar to HPeV infections, 66% of the total of EV-positive samples were detected between April and July (Fig. 2). Twenty-nine of the 32 EV (91%) were successfully characterized. All serotypes found belonged to EV species B. Echovirus (E)-18 was the most frequently detected type (7/29, 24%), followed by E-3 and E-5 (4/29, 14% each) and coxsackievirus (CV)-B3 (3/29, 10%). Other types also detected in minor proportion were E-11, E-30, E-6, E-9, E-7, E-17 and CV-B2. The diagnoses for EV-infected patients were meningitis in 19 cases (59.4%), FWS in 12 cases (37.5%), and clinical sepsis in 1 case (3%). Clinical data of the infants are shown in Table 1.

When clinical characteristics of EV-infected patients were compared with those of HPeV-positive children, we observed significant differences in the pleocytosis in CSF ($250 \pm 415$ cells/mm³) $p=0.015$. The odds ratio (OR) for pleocytosis and EV infection was $OR=1.5$ (confidence interval CI:1.1–2.1). CSF protein were also higher in EV group ($104 \pm 117$ vs. 49 ±17 mg/dL, $p=0.03$). Furthermore, meningitis diagnosis was significantly more frequent in EV-positive patients than in HPeV-infected children (19/32 vs. 0/9, $p<0.001$). Out of the 32 EV-positive patients, 23 (72%) received antibiotic treatment. Only three EV-positive patients (9%) needed admission to the NICU and no sequelae were described in this group. However, HPeV infection was a risk factor for admission to the NICU (OR=$5.2$ (CI: 1.9–14)) (3/32 vs. 4/9, $p<0.001$).

### Table 1 Clinical characteristics of infants with human enterovirus (EV) and parechovirus (HPeV) infections

| Clinical feature               | EV (n=32) | HPeV (n=9) | $p$   |
|-------------------------------|-----------|------------|-------|
| Male                          | 17 (53 %) | 4 (44 %)   | NS    |
| Age average (days)            | 17.2±7.9  | 17.6±9     | NS    |
| Temperature >37.9 °C          | 31 (97 %) | 9 (100 %)  | NS    |
| Highest temperature           | 38.5±0.6  | 38.6±0.4   | NS    |
| Fever duration (days)         | 1.6±1.1   | 1.3±1.1    | NS    |
| Rash                          | 4 (12.5 %)| 1 (11 %)   | NS    |
| Antibiotic treatment          | 23 (72 %)| 9 (100 %)  | NS    |
| NICU admission                | 3 (9 %)   | 4 (44 %)   | 0.001*|
| Leucocytes (cells/mm³)        | 10147±4693| 8500±5696  | NS    |
| Serum CRP (mg/L)              | 10.8±2.7  | 2.6±3.9    | NS    |
| Procalcitonine (ng/mL)        | 0.8±2.7   | 0.2±0.9    | NS    |
| CSF cells/mm³                 | 250±415*  | 5.7±7      | 0.015*|
| CSF proteins (mg/dL)          | 104±117*  | 49±17      | 0.030*|
| CSF glucose (mg/dL)           | 53±10³    | 50±8       | NS    |
| Diagnosis                     |           |            |       |
| Fever without source          | 12 (37.5 %)| 6 (67 %)  | NS    |
| Clinical sepsis               | 1 (3 %)   | 2 (22 %)   | NS    |
| Meningitis                    | 19 (59.4 %)| 0         | 0.001*|
| Encephalitis                  | 0         | 1 (11 %)   | NS    |

Quantitative variables are expressed as mean and standard deviation

*a Excluded a patient coinfect with EV and *Pasteurella multocida*

NS not significant, NICU neonatal intensive care unit, CSF cerebrospinal fluid, CRP C-reactive protein, OR odds ratio, CI confidence interval

*p values <0.05
Discussion

We present a prospective, multicentre study performed during 1 year in Spanish infants up to 1 month of age with suspected sepsis or meningo-encephalitis or fever without source. EV was detected in 38 % of the patients and HPeV in 11 %. With respect to EV infections, other authors found EV incidences in neonates between 9.5 and 24 % [2, 16, 18, 26]. The high detection frequency observed in our series might be due to the study being performed in clinically ill infants with a high suspicion of viral infection and not as active surveillance. Besides, not all EV-negative samples collected by the hospitals could be included in the final study, due to absence of available samples or signed consent forms.

