RESEARCH ARTICLE

A European multicentre evaluation of detection and typing methods for human enteroviruses and parechoviruses using RNA transcripts

A. Hayes1 | D. Nguyen1 | M. Andersson2 | A. Antón3 | J.-L. Bailly4,5 | S. Beard6 | K. S. M. Benschop7 | N. Berginc8 | S. Blomqvist9 | E. Cunningham10 | D. Davis11 | J. L. Dembinski12 | S. Diedrich13 | S. G. Dudman14 | R. Dyrdak15,16 | G. J. A. Eltringham17 | S. Gonzalez-Goggia18 | R. Gunson19 | H. C. Howson-Wells20 | A. J. Jääskeläinen21 | F. X. López-Labrador22,23 | M. Maier24 | M. Majumdar25 | S. Midgley26 | A. Mirand27 | U. Morley28 | S. A. Nordbø29,30 | S. Oikarinen31 | H. Osman32 | A. Papa33 | L. Pellegrinelli34 | A. Piralla35 | N. Rabella36 | J. Richter37 | M. Smith34,38 | A. Söderlund Strand39 | K. Templeton40 | B. Vipond41 | T. Vuorinen42 | C. Williams43 | E. Wollants44 | K. Zakikhany45 | T. K. Fischer23,26 | H. Harvala46 | P. Simmonds1

1Nuffield Department of Medicine, University of Oxford, Oxford, UK
2Microbiology Laboratory, John Radcliffe Hospital, Headington, Oxford, UK
3Respiratory Viruses Unit, Virology Section, Microbiology Department, Hospital Universitari Vall d’Hebron, Vall d’Hebron Research Institute, Universitat Autònoma de Barcelona, Passeig Vall d’Hebron, Barcelona, Spain
4Université Clermont Auvergne, LMGE UMR CNRS, UFR Médecine, Clermont-Ferrand, France
5CHU Clermont-Ferrand, National Reference Center for EV and Parechovirus-Associated Laboratory, Clermont-Ferrand, France
6Enteric Virus Unit, Virus Reference Department, National Infection Service, Public Health England, London, UK
7National Institute for Public Health and the Environment (RIVM), Bilthoven, The Netherlands
8Department for Public Health Virology, National Laboratory of Health, Environment and Food, Ljubljana, Slovenia
9National Institute for Health and Welfare, Mannerheimintie, Helsinki, Finland
10Viapath Infection Sciences, St. Thomas’ Hospital, London, UK
11Microbiology, Virology and Infection Prevention & Control, Great Ormond Street Hospital for Children NHS Foundation Trust, London, UK
12Department of Virology, Norwegian Institute of Public Health, Oslo, Norway
13National Reference Center for Poliomyelitis and Enteroviruses, Robert Koch Institute, Berlin, Germany
14Department of Microbiology, Oslo University Hospital Rikshospitalet, Inst. Clinical Medicine, University of Oslo, Oslo, Norway
15Department of Clinical Microbiology, Karolinska University Hospital, Stockholm, Sweden
16Department of Microbiology, Tumor and Cell Biology, Karolinska Institute, Stockholm, Sweden
17Molecular Diagnostics Laboratory, Microbiology, Freeman Hospital, Newcastle Upon Tyne, UK
18Public Health England Poliovirus Reference Laboratory, National Infection Service, Public Health England, London, UK
19West of Scotland Specialist Virology Centre, Glasgow Royal Infirmary, Glasgow, UK
20Nottingham University Hospitals NHS Trust, Clinical Microbiology, Queens Medical Centre, Nottingham, UK
21University of Helsinki and Helsinki University Hospital, HUSLAB, Virology and Immunology, Helsinki, Finland

