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Whole Genome Sequencing of Enteroviruses Species A to D by High-Throughput Sequencing: Application for Viral Mixtures

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Human enteroviruses (EV) consist of more than 100 serotypes classified within four species for enteroviruses (EV-A to -D) and three species for rhinoviruses, which have been implicated in a variety of human illnesses. Being able to simultaneously amplify the whole genome and identify enteroviruses in samples is important for studying the viral diversity in different geographical regions and populations. It also provides knowledge about the evolution of these viruses. Therefore, we developed a rapid, sensitive method to detect and genetically classify all human enteroviruses in mixtures. Strains of EV-A (15), EV-B (40), EV-C (20), and EV-D (2) viruses were used in addition to 20 supernatants from RD cells infected with stool extracts or sewage concentrates. Two overlapping fragments were produced using a newly designed degenerated primer targeting the conserved CRE region for enteroviruses A-D and one degenerated primer set designed to specifically target the conserved region for each enterovirus species (EV-A to -D). This method was capable of sequencing the full genome for all viruses except two, for which nearly 90% of the genome was sequenced. This method also demonstrated the ability to discriminate, in both spiked and unspiked mixtures, the different enterovirus types present.

Keywords: human enteroviruses, whole-genome sequences, high-throughput sequencing, viral mixtures, enterovirus identification

INTRODUCTION

The common human viruses, human enteroviruses (EV), consist of more than 100 serotypes most classified within four species for enteroviruses (EV-A to -D) and three species for rhinoviruses. Enteroviruses are ubiquitous and resilient in the environment and primarily transmitted fecal-orally. Human enteroviruses have been implicated in a variety of human diseases, including the common cold, hand foot and mouth disease, acute hemorrhagic conjunctivitis, myocarditis, encephalitis, and poliomyelitis.

Enteroviruses are non-enveloped viruses, approximately 7500 nucleotides (nt) in length with a positive, single-stranded RNA genome. There are two untranslated regions (5′ and 3′-UTR) flanking a large open reading frame encoding a polyprotein that is cleaved to give three precursors...
Viral mixtures. Enteroviruses of species A to D present in samples containing of viral typing and sequencing the whole genome for all enteroviruses can be found in human stool or sewage. Sequencing of enterovirus A71 was published (Tan et al., 2015). Additionally, we used 20 enterovirus positive supernatants from human rhabdomyosarcoma cells (RD cells) from the Madagascar National Polio Laboratory. These cells were infected with either 10 human stool extracts or 10 sewage concentrates collected during poliomyelitis surveillance in Madagascar and prepared according to WHO protocols (World Health Organization Polio Laboratory manual, 4th edition, 2004). Infections of RD cells with these extracts were followed by full cytopathogenic effect, resembling that induced by enteroviruses.

For the collection of stool from Madagascar, the protocol was approved by the National Ethics Committee of the Ministry of Public Health in Madagascar (Agreement N° 22-MSANP/CE). Parental written informed consent was given.

In addition to typing, studies dedicated to evaluating the enterovirus genomic diversity have shown that besides mutations, intra- and inter-typic genetic recombination is also a frequent mechanism of viral evolution (Lukashev et al., 2003; Simmonds, 2006; Combelas et al., 2011; Joffret et al., 2012). Enterovirus genomes frequently display mosaicism due to genetic exchanges among different enterovirus strains and types. Since this kind of genetic diversification can be at the source of genotypic and phenotypic diversity (Sadeuh-Mba et al., 2013; Bessaud et al., 2016) it is essential to determine the whole genomic sequences of enteroviruses for surveillance and public health purposes, as well as for basic research.

Next generation sequencing (NGS) methods offer a new powerful sequencing tool for the identification and characterization of enteroviruses. This has been successfully used to sequence partial or whole genome sequences of poliovirus and of enteroviruses species C (Bessaud et al., 2016; Montmayeur et al., 2017; Sahoo et al., 2017). Additionally, a generic assay for whole genome amplification and deep sequencing of enterovirus A71 was published (Tan et al., 2015). Despite advances in molecular methods, none of these assays were designed to simultaneously amplify the whole genome for all four enterovirus species, and it is known that mixtures of enteroviruses can be found in human stool or sewage. Hence, the goal of this research was to design generic and species-specific primers in order to develop an assay capable of viral typing and sequencing the whole genome for all enteroviruses of species A to D present in samples containing viral mixtures.