On the other hand, few HPeV epidemiological studies focused on newborns (infants <30 days) have been published. Most of them were performed in children (<18 years), and the presence of the virus in CSF varies from 2.3 to 7 %, whereas in the general population the percentage decreases to 0.5–0.8 % [9–11, 23]. As far as we know, there is only one study from Italy in which 7 (11.6 %) cases of EV and 3 (5 %) cases of HPeV infections were detected out of 60 neonates with suspected sepsis or neurological infection [18]. Our higher HPeV prevalence in the same type of patients (11 %) might be due to the fact that the Italian study [18] was conducted at the NICU, reducing the study population and leaving out the analysis of children who do not require admission to the NICU (56 % in our series).

In most of the previous studies, HPeV-3 infections have been described as clearly associated with central nervous system infections and sepsis-like illness in neonates and young infants [4, 15, 22]. Our data supported these findings, as HPeV-3 was also the predominant genotype identified, representing 89 % of the total HPeV detected. Only one strain was identified as HPeV-5, which is an infrequent type reported in the literature [9–11, 18, 23]. In this study, this type was detected in a case with FWS, so we cannot conclude that HPeV type 5 has a proved association with neurological or systemic diseases in newborns.

Regarding seasonal distribution, in Europe, HPeV infections are more common in spring and summer, similar to the EV infections [9–11, 23]. In our series, there is also a predominance of HPeV detection in spring and summer, although we have only 1 year of enrollment period. Besides, HPeV has been described to present in biannual cycles in Europe [9, 11, 15]. Previous data in Spain (not published) are consistent with those findings as the detection frequency was significantly higher in 2011 and 2013 than in 2012, but further longer surveillance studies are necessary to confirm the seasonality and periodicity of the HPeV-3 infections. This circulation pattern might also explain the high HPeV incidence found in this study, in comparison with the previously mentioned Italian one [18].

Recently, Sharp et al [23], in Kansas, found that up to 17 % of the CSF-positive samples of children (<18 years) tested between June and October 2009 were positive for HPeV, with HPeV-3 being the predominant type. Clinically, they described HPeV infections with irritability and longer fever, lower peripheral leucocytes, absence of pleocytosis, lower CSF protein and higher CSF glucose compared to EV infections.

In our series, some of these clinical characteristics were also observed as HPeV-infected patients had lower blood leucocytes, CSF protein and CSF cells values than those with EV infection. Furthermore, several studies including the American one mentioned above reported that admission to intensive care units were more frequent in HPeV-positive cases than in those infected by EV, but the prognosis was generally good [9, 22, 23]. We observed the same in our series. The higher proportion of HPeV-infected children who required admission to NICU was probably determined by the low age in an infant with an unwell appearance rather than by the clinical severity, as there was only one patient with encephalitis who had sequelae. In the literature, however, some fatal cases were described [19, 20], encephalitis being the major complication that caused sequelae and deaths.

Another recent study reported that between 50 and 100 % of HPeV-positive cases had been associated with erythematosous rash on the extremities, especially palms and soles, not always present in the first hours but during evolution [24]. This dermatologic manifestation would be very suspicious data of HPeV infection in neonates and very young infants with febrile syndrome. However, these data are not consistent in the literature; the rash might have gone unnoticed or showed a different prevalence depending on the area. In the present study, only 12.5 % of patients showed rash.

In summary, HPeV-3 infections should be suspected in infants younger than 1 month, with fever but without leucocytosis and pleocytosis in CSF. In these cases, HPeV screening in CSF or blood should be incorporated in the routine viral diagnosis to prevent unnecessary antibiotic treatment and prolonged hospitalization. The low number of samples included in this study was a limitation that affects the incidence data of the EV and HPeV infections, but not the results obtained in relation to the clinical and epidemiological differences found between both infections. Further studies could provide more information about the burden of HPeV infections in newborns.