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Abstract

Polymerase chain reaction (PCR) detection has become the gold standard for diagnosis and typing of enterovirus (EV) and human parechovirus (HPeV) infections. Its effectiveness depends critically on using the appropriate sample types and high assay sensitivity as viral loads in cerebrospinal fluid samples from meningitis and sepsis clinical presentation can be extremely low. This study evaluated the sensitivity and specificity of currently used commercial and in-house diagnostic and typing assays. Accurately quantified RNA transcript controls were distributed to 27 diagnostic and 12 reference laboratories in 17 European countries for blinded testing. Transcripts represented the four human EV species (EV-A71, echovirus 30, coxsackie A virus 21, and EV-D68), HPeV3, and specificity controls. Reported results from 48 in-house and 15 commercial assays showed 98% detection frequencies of high copy (1000 RNA copies/5 µL) transcripts. In-house assays showed significantly greater detection frequencies of the low copy (10 copies/5 µL) EV and HPeV transcripts (81% and 86%, respectively) compared with commercial assays (56%, 50%; \( P = 7 \times 10^{-5} \)). EV-specific PCRs showed low cross-reactivity with human rhinovirus C (3 of 42 tests) and infrequent positivity in the negative control (2 of 63 tests). Most or all high copy EV and HPeV controls were successfully typed (88%, 100%) by reference laboratories, but showed reduced effectiveness for low copy controls (41%, 67%). Stabilized RNA transcripts provide an effective, logistically simple and inexpensive reagent for evaluation of diagnostic assay performance.
1 | INTRODUCTION

Enteroviruses (EVs) belonging to the family *Picornaviridae* are currently classified into 116 serologically distinct enterovirus types, which can be assigned into four genetically distinct species, HEV-A to D1,2 and over 250 human rhinovirus (HRV) types divided into three species (A-C). Species B EV types including echoviruses and coxsackieviruses (CBVs) are the most frequently identified viral causes of meningitis and other central nervous system (CNS)-associated infections in western countries, while the species A serotype, EV-A71 is an important cause of hand, foot, and mouth disease and encephalitis in South-East Asia and EV-D68 within species D has recently emerged as a respiratory pathogen occasionally leading to acute flaccid myelitis (AFM).3 Infections with human parechoviruses (HPeVs) in the genus *Parechovirus* are enteric, usually asymptomatic apart from those of HPeV type 3, which is associated with sepsis-like illness, meningitis, and encephalitis in young children.4-6

Although there is no effective antiviral treatment available for EV infections, detection and identification of EV and HPeV infections are vital for informing other treatment options, supportive care and prognosis of affected individuals. The reverse-transcriptase polymerase chain reaction (RT-PCR) is now the "gold standard" for diagnosing EV and HPeV infections due to its advantages of fast turn-around time and high sensitivity over virus isolation.7 Even in severe cases, viral loads are relatively low in cerebrospinal fluid (CSF) samples that are typically tested in patients presenting with meningitis or encephalitis, and may be missed by less sensitive methods.

To detect all EV types, RT-PCR assays for the detection of EV RNA usually target the highly conserved 5′ non-translated region (5′NTR). Depending on the primer and probe design, some molecular detection methods may fail to detect certain EV types such as EV-D68, whereas some assays may also detect HRVs (reviewed in Holm-Hansen).8 EV and HPeV serotypes are defined serologically and genetically by their capsid region sequences; virus typing, therefore, requires amplification and sequencing of regions within this structural gene block, typically VP1.8,9

Evaluation of sensitivity and specificity of the diagnostic assays used for EV and HPeV detection and typing is essential. We have previously evaluated the use of RNA transcripts of several EV and HRV serotypes and HPeV1 for quality control purposes in six expert clinical virology laboratories in Europe.10 Following this study, we have now produced a further set of RNA transcript standards for selected representative serotypes from EV species A-D and HPeV3. The RNA standards were distributed via the European non-polio enterovirus network (ENPEN) to members in diagnostic and reference laboratories for evaluation of the sensitivity of their routinely used assays for detection and typing of enteroviruses.

2 | MATERIALS AND METHODS

2.1 | RNA transcript synthesis

Available full-length cDNA clones of EV species A (EV-A71 genogroup B4 strain, accession number AF316321)11, B (Bastianni prototype strain of echovirus 30 [E30], AF162711), C (Coe strain of coxsackievirus A 21 [CVA21], D00538), and D (Fermon strain of EV-D68, NC_038308), and parechovirus (HPeV3, GQ183026) were selected for this study. For rhinovirus species C (HRV-C49, MF775365), a partial 5′-UTR-VP4-VP2 clone was assembled from amplified sequences. All plasmids were transformed into DH5α competent cells by heat shock, with single colonies picked and grown in liquid medium before plasmid extraction using the QIAprep Spin Miniprep Kit (Qiagen), according to the manufacturer's instructions. Plasmids were linearized at the 3′ end, and RNA transcripts were produced using the MEGAscript T7 Transcription Kit (Ambion), followed by DNase treatment to remove template DNA. RNA was purified using the RNA Easy Mini Kit (Qiagen), according to the manufacturer's instructions.