MATERIALS AND METHODS

Viruses

For this study 15-EVA, 40-EBV, 20-EVC, and 2-EVD viruses were used (Table 1) to test and analyze whole-genome sequencing. The prototype strains selected were from the European Virus Archive (EVAg)1. Additionally, we used 20 enterovirus positive supernatants from human rhabdomyosarcoma cells (RD cells) from the Madagascar National Polio Laboratory. These cells were infected with either 10 human stool extracts or 10 sewage concentrates collected during poliomyelitis surveillance in Madagascar and prepared according to WHO protocols (World Health Organization Polio Laboratory manual, 4th edition, 2004). Infections of RD cells with these extracts were followed by full cytopathogenic effect, resembling that induced by enteroviruses.

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Primer Design

For each enterovirus group, complete genome sequences were separately aligned using CLC Main Workbench. To achieve the two overlapping fragments, the first half of the genome was amplified using the previously described C004 primer (Bessaud et al., 2016) in combination with a newly designed degenerated primer targeting the conserved CRE (cis-acting replication element) region for enteroviruses A-D. The second half of the genome was amplified using one degenerated primer set designed to specifically target the conserved region for each enterovirus species. Therefore, we performed five PCR mixes for each isolate and we pooled the PCR products for sequencing. All primers used in this study are described in Table 2 and Figure 1 illustrates the primer locations.

RNA Extraction

Viral RNA was extracted from supernatants of infected cells, using the High Pure Viral RNA kit (Roche Diagnostics, Meylan, France) per manufacturer’s protocol. All RNA was either immediately used for PCR amplification or stored at −80°C for further analysis.

One-Step RT-PCR Amplification

For entire length genome amplification, we used two primer sets (Table 2) per enterovirus group (A, B, C, and D). This allowed us to synthesize the two overlapping amplicons by using the One-Step RT-PCR kit (ref G174 Applied Biological Materials Inc.). The reaction mixture contained 1.5 µl of purified RNA, 12.5 µl of 2× One-step RT-PCR buffer, 1 µl of each forward and reverse primer (20 µM), 0.5 µl of EasyScript RTase (200 U/µl), 1 µl of Bestaq DNA Polymerase (5 U/µl), and 9 µl of DNAse free water. PCR amplification was performed using a thermodenat with the following protocol: 42°C for 30 min, 94°C for 3 min, 35 cycles

1https://www.european-virus-archive.com
at 94°C for 30 s, 55°C for 30 s, 72°C for 4 min 30 s, and a last step at 72°C for 10 min, ending with 2 min at 4°C. All PCR products were visualized on ethidium bromide-stained agarose gels to ensure appropriate size products.

The sensitivity of this assay was evaluated using serial fourfold dilutions of four viruses (EV-A71 strain MAD-72341, CV-B4 strain E2, CV-A13 strain MAD 67001, and EV-D 68 strain Fermon) representing each species (Table 3). To maintain similar amounts of cellular nucleic acids throughout dilutions, viral stocks were diluted using a supernatant of confluent non-infected HEp-2c cell monolayers frozen and thawed twice and clarified by centrifugation.

This RT-PCR amplification method was developed for the analysis of certain amount of viruses present following amplification in infected cells. It could probably be applied to the direct analysis of human or environmental samples, including different compartment-specific human fluids, provided that the sensitivity of the method has been adapted to the amount of virus present in these samples.

**Virus Mixture Detection**

To confirm the ability to detect viruses in a mixture, different samples containing known viral isolates under the following conditions were prepared: (1) equal amounts of four viral isolates representing enterovirus species A, B, C, and D and (2) four viral isolates belonging to enterovirus species B. The viral isolates used to perform these mixture experiments are listed in Table 4.

To confirm the ability to detect a mixture of viruses from “real life” conditions, we used 20 supernatants from RD cells which were infected with 10 stool extracts and 10 sewage concentrates, respectively (Table 5). RNA extraction, RT-PCR, and NGS sequencing were performed as described in the related sections.

**PCR Purification and Next Generation Sequencing (NGS)**

For each sample, the five PCR products were pooled and purified using a vacuum method and then sent to the sequencing platform PIBNET (Pasteur International Bioresources Network, Institute Pasteur Paris). The libraries were created using 1 ng of DNA with the Nextera XT DNA Library Preparation Kit in a SureCycler 8800 thermocycler (Agilent). Following purification on AMPure beads (Beckman), the libraries were controlled using the High Sensitivity D1000 assay (Agilent) on a TapeStation 2200. The products were sequenced using Illumina NextSeq HiSeq. All kits were used following manufacturer's instructions.

**Data Analysis**

Using CLC Genomics Workbench 8.5 (CLCbio), we paired and assembled contigs from the raw reads. Next, de novo assembly was performed using CLC Main Workbench (CLCbio) with the following parameters: Mismatch cost = 2; Insertion cost = 2; Deletion cost = 2; Length Fraction = 0.5; Similarity Fraction = 0.95. All contigs longer than 200 nt were submitted to National Center for Biotechnology Information (NCBI) for BLAST analysis.