Acknowledgments This study was partially supported by a grant from the Spanish National Health Institute PI12-00904.

Conflict of interest The authors declare that they have no conflict of interest and they do not have a financial relationship with the organization that sponsored the research.

Author’s contribution María Cabrerizo and Gloria Trallero performed the detection and genotyping of HPeV. María Cabrerizo and Cristina
Calvo designed the study, analysed the clinical data and wrote the manuscript. María José Pena Gregoria Megías, María Carmen Muñoz Almagro, María Pilar Romero, Sara Sanbonmatsu-Gámez, Mercedes Pérez-Ruiz, Manuel Omeñaca, Juan Garcia Costa, Antonio Moreno-Docón and Nuria Rabella performed the local virological study, drafted the initial manuscript and approved the final manuscript as submitted. Amaia Cilla Eva del Amo, Diana Roda, María José Santos-Muñoz, Ana Isabel Mensalvas and Cristina Calvo recruited the patients and performed the data collection, contributed to the discussion of the results, and approved the final manuscript as submitted.

Open Access This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made.

References

1. Aguado I, Calvo C, Wilhelm I, Pablo-Hernando ME, Medina MJ, Sáez-Nieto JA, Cabreroz M (2014) Sepsis and meningitis caused by pasturella multocida and echovirus 9 in a neonate. Pediatr Infect Dis J 33:1308–1309
2. Ahmad S, Dalwai A, Al-Nakib W (2013) Frequency of enterovirus detection in blood samples of neonates admitted to hospital with sepsis-like illness in Kuwait. J Med Virol 85:1280–1285
3. Baumgarte S, de Souza-Luna LK, Grywna K, Panning M, Drexler JF, Karsten C, Huppertz HI, Drosten C (2008) Prevalence, types, and RNA concentrations of human parechoviruses, including a sixth parechovirus type, in stool samples from patients with acute enteritis. J Clin Microbiol 46:242–248
4. Benschop KS, Schinkel J, Minnaar RP, Pajkrt D, Spanjerberg L, Kraakman HC, Berkhout B, Zaaijer HL, Beld MG, Wolthers KC (2006) Human parechovirus infections in Dutch children and the association between serotype and disease severity. Clin Infect Dis 42:204–210
5. Boavin G, Abed Y, Boucher FD (2005) Human parechovirus 3 and neonatal infections. Emerg Infect Dis 11:103–107
6. Cabreroz M, Echevarria JE, González I, de Miguel T, Trallero G (2008) Molecular epidemiological study of HEV-B enteroviruses involved in the increase in meningitis cases occurred in Spain during 2006. J Med Virol 80:1018–1024
7. Cabreroz M, Calvo C, Rabella N, Muñoz-Almagro C, del Amo E, Pérez-Ruiz M, Sanbonmatsu-Gámez S, Moreo-Docón A, Otero A, Trallero G et study group of Enterovirus and parechovirus infections in children under 3 years-old, Spain (PI12-00904) (2014) Design and validation of a real-time RT-PCR for the simultaneous detection of enteroviruses and parechoviruses in clinical samples. J Virol Methods 208:125–128
8. Casas I, Tenorio A, Echevarria JM, Klapper PE, Cleator GM (1997) Detection of enteroviral RNA and specific DNA of herpesviruses by multiplex genome amplification. J Virol Methods 66:39–50
9. Esposito S, Rahamat-Langendoen J, Ascolese B, Senatore L, Castellazzi L, Niesters HG (2014) Pediatric parechovirus infections. J Clin Virol 60:84–89
10. Fischer TK, Midgley S, Dalgaard C, Nielsen AY (2014) Human parechovirus infection. Denmark Emerg Infect Dis 20:83–87