2.2 | RNA quantification and stability assessment

Quantification of RNA transcripts was carried out using the NanoDrop ND-1000 UV-Vis Spectrophotometer (Thermo Fisher Scientific) and the Qubit 4 Fluorometer (Thermo Fisher Scientific). The concentrations obtained were used to calculate copy numbers of transcripts produced, assuming a mean molecular mass for each base of 330 g/mol. A serial dilution of RNA transcripts (105 to 10−2 copies/µL) was prepared using the RNA storage solution (Thermo Fisher Scientific; 1 mM sodium citrate, pH 6.4) containing herring sperm carrier RNA (50 µg/mL) and RNasin (New England BioLabs UK, 100 U/mL). Dilutions were aliquoted and stored at −80°C before testing and distribution to the participating laboratories.

EV species A (EV-A71) and C (CVA21) transcripts were investigated for stability at different temperatures. Transcripts were incubated in storage solution for up to 30 days at ambient temperature, 4°C and 37°C. A further aliquot of each was...
freeze-thawed three times. The amount of RNA was quantified by RT-PCR and values compared to those of the original preparations.

2.3 | Transcript amplification by real-time RT-PCR

For quantification of RNA sequences before distribution, In-house quantitative real-time RT-PCR was carried out using the StepOne-Plus Real-Time PCR System (Thermo Fisher Scientific), according to the manufacturer’s instructions. The following reaction conditions were used: 50°C for 30 minutes, 95°C for 15 minutes followed by 45 cycles of 95°C for 10 seconds then 60°C for 1 minute. A total of 20 μL reaction volume containing 2 μL of the diluted transcript was used. PCRs used primers and probes as previously described for EV, HRV and HPeV (Table S1).

2.4 | Laboratory evaluation

RNA from EV species A-D and HPeV was distributed in 200 μL volumes of storage buffer at concentrations of 10 and 10^3 copies/5 μL. The HRV species C, as a negative transcript control, was distributed only at the higher concentration (10^3 copies in 5 μL) to investigate cross-reactivity of EV assays with rhinoviruses. Storage buffer was included as another negative transcript control. Sample labeling was coded, and the details of coding were provided to participating laboratories only after all results were received. Coded transcript panels were sent by standard registered post to the participating laboratories in February 2019.

2.5 | Participating laboratories

The RNA transcripts were distributed via ENPEN to member diagnostic and reference laboratories for evaluation of the sensitivity of their routinely used assays for detection and typing of EV and HPeV. Laboratories were identified by a standardized code (L1, L3...). Coded transcripts panels were shipped by standard registered UK post in February 2019. Participating labs were asked to test 5 μL of each sample sent using their routine detection and/or typing assay. Results were reported through an EU survey tool (https://ec.europa.eu/eusurvey/runner/ENPEN_transcript_study_panel_A_2019). Collected information comprised the results of testing in terms of positivity or negativity, Ct values were obtained, virus type and further technical information such as volume tested, extraction method (if used) and additional information on the testing methods used (listed in Supplementary Methods).

3 | RESULTS

3.1 | Validation of the transcript panel

RNA concentrations of EV-A71, E30, CVA21, EV-D68, HPeV3, and HRV-C49 transcripts were determined by two different physicochemical methods (Nanodrop and Qubit). The values obtained by the two methods’ values were similar (within a factor of two in all cases; data not shown) and the mean value was used for calibration. Serial dilutions of each RNA transcript ranging from 10^6 to 10^{-1} copies in 5 μL were assayed in replicate by quantitative EV and HPeV RT-PCR assay. Amplification was highly reproducible between replicates and between transcripts of different EV species and HPeV3 (Figure 1). All samples were positive with an input of 10 copies, while detection of single copies of RNA was stochastic (2/5 replicates detected). The end-point detection of the RNA transcripts and their similarity in Ct values to each demonstrates that quantitation was reproducible between the transcripts of the five different viral species (all Ct values were within a factor of two from each other). Finally, the measured gradients of the lines of best fit between log viral load and Ct-value were between 3.1 and 3.6, consistent with efficient amplification.