**RESULTS**

**Sequence Analysis and Primer Design**

To sequence all EVs, we designed primers targeting conserved genomic regions that allowed the synthesis of overlapping amplicons. To amplify the first half of the genome primer C004 (Bessaud et al., 2016) was used in combination with a newly designed generic primer (EV-CRE-R), targeting the CRE region. This primer set was used to amplify the first half of the genome for the four EV species (EV A-D). To ensure the best amplification of the second half of the genome, we designed primers specifically targeting species A, B, C, and D. The combination of the two primer sets (Table 2), allowed for the amplification of the 5′ and 3′ parts of the genomes, which led to the synthesis of two overlapping DNA fragments per virus (approximately 400-nt long see Figure 1). This method resulted in the amplification of the whole genome for samples containing a single virus or mixture and the amplicon products were used for NGS.
To validate the proposed primer sets capability of generating whole genome sequencing data, we tested 15-EVA, 40-EVB, 20-EVC, and 2-EVD viruses (Table 1). We obtained the full genome sequencing data for all viruses except two (EV-C95 T08-083 and EV-D70), for which we obtained 93.6 and 87.5% of the genomic sequences, respectively. In all cases the sequences of the VP1 capsid protein could be used to confirm the type of virus. These results indicated the effectiveness of the proposed primers in conjunction with NGS sequencing to correctly identify the viruses and to reconstruct the whole genome using de novo assembly.

Additionally, to evaluate the sensitivity of this assay, we tested a fourfold serial dilution of one representative of each enterovirus species (Table 3). For undiluted viruses, the number of reads that mapped against the reference strains ranged between 86.9 to 99.3%, depending on the species. The full-length genome was reconstructed, using de novo assembly, for the majority of viruses and dilutions (ranging from 1.0e8 to 1.5e3 TCID50/ml). For some strains, long contigs overlapping almost the whole genomic sequences were recovered when high amounts of viruses were used (1.6e5 for CV-B4 and 1.6e6 for CV-A13).

**TABLE 1 | Viruses used in this study.**

| Type       | Strain or isolate | n° Databank | Consensus length |
|------------|------------------|-------------|-----------------|
| **SPECIES A** |                  |             |                 |
| CV-A2      | Fleetwood        | AY421760    | 7411            |
| CV-A3      | Olson            | AY421761    | 7411            |
| CV-A6      | Gdula            | AY421764    | 7551            |
| CV-A6      | MAD-2628-11      | LT719047    | 7503            |
| CV-A7      | Parker           | AY421765    | 7395            |
| CV-A10     | Kowalik          | AY421767    | 7420            |
| CV-A10     | MAD-9856-11      | LT719059    | 7402            |
| CV-A10     | MAD-3995-11      | LT719056    | 7415            |
| CV-A12     | Texas-12         | AY421768    | 7623            |
| CV-A14     | G14              | AY421769    | 7540            |
| CV-A14     | MAD-2718-11      | LT719062    | 7521            |
| CV-A16     | G10              | U05074      | 7500            |
| EV-A71     | MAD-3126-11      | LT719063    | 7400            |
| EV-A71     | MAD-72341-04     | LT719065    | 7326            |
| EV-A71     | CAE-146-08       | LT719066    | 7511            |
| **SPECIES B** |                  |             |                 |
| CV-A9      | RO-609-4-80      | LS451285    | 7398            |
| CV-B1      | RO-98-1-74       | LS451286    | 7410            |
| CV-B2      | Ohio             | AF081485    | 7350            |
| CV-B3      | RO-123-1-95      | LS451287    | 7519            |
| CV-B3      | RO-69-1-89       | LS451288    | 7327            |
| CV-B4      | E2               | AF311399    | 7173            |
| CV-B4      | RO-69-1-86       | LS451289    | 7351            |
| CV-B5      | RO-14-5-70       | LS451290    | 7273            |
| CV-B5      | Faulkner         | AF114383    | 7404            |
| CV-B6      | Schmitt          | AF039205    | 7209            |
| CV-B6      | RO-86-1-73       | LS451291    | 7345            |
| E-1        | Farouk           | AF029859    | 7313            |
| E-1        | RO-122-1-74      | LS451292    | 7351            |
| E-3        | Morrissey        | AY302553    | 7366            |
| E-4        | Pesacek          | AY302557    | 7359            |
| E-5        | Noyce            | AF083069    | 7340            |
| E-6        | RO-79-2-71       | LS451293    | 7444            |
| E-6        | D’Amor           | AY302558    | 7436            |
| E-6        | RO-24-9-79       | LS451294    | 7421            |
| E-7        | RO-434-2-81      | LS451295    | 7355            |
| E-7        | RO-141-2-95      | LS451296    | 7386            |
| E-9        | Hill             | X64981      | 7379            |
| E-9        | RO-116-6-82      | LS451297    | 7369            |
| E-11       | Gregory          | X80059      | 7564            |