11. Ghanem-Zoubi N, Shiner M, Shulman LM, Sofer D, Wolf D, Marva E, Kra-Oz Z, Shachor-Meyouhas Y, Averbuch D, Bechhor-Fellner A, Barkai G, Kinarty A, Gershstein V, Ephros M (2013) Human parechovirus type 3 central nervous system infections in Israeli infants. J Clin Virol 58:205–210
12. Harvala H, Robertson I, McWilliam Leitch EC, Benschop K, Wolthers KC, Templeton K, Simmonds P (2008) Epidemiology and clinical associations of human parechovirus respiratory infections. J Clin Microbiol 46:3446–3453
13. Harvala H, Simmonds P (2009) Human parechoviruses: biology, epidemiology and clinical significance. J Virol 45:1–9
14. Harvala H, McLeish N, Kondracka J, McIntyre CL, McWilliam Leitch EC, Templeton K, Simmonds P (2011) Comparison of human parechovirus and enterovirus detection frequencies in cerebrospinal fluid samples collected over a 5-year period in Edinburgh: HEV Type 3 identified as the most common picornavirus type. J Med Virol 83:889–896
15. Harvala H, Griffiths M, Solomon T, Simmonds P (2014) Distinct systemic and central nervous system disease patterns in enterovirus and parechovirus infected children. J Infect 69:67–74
16. Khetsuriani N, LaMonte A, Oberste MS, Pallansch M (2006) Neonatal enterovirus infections reported to the national enterovirus surveillance system in the United States, 1983–2003. Pediatr Infect Dis J 25:889–893
17. Piñeiro L, Vicente D, Montes M, Hernández-Dorronsoro U, Cilla G (2010) Human parechoviruses in infants with systemic infection. J Med Virol 82:1790–1796
18. Piralla A, Mariani B, Stronati M, Maroni P, Baldante F (2014) Human enterovirus and an enterovirus infections in newborns with sepsis-like illness and neurological disorders. Early Hum Dev 90(Suppl1):S75–S77
19. Schuffenecker I, Javouhey E, Gillet Y, Kugener B, Billaud G, Floret D, Lina B, Morfin F (2012) Human parechovirus infections, Lyon, France, 2008–10: evidence for severe cases. J Clin Virol 54: 337–341
20. Sedmak G, Nix WA, Jentzen J, Haupt TE, Davis JP, Bhattacharyya S, Pallansch MA, Oberste MS (2010) Infant deaths associated with human parechovirus infection in Wisconsin. Clin Infect Dis 50: 357–361
21. Selva L, Martinez-Planas A, Garcia-Garcia JJ, Casadevall R, Luaces C, Muñoz-Almagro C (2012) Comparison of an in-house real-time RT-PCR assay with a commercial assay for detection of enterovirus RNA in clinical samples. Eur J Clin Microbiol Infect Dis 31:715–719
22. Selvarangan R, Nzabi M, Selvaraju SB, Ketter P, Carpenter C, Harrison CJ (2011) Human parechovirus 3 causing sepsis-like illness in children from midwestern United States. Pediatr Infect Dis J 30:238–242
23. Sharp J, Harrison CJ, Puckett K, Selvaraju SB, Penaranda S, Nix WA, Oberste MS, Selvarangan R (2013) Characteristics of young infants in whom human parechovirus, enterovirus or neither were detected in cerebrospinal fluid during sepsis evaluations. Pediatr Infect Dis J 32:213–216
24. Shoji K, Komuro H, Miyata M, Miyairi I, Saitoh A (2013) Dermatologic manifestations of human parechovirus type 3 infection in neonates and infants. Pediatr Infect Dis J 32:233–236
25. Stanway G, Joki-Korpela P, Hyypia T (2000) Human parechovirus infections in Dutch children and the epidemiology and clinical significance. J Clin Virol 20:83–97
26. Verboon-Maciolek MA, Krediet TG, van Looon AM, Kaan J, Galama JM, Gerard R, Fleer A (2002) Epidemiological survey of neonatal non-polio enterovirus infection in the Netherlands. J Med Virol 66:241–245