To investigate the stability of RNA transcript, two representative EV transcripts (EV-A71 and CVA21) were subjected to a range of temperatures and freeze-thaw cycles and their RNA content was assessed by real-time PCR (Figure 2). No or minimal changes in Ct values (reflecting residual RNA concentrations) were observed on freezing-thawing or incubation for up to 30 days at 4°C or ambient temperate, while there was an approximately 10-fold reduction in RNA levels on incubation at 37°C for 30 days. No decline in Ct values was observed for any transcript stored at 4°C or room temperate.
times ranged from one to 4 weeks providing reassurance for the RNA using standard registered post at ambient temperature; delivery 1000 copies in 5 µL), HRV SEMS of three assay repeats reductions of RNA relative to the starting amount; error bars show transcripts were additionally subjected to three freeze/thaw cycles were compared to those of RNA transcripts stored at for various durations at different temperatures. Detected viral loads were incubated for numerous at different temperatures. Detected viral loads were compared to those of RNA transcripts stored at −80°C. RNA transcripts were additionally subjected to three freeze/thaw cycles (rapid cooling and thawing; right hand panel). Bar heights show fold reductions of RNA relative to the starting amount; error bars show SEMS of three assay repeats over the month period, providing reassurance that transcripts received by the participating laboratories were not degraded.

A panel of 12 RNA transcripts was constructed containing two concentrations of EV-A71, E30, CVA21, EV-D68, and HPeV3 (10 and 1000 copies in 5 µL), HRV-C49 (1000 copies in 5 µL) and a water control. These panels were sent to the participating laboratories using standard registered post at ambient temperature; delivery times ranged from one to 4 weeks providing reassurance for the RNA quality.

3.2 | Participating laboratories

A total of 39 laboratories from 17 European countries including Belgium (number of participating laboratories = 1), Cyprus (1), Denmark (1), France (1), Finland (4), Germany (2), Greece (1), Italy (2), Ireland (1), Netherlands (1), Norway (3), Slovenia (1), Spain (3), Sweden (3), and UK (14) participated to this study. From these, 12 were classified as national reference laboratories and the remaining 27 as primary diagnostic laboratories (Table S2).

Most laboratories (n = 36) participated in the evaluation of detection assays, some evaluating multiple assays; this produced a total of 63 sets of results for detection assays. Reference Laboratories showed greater investment in in-house detection methods for EV and HPeV detection; in-house assays were used for 19 of the 22 results sets provided by 12 reference laboratories for the study (86%), with only one reference laboratory located within a local hospital contributing results from three different commercial assays). This compares 29/41 (71%) of in-house result sets provided by 27 diagnostic laboratories. Commercial kits used included assays from BioMérieux, Seegene, Progenie, Luminex, Fast Track Diagnostics, Elite Ingenius, Biofire, Altona, and AusDiagnostics (Table 1).

A total of 37 sets of in-house typing results were provided, from these 18 sets (14 for EV typing, 3 for HPeV, and 1 for both) were reported by reference laboratories and 12 sets by diagnostic laboratories from 7 different countries including Spain, Sweden, Greece, Italy, Germany, Norway, and UK. Most of these results were reported for EV typing (n = 29) but HPeV typing was also performed (n = 10).

3.3 | Sensitivity and specificity of screening methods

The sensitivity and specificity of detection assay results were calculated, and totals adjusted for the declared target range of the tests. Intended assay targets included combined EV and HPeV detection (n = 23), EV detection only (n = 17), HPeV detection only (n = 10) and combined EV and HRV detection (n = 7) as well as monospecific assays for EV-D68 (n = 1) and EV-A71 (n = 1). Two HRV-only assays were evaluated for specificity only.

In general, laboratories reported high rates of detection (98% for CVA21; 100% for EV-A71, E30, and EV-D68) of the EV transcripts at the higher concentration (10⁴ RNA copies in 5 µL) (Figure 3). More variable detection of the low concentration transcripts (10 RNA copies in 5 µL) was reported, ranging from 62% (EV-D68) to 90% (EV-A71). Detection frequencies of the HPeV3 transcripts were comparable; 97% for higher concentration and 75% for lower concentration. For assays reporting Ct values for the higher and lower concentration transcripts, values were compared to evaluate viral load ratios (Figure S1). Although no assay produced quantitative results, reported results showed a 58 to 106 fold differences in geometric mean viral loads, close to the expected 100-fold difference. Assays were therefore reasonably quantitative in relative terms in this concentration range.