**SPECIES B continued**

| Type       | Strain or isolate | n° Databank | Consensus length |
|------------|------------------|-------------|-----------------|
| E-11       | RO-91-91         | AJ577594    | 7460            |
| E-12       | RO-78-3-74       | LS451298    | 7351            |
| E-13       | DelCarmen        | AY302539    | 7410            |
| E-14       | Tow              | AY302540    | 7352            |
| E-14       | RO-81-1-79       | LS451299    | 7457            |
| E-15       | CH-96-51         | AY302541    | 7364            |
| E-20       | J-1              | AY302546    | 7321            |
| E-21       | Farina           | AY302547    | 7000            |
| E-24       | DeCamp           | AY302548    | 7239            |
| E-25       | JV-4             | AY302549    | 7443            |
| E-26       | Coronel          | AY302550    | 7459            |
| E-27       | Bacon            | AY302551    | 7339            |
| E-29       | JV-10            | AY302552    | 7362            |
| E-30       | Bastianni        | AY311938    | 7440            |
| E-32       | PR-10            | AY302555    | 7448            |
| EV-B69     | Toluca-1         | AY302560    | 7338            |

**SPECIES C**

| Type       | Strain or isolate | n° Databank | Consensus length |
|------------|------------------|-------------|-----------------|
| CV-A1      | Tompkins         | AF499635    | 7351            |
| CV-A11     | Belgium          | AF499636    | 7326            |
| CV-A11     | MAD-66122        | JF260917    | 7296            |
| CV-A11     | MAD-66990        | JF260918    | 7000            |
| CV-A13     | Flores           | AF655511    | 7368            |
| CV-A13     | G13              | AF499640    | 7280            |
| CV-A13     | MAD-67001        | JF260920    | 7298            |
| CV-A13     | MAD-67900        | JF260921    | 7000            |
| CV-A17     | G12              | AF499639    | 7301            |
| CV-A17     | MAD-67610        | JF260924    | 7388            |
| CV-A17     | MAD-68154        | JF260925    | 7541            |
| CV-A19     | 8663             | AF499641    | 7449            |
| CV-A20     | IH35             | AF499642    | 7415            |
| CV-A20a    | Tulane           | X87601      | 7305            |
| CV-A20b    | Cecil            | X87602      | 7360            |
| CV-A21     | Coe              | D00588      | 7393            |
| CV-A24     | DOLU-054         | JX417873    | 7366            |
| CV-C99     | MAD-69412-03     | LS451300    | 7438            |
| CV-C99     | MAD-69558-03     | LS451301    | 7383            |

**SPECIES D**

| Type       | Strain or isolate | n° Databank | Consensus length |
|------------|------------------|-------------|-----------------|
| EV-D68     | Fermon           | AY426531    | 7369            |
| EV-D70     | J670/71          | D00820      | 6467            |
**Mixture Detection**

To assess the capacity of the method to detect and identify different enteroviruses in samples containing mixtures, we performed two types of experiments: (1) mixed samples containing different mixtures of known viral isolates under controlled conditions (spiked) and (2) supernatants of cells infected with stool or sewage extracts in which the mixture of viruses was unknown and the conditions were not controlled (unspiked).

**Spiked Experiments**

To simulate true clinical and/or environmental conditions, four representative viral isolates for each enterovirus species were used to make mixture 1 (similar quantities or viral RNA); and four viral isolates of species B were used for mixtures 2 and 3. For all mixtures, RNA was extracted and DNA amplification was performed using the newly designed primer sets.

For mixture 1, we were successful in generating one contig per virus with 99.9% identity with the corresponding sequences in GeneBank using BLAST algorithm (Table 4). We were able to recover data about the four viral isolates and to consider the VP1 region used to identify the virus(es) present, either in its full or partial length.

For mixtures 2 and 3 we were able to correctly isolate and identify all four serotypes from species B with 99.7–99.9% identity and we recovered the full length of the VP1 region for the four B viral isolate strains under both conditions (half and full genome amplification). Additionally, for all three experiments, we confirmed that the parameters used by de novo could assemble the reads into separate contigs, resulting in the expected viruses.