Assays were also generally highly specific, with only 2 high Ct value (weak positive) results reported falsely positive from the 62 tests performed. A larger number of tests specific for EVs reported HRV detection, with five tests designed for the detection of EV (n = 2), EV and HPeV (n = 2) and EV-D68 (n = 1) reporting positive results with the HRV-C49 RNA transcript.

Methodology differences contributed substantially to the sensitivity of the screening assays (Figure 3B). In particular, commercial assays, often highly multiplexed for other viral targets in CSF, showed significantly reduced sensitivity for the detection of RNA transcripts at a lower concentration compared to in-house methods.
There was a comparable difference in HPeV detection rates, with 50% of the lower concentration transcript detected by commercial and 86% by in-house assays. There were significant differences in assay sensitivity in results reported by diagnostic and by reference laboratories (Figure 3C). This is largely accounted for by the greater use of in-house assays by reference laboratories.

Commercial assays included a variety of platforms and assay specificities (Table 1); several including Biofire, AusDiagnostics, Progenie and some assays from Seegene and bioMérieux were unable to detect the lower concentration RNA control (10 RNA copies in 5 µL), and in some cases, even the higher concentration RNA transcript (1000 copies in 5 µL) (Biofire, AusDiagnostics). We further investigated the sensitivity of the Biofire assay with intermediate RNA concentrations; assay sensitivity lay between 400 and 1000 RNA copies for most of the EV transcripts and between 1000 and 40 000 copies for HPeV RNA (Table 2).

### 3.4 Sensitivity and accuracy of EV and HPeV typing methods

EV and HPeV typing performed by 22 participating laboratories were based upon amplification by PCR and Sanger sequencing of VP1 and/or VP3/VP1 (EV) or VP3/VP1 (HPeV) regions either using species- or genus-specific assays. Half of the typing was performed at the reference laboratories and half at the diagnostic laboratories resulting in the use of 37 different typing assays (Figure 4). The higher concentration EV RNA transcripts were successfully typed by 88% of reference laboratories (45 of 51) and 71% of diagnostic laboratories (36 of 57; P = .02 by Fisher’s Exact Test) whereas the lower concentration EV transcripts were successfully typed by 41% of reference laboratories (21 of 51) and 31% of diagnostic laboratories (16 of 51; P = 0.2). High concentration HPeV RNA transcripts were successfully typed by all reference (3 of 3) and diagnostic laboratories (6 of 6) but one of each failed detection of the low copy HPeV transcript.

### 4 DISCUSSION

This study describes a quality control exercise for EV and HPeV detection and typing in 43 virus diagnostic and reference laboratories in 17 European countries. Intrinsic to the study design was the use of stabilized RNA transcripts of representative serotypes and species of EV and HPeV.

#### 4.1 The suitability of RNA transcripts for quality assurance

RNA transcripts were used to evaluate the sensitivity and specificity of commercial and in-house detection and characterization assays for EVs and HPeV. Multiple EV types were used in the panel to estimate the range of serotypes and species that could be detected; enteroviruses are substantially diverse genetically, and conserved
regions of the genome between EV species suitable for amplification by PCR are largely confined to short motifs in the 5′-UTR.\textsuperscript{13} EV and HPeV RNA transcripts possess a number of attributes that make them suitable as reagents for quality assurance (QA) and evaluation of diagnostic assays. Most importantly, their content can be quantified in absolute terms, so it is possible to express assay sensitivities in terms of RNA copies. This provides a stable quantitative standard for longer-term evaluation of assay sensitivities. Secondly, the transcripts are based on uniform, defined sequences enabling assay failures to be investigated through comparisons of primer/probe and target sequences. Thirdly, unlimited amounts of RNA transcripts can be generated from cDNA clones of each target guaranteeing long-term stability and reproducibility of the control materials. Fourthly, the RNA transcripts are highly stable at ambient temperature and on repeated freeze/thawing (Figure 2). Finally, they are noninfectious, enabling their international distribution by ordinary post, providing considerable cost savings, assured biological safety and simple logistics. RNA transcripts can indeed be very easily prepared on a large scale for little cost—in the current study, the consumable costs for laboratory production and packaging of the transcripts was approximately €120, while postage cost to the participating laboratories was approximately €390. This low cost enables participation in this and future RNA transcript-based QA exercises to be uncharged. The wide availability and insignificant cost will facilitate the use of RNA transcripts in the development of screening and typing capacity in Eastern Europe, where EV detection is largely restricted to virus isolation and neutralization assays.\textsuperscript{16}

There are however also some potential disadvantages to the use of RNA transcript controls and limitations of the study. Firstly, the RNA controls were directly used without RNA/DNA extraction in some test formats, so the efficiency of this step, which may be critical in clinical sample handling, and its downstream effects on overall assay performance was not evaluated for all assays. In practice, however, this can be evaluated through comparison of \( C_t \) values of transcript samples with and without an extraction step. The second potential disadvantage of RNA transcripts was the use of carrier RNA for stabilization. Although the concentrations used were relatively low in molecular terms (a 5 µL aliquot used for testing contained 250 ng of RNA), its presence could interfere with high throughput sequencing methods that may become increasingly used for virus typing in the

![Figure 3](image-url)  
**FIGURE 3** Detection frequencies of EV and HPeV transcripts. Frequencies of detection of the 10 and 1000 RNA copy/5 µL dilutions of each transcript by participant laboratories, divided by (A) transcript sequence, including detection frequencies of the HRV-C and water negative controls. The detection frequency of the EV-A71 transcript was significantly higher than achieved for the other EV species (B) Assay type—in-house for commercially available (results for individual commercial assays are shown in Table 1), and (C) laboratory type, diagnostic or national reference laboratory both shown separately for the EV and HPeV transcripts. \( P \) values above bars show frequency comparisons using Fisher’s Exact Test. EV, enterovirus; HPeV, human parechovirus; HRV-C, human rhinovirus species C

| TABLE 2 | Sensitivity of biofire film array assay |
|-----------------|-------------------|-------------------|-------------------|-------------------|
| **Transcript** | **RNA copies/5 µL** | **10** | **400** | **1000** | **40000** |
| EV-A71 | N\textsuperscript{1} | P\textsuperscript{2} | P | P |
| E30 | - | N | P | P |
| CAV21 | - | N | P | P |
| EV-D68 | - | N | P | P |
| HPeV3 | - | N | N | P |

\textsuperscript{1}Negative in assay.  
\textsuperscript{2}Positive in assay.  
\textsuperscript{3}Not done.
future. However, the RNA amount used was well within the capacity of current library preparation methods for Illumina and Nanopore methods, and indeed cDNA in these concentrations may be required for efficient template preparation. We are currently investigating the stabilizing properties of alternative carriers such as linear acrylamide to avoid this currently theoretical problem in the future. Finally and more generally, RNA transcripts or cloned viral DNA sequences are available from a relatively narrow range of viruses, typically those that possess relatively small, non-segmented genomes, and for which full-length cDNA clones representative of currently circulating strains are available. Although RNA transcripts for EVs and HPeV used in the current study can be readily derived from cloned sequences of a wide range of contemporary circulating virus strains, development of equivalent RNA (or DNA)-based standards would be problematic or impossible for many respiratory and enteric viruses, and for adenoviruses, herpesviruses and other large DNA viruses with genomes that are too large for conventional cloning strategies.

4.2 Analytical sensitivity of EV and HPeV detection assays

As described, the transcripts enabled assay sensitivity to be determined in absolute RNA copy numbers, with almost all assays able to detect 1000 copies of RNA transcripts but with more variable detection of 10 RNA copy controls for EVs and HPeV. There are no current statutory guidelines for EV or HPeV detection sensitivity for diagnostic assays although previous evaluations of widely used EV PCRs found detection limits of 10 to 50 copies for EV and HPeV.

This corresponds to 50 to 200 RNA copies/mL using standard 200 µL extraction volumes of CSF. In the current study, positive results from the low copy number EV or HPeV controls equated to an analytical sensitivity of greater than 50 EV/HPeV RNA genomes/mL for a standard 200 µL CSF extraction volumes. Almost all reference laboratories achieved this sensitivity using in-house PCR methods, but the 10 copy control was frequently negative in testing with commercial assays. In the case of the biofire film array assay, the mean limit of detection for the 4 EV species was greater than 2000-4000 RNA copies/mL, with an even lower sensitivity for HPeV3 (Table 2). These findings are consistent with previous evaluations of the Biofire film assay, with reported limits of detection for EV detection of >500 RNA copies/mL, and some reduction in rates of detection of (unquantified) EV-positive clinical samples compared to conventional diagnostic assays.