**Unspiked Experiments**

To validate this method for field studies, we tested unknown mixtures of viral isolates present in the supernatant of RD cells infected with extracts from stools and sewage samples. Ten samples from stools and ten samples from sewage were used, following the protocols described in the materials and methods section. Full genomic sequences were identified for 8 viruses present in RD supernatants from stool samples and 5 viruses present in those from sewage samples. However, for certain viruses, contigs covering only partial genomic sequences could be obtained. In these cases, we could use the VP1 contigs to identify the viruses present. In cell supernatants of infected cells, we were able to recover 1 to 2 viruses per stool sample and 2 to 7 viruses per sewage sample (Table 5). Among twenty samples, we were able to find 48 enteroviruses based on the VP1 region. For 21 viruses, more than 6100 nt per genome could be determined, whereas, for 27 other viruses, genomic contigs were not as long. The genome of 18 viruses was higher than 90% in length. Overall, 41 EV-Bs, 3 EV-As, 3 EV-Cs, and no EV-Ds were identified.

**DISCUSSION**

The goal of this research was to genetically sequence the entire genome for all enteroviruses (EV-A, -B, -C, and -D) present in a given sample, using a simple method for detection and genetic characterization. To accomplish this, we developed an RT-PCR assay where we designed degenerate primers targeting conserved regions of the genome, which allowed the DNA fragments to be amplified, followed by NGS. This paper described our approach and demonstrated the feasibility of our methods to successfully identify EV in mixtures.

The effectiveness of this assay was achieved by selecting two primer sets per enterovirus species that generated two overlapping fragments decreasing labor, time, and cost per sample tested. The first generic primer set (C004-F and CRE-R) corresponds to two conserved regions for all enteroviruses (-A through -D) and was capable of successfully amplifying the first half of the genome (~4500 nt). The second half of the genome was amplified using generic primers that were designed to specifically target all viruses within the respective species (EV-A, -B, -C, and -D). We were able to amplify nearly the entire genome for all enteroviruses, using only five PCR mixtures. The sensitivity of the primers was determined and the EV-A primers appeared to be more sensitive than the others (Table 3). However, the primers selected for this project were sensitive enough to detect all enteroviruses and specific enough to detect only one species type in the presence of the other species.

To validate our sequencing method, we performed it using twenty RD isolate samples from human stools and sewage concentrates. The consensus lengths after de novo assembly were

| Primer set | Enterovirus species | Primer name | 5′–3′ Sequence | Genome position1 |
|------------|--------------------|-------------|----------------|-----------------|
| 1          | A, B, C, and D     | C0042       | TTAAACAGCYYKDGGGTTG | 1–20           |
|            | A, B, C, and D     | EV-CRE-R    | CGBRRTTTGGTCGCTAGTGC | ~4500         |
| 2          | A                  | EVA-4110-F  | AARAAITTYAAYGATGGGC | 4110–4130      |
|            | A                  | EVA-7410-R  | TTTGCTATTCGTTAATAC | 7410–7390      |
|            | B                  | EVB-4110-F  | GGCNTGGCTAYACRRAARG | 4110–4130      |
|            | B                  | EVB-7400-R  | GACCGAGTCCGAGAATTTACC | 7400–7378   |
|            | C                  | EVC-4220-F  | GARCNTGYYAGCNGCNAA | 4220–4242      |
|            | C                  | C0052       | CCAATYAAAAAATTTACCC | 7347–7415     |
|            | D                  | EVD-4112-F  | GCCCTKMCCAAAAGATWGAC | 4112–4132     |
|            | D                  | EVD-7362-R  | CCAAKTRACCCAAATTTACC | 7362–7342     |

1The specific genome position. Position may vary depending on the strain of the virus. 2Primer described in Bessaud et al. (2016).
### TABLE 3 | Sensitivity of the NGS assay.