Viral loads in CSF are low, often at the limit of assay sensitivity of PCR, so variability in assay sensitivity could substantially influence diagnostic target detection rates in diagnostic samples. The development of guidelines for assay sensitivity for EV and HPeV RNA detection in CSF would be of considerable value in the future quality control of these assays. The legislation for the use of CE-marked tests is driving the replacement of in-house assays with commercial tests, many of which offer syndromic testing with multiplexed detection of a large range of viruses and bacteria. Although such assays often have operational advantages as point-of-care tests in emergency rooms requiring rapid results, the effects of their potentially reduced sensitivity on their clinical utility need to be evaluated. Validation of their performance can be challenging for multiple analytes and existing studies typically do not include samples with defined viral loads or identified EV species or serotypes, nor investigation of samples that have failed detection. In the future, validation of such assays with calibrated RNA controls is of particular value for evaluation of their performance, particularly when the availability of control material derived from traditional virus isolation methods becomes increasingly restricted.
4.3 Virus typing

In the current study, EV and HPeV typing were performed in both reference and diagnostic laboratories (Figure 4). Although all results reported the correct EV-type identification and most laboratories successfully typed the high concentration (1000 copies in 5 µL) transcripts, fewer than 50% of either reference or diagnostic laboratories could successfully amplify and sequence the low concentration controls. The observed restriction in assay sensitivity underpins the importance of obtaining multiple samples including blood, respiratory samples and feces for EV diagnostic and typing assays where viral loads are higher during acute infections.22-24 It is also time to consider how EV and HPeV typing data can be centrally collected and analyzed at the time when increasing numbers of diagnostic laboratories are starting to introduce typing within the hospital premises.7,16

In conclusion, effective EV and HPeV detection and type identification are integral to clinical management, public health surveillance and outbreak preparedness for emerging strains. However, their genetic diversity, and often low viral loads in diagnostic specimens places stringent demands on the analytical sensitivity and breadth of detection and typing assays. RNA transcripts provide the means to independently evaluate these aspects of their performance. In the future, they can provide objective and fixed standards needed for a more critical assessment of the effectiveness of the numerous, newly developed and currently largely unevaluated testing platforms for syndromic testing. We would be delighted to provide EV, HPeV and further RNA transcript controls for a wider range of viruses to laboratories for QA purposes in the future.

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REFERENCES

1. Stanway G, Brown F, Christian P, et al. Family Picornaviridae. In: Fauquet CM, Mayo MA, Maniloff J, Desselberger U, Ball LA, eds. Virus Taxonomy. Eighth Report of the International Committee on Taxonomy of Viruses. London: Elsevier/Academic Press; 2005:757-777.

2. Mulders MN, Salminen M, Kalkkinen N, Hovi T. Molecular epidemiology of coxsackievirus B4 and disclosure of the correct VP1/2A(pro) cleavage site: evidence for high genomic diversity and long-term endemicity of distinct genotypes. J Gen Virol. 2000;81(Pt 3):803-812.

3. Holm Hansen CC, Midgley SE, Fischer TK. Global emergence of enterovirus D68: a systematic review. Lancet Infect Dis. 2016;16(5): e64-e75.

4. Boivin G, Abed Y, Boucher FD. Human parechovirus 3 and neonatal infections. Emerg Infect Dis. 2005;11(3):103-105.

5. Benschop KSM, Schinkel J, Minnaar RP, et al. Human parechovirus infections in Dutch children and the association between serotype and disease severity. Clin Infect Dis. 2006;42(2):204-210.

6. Harvala H, Robertson I, McWilliam Leitch C, Chieochansin T, Templeton K, Simmonds P. Aetiological role of human parechovirus type 3 in neonatal sepsis identified by direct typing assay on cerebrospinal fluid. J Infect Dis. 2009;199(12):1753-1760.

7. Harvala H, Broberg E, Benschop K, et al. Recommendations for enterovirus diagnostics and characterisation within and beyond Europe. J Clin Virol. 2018;101:11-17.

8. Nix WA, Oberste MS, Pallansch MA. Sensitive, seminested PCR amplification of VP1 sequences for direct identification of all enterovirus serotypes from original clinical specimens. J Clin Microbiol. 2006;44(8):2698-2704.

9. Harvala H, Robertson I, McWilliam Leitch EC, et al. Epidemiology and clinical associations of human parechovirus respiratory infections. J Clin Microbiol. 2008;46:3446-3453.

10. McLeish NJ, Witteveldt J, Clasper L, et al. Development and assay of RNA transcripts of enterovirus species A to D, rhinovirus species A to C, and human parechovirus: assessment of assay sensitivity and specificity of real-time screening and typing methods. J Clin Microbiol. 2012;50(9):2910-2917.

11. Tan CW, Tee HK, Lee MH, Sam IC, Chan YF. Enterovirus A71 DNA-launched infectious clone as a robust reverse genetic tool. PLoS One. 2016;11(9):e0162771.

12. Benschop KSM, de Vries M, Minnaar RP, et al. Comprehensive full length sequence analyses of human parechoviruses; diversity and recombination. J Gen Virol. 2010;91:151-154.

13. Dierssen U, Rehren F, Henke-Gendo C, Harste G, Heim A. Rapid routine detection of enterovirus RNA in cerebrospinal fluid by a one-step real-time RT-PCR assay. J Clin Virol. 2008;42(1):58-64.

14. Scheltinga SA, Templeton KE, Beersma MF, Claas EC. Diagnosis of human metapneumovirus and rhinovirus in patients with respiratory tract infections by an internally controlled multiplex real-time RNA PCR. J Clin Virol. 2005;33(4):306-311.

15. Benschop K, Molenkamp R, van der Ham A, Woltmers K, Beld M. Rapid detection of human parechoviruses in clinical samples by real-time PCR. J Clin Virol. 2008;41(2):69-74.

16. Harvala H, Jasir A, Penttinen P, Pastore Centelano L, Greco D, Broberg E. Surveillance and laboratory detection for non-polio enteroviruses in the European Union/European Economic Area, 2016. Euro Surveill. 2017;22(45):1-13.

17. Bennett S, Harvala H, Witteveldt J, et al. Rapid simultaneous detection of enterovirus and parechovirus RNAs in clinical samples by one-step real-time reverse transcription-PCR assay. J Clin Microbiol. 2011;49(7):2620-2624.

18. Piccirilli G, Chiereghin A, Gabrielli L, et al. Infectious meningitis/encephalitis: evaluation of a rapid and fully automated multiplex PCR in the microbiological diagnostic workup. New Microbiol. 2018;41(2):118-125.

19. Leber AL, Everhart K, Balada-Llasat JM, et al. Multicenter evaluation of BioFire filmarray meningitis/encephalitis panel for detection of bacteria, viruses, and yeast in cerebrospinal fluid specimens. J Clin Microbiol. 2016;54(9):2251-2261.

20. Messacar K, Breazeale G, Robinson CC, Dominguez SR. Potential clinical impact of the film array meningitis encephalitis panel in children with suspected central nervous system infections. Diagn Microbiol Infect Dis. 2016;86(1):118-120.

21. Liesman RM, Strasburg AP, Heitman AK, Theel ES, Patel R, Binnicker MJ. Evaluation of a commercial multiplex molecular panel for diagnosis of infectious meningitis and encephalitis. J Clin Microbiol. 2018;56(4):1-9. e01927-17.

22. Harvala H, Griffiths M, Solomon T, Simmonds P. Distinct systemic and central nervous system disease patterns in enterovirus and parechovirus infected children. J Infect. 2014;69(1):69-74.

23. Cordey S, Schibler M, L’Huillier AG, et al. Comparative analysis of viral shedding in pediatric and adult subjects with central nervous system-associated enterovirus infections from 2013 to 2015 in Switzerland. J Clin Virol. 2017;89:22-29.

24. Lafolie J, Labbé A, L’Honneur AS, et al. Assessment of blood enterovirus PCR testing in paediatric populations with fever without source, sepsis-like disease, or suspected meningitis: a prospective, multicentre, observational cohort study. Lancet Infect Dis. 2018;18(12):1385-1396.

SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.