| Species (reference strain)†  | Dilution factor | Viral titer | rRT-PCR cycle threshold² | DNA concentration (ng/µL)³ | Number of reads⁴ | Number of reads mapping against the reference (%)⁵ | Contig length⁶ |
|-----------------------------|-----------------|-------------|--------------------------|-----------------------------|------------------|--------------------------------------------------|--------------|
| EV-A (EV-A71 MAD-72341-04)  |                 |             |                          |                             |                  |                                                  |              |
| 4⁰                          | 6.3 10e6        | 18.4        | 50.6                     | 437 922                     | 434360           | (99.2%)                                          | 7 336        |
| 4¹                          | 1.6 10e6        | 21.5        | 31.8                     | 483 490                     | 471138           | (97.5%)                                          | 7 410        |
| 4²                          | 3.9 10e5        | 23.1        | 20.1                     | 640 522                     | 565731           | (88.3%)                                          | 7 470        |
| 4³                          | 9.8 10e4        | 30.2        | 11.7                     | 478 940                     | 357168           | (74.6%)                                          | 7 405        |
| 4⁴                          | 2.5 10e4        | 31.1        | 10.2                     | 270 790                     | 109731           | (40.5%)                                          | 7 405        |
| 4⁵                          | 6.2 10e3        | 29.6        | 9.5                      | 500 780                     | 85344            | (17.0%)                                          | 7 405        |
| 4⁶                          | 1.5 10e3        | 32.6        | 10.6                     | 349 396                     | 24072            | (6.9%)                                           | 7 408        |
| EV-B (CV-B4 E2)            |                 |             |                          |                             |                  |                                                  |              |
| 4⁰                          | 4.0 10e7        | 21.3        | 58.1                     | 526 474                     | 457529           | (86.9%)                                          | 7 442        |
| 4¹                          | 1.0 10e7        | 25.3        | 36.5                     | 529 664                     | 475725           | (89.8%)                                          | 7 430        |
| 4²                          | 2.5 10e6        | 28.3        | 22.7                     | 657 448                     | 539309           | (82.0%)                                          | 7 450        |
| 4³                          | 6.3 10e5        | 29.3        | 15.7                     | 365 820                     | 222881           | (60.9%)                                          | 7 410        |
| 4⁴                          | 1.6 10e5        | 33.5        | 15.1                     | 400 011                     | 127689           | (31.9%)                                          | 7 434        |
| 4⁵                          | 3.9 10e4        | Undeter⁷    | 17.6                     | 517 449                     | 9656             | (2.0%)                                           | 4 294        |
| 4⁶                          | 9.8 10e3        | Undeter     | 10.4                     | 338 670                     | 2375             | (0.7%)                                           | 6 872        |
| EV-C (CV-A13 MAD 67001)    |                 |             |                          |                             |                  |                                                  |              |
| 4⁰                          | 1.0 10e8        | 18.6        | 29.3                     | 538 236                     | 50299            | (93.5%)                                          | 7 450        |
| 4¹                          | 2.5 10e7        | 22.3        | 12.3                     | 551 962                     | 462287           | (83.8%)                                          | 7 374        |
| 4²                          | 6.3 10e6        | 23.7        | 10.4                     | 369 904                     | 255970           | (69.2%)                                          | 7 381        |
| 4³                          | 1.6 10e6        | 26.5        | 9.4                      | 418 630                     | 60066            | (14.4%)                                          | 7 442        |
| 4⁴                          | 3.9 10e5        | 28.7        | 10.8                     | 383 840                     | 15414            | (4.0%)                                           | 3 279        |
| 4⁵                          | 9.8 10e4        | 31.2        | 10.8                     | 454 272                     | 10492            | (2.5%)                                           | 3 371        |
| 4⁶                          | 2.4 10e4        | 33.6        | 8.8                      | 467 808                     | 982              | (0.2%)                                           | 3 262        |
| EV-D (EV-68 Fermon)        |                 |             |                          |                             |                  |                                                  |              |
| 4⁰                          | 1.3 10e6        | 24.3        | 23.8                     | 468 514                     | 465107           | (99.3%)                                          | 7 367        |
| 4¹                          | 3.3 10e5        | 28.2        | 14.1                     | 585 648                     | 494846           | (84.5%)                                          | 7 359        |
| 4²                          | 8.1 10e4        | 32.0        | 14.2                     | 496 292                     | 408854           | (82.4%)                                          | 7 282        |
| 4³                          | 2.0 10e4        | 35.6        | 8.3                      | 338 906                     | 178190           | (52.6%)                                          | 7 277        |

(Continued)
obtained for the ten isolate samples from stool tested, indicating that the method was sensitive enough to determine the entire genome. The results were different for sewage because mixtures of strains appeared to be more complex (up to 7 different strains) than those present in stools (two strains maximum). Only three of the ten isolate samples resulted in sequencing data for the whole genome. However, the sequencing data for the other seven provided long genomic contigs and the coverage necessary to identify the type of virus(es) present using VP1 de novo assembly generally span all the capsid-encoding region.

In this case, de novo assembly can fail to build full-length contigs because it is impossible for the software to determine which reads are from virus 1 or from virus 2. This limitation is not related to our method used to generate DNA from RNA genomes, but it is due to the fact that most NGS methods (i.e., Illumina) cannot deal with long DNA fragments, requiring the shearing of the DNA amplicons prior to sequencing. Other NGS methods (such as PacBio or MinION) can sequence long DNA fragments and could be used to overcome the assembly problems but these methods have a high error rate compared to Illumina. Therefore, there is a balance between the accuracy of the sequences and the length of the contigs that can be generated in cases of mixed recombinant genomes. Nonetheless, this challenge did not compromise the main objective of our method, which was to allow the identification of the enterovirus lineages found in a given sample. The typing of EVs relies on their capsid sequence and intertypic recombination is very uncommon inside this genomic region, the contigs generated through de novo assembly generally span all the capsid-encoding region.

TABLE 4 | NGS analysis in three mixtures containing four viruses.

| Mixture | Contig name | Contig length | Annotation (virus) | Percent identity | VP1 | VP1 length |
|---------|-------------|---------------|-------------------|-----------------|-----|-----------|
| 1       | contig 3    | 6620          | Coxsackievirus A10, strain Kowalk | 99.9            | full | 894       |
|         | contig 1    | 7435          | Echovirus 11, isolate ROU-9191 | 99.9            | full | 876       |
|         | contig 8    | 4681          | Coxsackievirus A13, isolate 67001 | 99.8            | partial | 343 |
|         | contig 2    | 7318          | Enterovirus 68, strain Fermon | 99.9            | partial | 536 |
| 2       | contig 1    | 4444          | Coxsackievirus B6, strain Schmitt | 99.7            | full | 846       |
|         | contig 2    | 4416          | Echovirus 5, strain Noyce | 99.9            | full | 876       |
|         | contig 4    | 4408          | Echovirus 7, strain Wallace | 99.9            | full | 876       |
|         | contig 3    | 4364          | Coxsackievirus B3, strain Nancy | 99.9            | full | 852       |
| 3       | contig 1    | 7416          | Echovirus 5, strain Noyce | 99.9            | full | 876       |
|         | contig 2    | 7404          | Echovirus 7, strain Wallace | 99.9            | full | 876       |
|         | contig 3    | 7383          | Coxsackievirus B6, strain Schmitt | 99.9            | full | 846       |
|         | contig 4    | 7376          | Coxsackievirus B3, strain Nancy | 99.9            | full | 852       |

1Mixture 1 was performed using four viruses (one strain from each species) with equal quantities of RNA. Mixture 2 and 3 were performed using four EV-B viruses. For mixture 2, PCR was performed using primers designed to amplify the first half of the genome, whereas mixture 3 was the PCR for the full genome. 2The identification of the contig that is provided in the NGS analysis. 3The length of the contig in nucleotides (∼7500 nt = enterovirus size). 4The name given for the virus in GeneBank (using BLAST in NCBI). 5The percent identity between the contig sequence and the nearest sequence of known virus in GeneBank. 6If the VP1 was retrieved in full or partial size.
### TABLE 5 | Detection of viral sequences in supernatants of RD cells infected with stool and sewage samples, using de novo assembly.

| Sample  | Enterovirus contig(s) | VP1 contig(s) | Contig(s) with VP1(nt) | Read Count | Viral type(s) | Enterovirus species |
|---------|-----------------------|---------------|------------------------|------------|---------------|-------------------|
| Stool 1 | 5                     | 1             | 6886                   | 450016     | E-21          | EV-B              |
| Stool 2 | 1                     | 1             | 7475                   | 557642     | E-4           | EV-B              |
| Stool 3 | 3                     | 2             | 7230                   | 452063     | E-11          | EV-B              |
|         |                       |               | 2069                   | 842        | EV-C99        | EV-C              |
| Stool 4 | 2                     | 2             | 6948                   | 371680     | E-2           | EV-B              |
|         |                       |               | 6932                   | 72007      | CV-A4         | EV-A              |
| Stool 5 | 3                     | 1             | 7247                   | 437921     | E-14          | EV-B              |
| Stool 6 | 2                     | 2             | 7426                   | 363551     | E-5           | EV-B              |
|         |                       |               | 5170                   | 31239      | EV-C99        | EV-C              |
| Stool 7 | 5                     | 2             | 7411                   | 523337     | E-13          | EV-B              |
|         |                       |               | 6606                   | 5567       | E-20          | EV-B              |
| Stool 8 | 4                     | 2             | 7109                   | 605697     | EV-B84        | EV-B              |
|         |                       |               | 1791                   | 75         | CV-A4         | EV-A              |
| Stool 9 | 11                    | 2             | 7581                   | 413225     | E-15          | EV-B              |
|         |                       |               | 1941                   | 359        | CV-A13p       | EV-C              |
| Stool 10| 4                     | 2             | 7109                   | 339135     | E-14          | EV-B              |
|         |                       |               | 5274                   | 41917      | EV-C99        | EV-C              |
| SEW 1   | 3                     | 2             | 7337                   | 435585     | E-6           | EV-B              |
|         |                       |               | 5095                   | 1657       | E-13          | EV-B              |
| SEW 2   | 2                     | 2             | 4444                   | 297319     | E-19          | EV-B              |
|         |                       |               | 4404                   | 8236       | E-7           | EV-B              |
| SEW 3   | 9                     | 5             | 7403                   | 102527     | E-20          | EV-B              |
|         |                       |               | 7380                   | 141133     | E-12          | EV-B              |
|         |                       |               | 7371                   | 290910     | E-7           | EV-B              |
|         |                       |               | 4094                   | 11083      | E-6           | EV-B              |
|         |                       |               | 3758                   | 390        | EV-A76        | EV-A              |
| SEW 4   | 5                     | 3             | 4574                   | 63109      | E-7           | EV-B              |
|         |                       |               | 3871                   | 13766      | CV-B5         | EV-B              |
|         |                       |               | 7369                   | 390728     | E-33          | EV-B              |
| SEW 5   | 9                     | 3             | 4476                   | 333848     | E-11          | EV-B              |
|         |                       |               | 3302                   | 107812     | E-19          | EV-B              |
|         |                       |               | 2607                   | 4608       | E-12          | EV-B              |
| SEW 6   | 8                     | 3             | 4400                   | 81721      | E-6           | EV-B              |
|         |                       |               | 4367                   | 359419     | CV-B5         | EV-B              |
|         |                       |               | 207                    | 32         | E-6p          | EV-B              |
| SEW 7   | 20                    | 2              | 5980                   | 409716     | E-6           | EV-B              |
|         |                       |               | 2663                   | 1321       | E-12          | EV-B              |
| SEW 8   | 3                     | 2             | 6949                   | 303796     | E-12          | EV-B              |
|         |                       |               | 6858                   | 17635      | E-24          | EV-B              |
| SEW 9   | 2                     | 2             | 4410                   | 56755      | E-11          | EV-B              |
|         |                       |               | 5174                   | 144696     | E-6           | EV-B              |
| SEW 10  | 28                    | 7              | 3838                   | 4963       | E-12          | EV-B              |
|         |                       |               | 6155                   | 13411      | E-6           | EV-B              |
|         |                       |               | 6176                   | 172800     | E-6           | EV-B              |
|         |                       |               | 4228                   | 182570     | E-11          | EV-B              |
|         |                       |               | 683                    | 100        | E-6p          | EV-B              |
|         |                       |               | 1266                   | 227        | E-6p          | EV-B              |
|         |                       |               | 3380                   | 1060       | E-33          | EV-B              |

1 Type of isolates: RD cells infected by stool extracts or sewage (SEW) concentrates. 2 Number of contigs identified per sample. 3 Number of contigs that contained the VP1 region (needed to identify the virus). 4 Size of the contig with the VP1 region included. Size varies but we are interested in the ones closest to the size of the whole genome (~7500 nb). 5 Number of reads used to generate the VP1 contigs. 6 Virus serotype (identity) using the VP1 region. Note: p indicates partial size of the VP1 based on the length. In SEW 6 and ~10 different E-6 genotypes were found. 7 Enterovirus species classification based on the virus identified.
Blomqvist, S., and Roivainen, M. (2016). Isolation and characterization of enteroviruses within the four species A to D that were a way to collect full genome sequencing data from mixtures within given populations. In addition, this method provided as indicated in our results. useful when analyzing human and environmental field samples during a single sequencing run. This type of assay would be the specific identification of all enteroviruses present in samples.

Our strategy is more suitable to specifically amplify the relevant sequences of viruses that are unexpected or unknown, they have the disadvantage of decreasing the number of relevant reads since the assay includes species-specific primers. Although strategies based on random primers, we can increase the amplification of the respective targeted species. This is not the case with random primers that cannot be ensured to capture all enteroviruses within a given species. In addition, the assay can be simplified to target one particular species. Although strategies based on random amplification have the advantage of being able to amplify copies of viruses that are unexpected or unknown, they have the disadvantage of decreasing the number of relevant reads since the sequence of non-viral origins is also amplified and sequenced. Our strategy is more suitable to specifically amplify the relevant sequences, limiting the requested depth of coverage, which favors multiplexing and thus reducing the cost per genome and per sample.

In conclusion, the method described in this study enables the specific identification of all enteroviruses present in samples during a single sequencing run. This type of assay would be useful when analyzing human and environmental field samples as indicated in our results.

This contributes to the study of EV diversity and ecosystem within given populations. In addition, this method provided a way to collect full genome sequencing data from mixtures of enteroviruses within the four species A to D that were present in cellular supernatants. These data become a necessity for surveillance purposes and when studying the relationships between the genetic characteristics, including the mosaic features of their genomes acquired through frequent intra- and intertypic recombination and their biological properties. Indeed, mutations and recombination events can be implicated in reintroducing virulence factors and have been involved in the evolution of enteroviruses (Riquet et al., 2008; Jegouic et al., 2009; Joffret et al., 2012; Holmblat et al., 2014).

AUTHOR CONTRIBUTIONS
M-LJ, PP, and FD conceived and designed the experiments and analyzed the data. M-LJ and PP performed the experiments. RR and J-MH contributed to samples and experiments in Madagascar. PP and M-LJ wrote the manuscript. MB, RR, J-MH, and FD critically revised.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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