Rapid Confirmation of the Zaire Ebola Virus in the Outbreak of the Equateur Province in the Democratic Republic of Congo: Implications for Public Health Interventions

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Ten days after the declaration of the Ebola outbreak in the Democratic Republic of Congo, rapid identification of the species Zaire Ebola virus using partial gene amplification and nanopore sequencing backed up the use of the recombinant vesicular stomatitis virus–Zaire Ebola virus vaccine in the recommended ring vaccination strategy.

Keywords. Ebola; Democratic Republic of Congo; nanopore sequencing.

The Democratic Republic of Congo (DRC) declared a new outbreak of Ebola virus disease (EVD) on 8 May 2018. Initially, a report from the Provincial Health Division of the Equateur Province at the end of April informed the national authorities of 21 cases of potential hemorrhagic fever, including 17 deaths in the Ikoko-Impenge health area. Deployed disease experts on site identified 5 possible cases on 5 May and a blood sample was taken from each individual, 2 who were hospitalized in Bikoro and 3 at the health centre of Iboko. Samples were sent to Institut National de Recherche Biomédicale (INRB) in Kinshasa, and the presence of Ebola virus was identified in 2 samples on 7 May [1]. The samples tested positive with the OraQuick Ebola Rapid Antigen test (OraSure Technologies) and with the RealStar Zaire Ebolavirus reverse-transcription polymerase chain reaction (PCR) kit 1.0 (Altona Diagnostics).

Proven emergency responses such as enhanced surveillance, active case finding, contact tracing, infection prevention, safe burials, and diagnosis with deployment of mobile laboratories on site were implemented rapidly. However, in 2017, the World Health Organization (WHO) recommended the deployment of the recombinant vesicular stomatitis virus–Zaire Ebola virus (rVSV-ZEBOV) candidate vaccine as an additional measure for prevention and control in the event of novel Zaire Ebola virus (EBOV) outbreaks [2]. This vaccine has been shown to be effective and safe and has been used during the epidemic decline of EBOV in Guinea in 2015 and during the flare-up of the disease in rural Guinea in 2016 [3, 4]. However, 4 species of Ebola viruses have been documented in Africa: Zaire (EBOV), Sudan (SUDV), Bundibugyo (BDBV), and Tai Forest [5]. Whereas 7 of the 8 previous EBOV epidemics in DRC were due to EBOV, BDBV was responsible for the outbreak in Isiro in 2012, and previous outbreaks in South Sudan were all close to the border with DRC and were due to SUDV [6]. Therefore, it is thus important to confirm rapidly the viral strain of any new outbreak to guarantee the utility of anti-EBOV vaccination. Here, we describe the rapid laboratorial response that led to the confirmation of EBOV in the latest outbreak and highlight the implication of rapid viral detection for public health interventions.

MATERIALS AND METHODS

PCR Amplification

Viral RNA was extracted from 140 μL of whole blood collected from the 5 clinically suspected Ebola virus–infected individuals from Bikoro/Iboko using the Nuclisens kit (bioMérieux) and following the manufacturer’s instructions. After preparation of complementary DNA with GoScript Reverse Transcriptase (Promega, France), amplification of a small fragment of the VP35 region was attempted in a seminested PCR with a modified protocol adapted from He and colleagues [7]. Three additional samples were included as controls: 2 EBOV positive and 1 negative. In the first round, a 217-bp fragment was targeted with primers VP35-F: 5'-ATGTGGATCRAAG-3' and VP35-R: AGCGR ATGTGGAATSACRGGT-3’. In the second round, an 184-bp product was amplified with primers VP35-R and VP35-in-F: 5'-GCTTTYCAAYCAAYTAGTCAAG-3'.

Sequence Analysis

First-round VP35 PCR products from positive samples were barcoded and pooled using the Native Barcoding Kit (Oxford
Nanopore Technologies). Sequencing libraries were generated from the barcoded products using the Genomic DNA Sequencing Kit EXP-NBD103/SQK-LSK208 (Oxford Nanopore Technologies) and were loaded onto a R9 flow cell. Genetic data was collected for 1 hour.

Bioinformatics and Phylogenetic Analysis

Basecalling of sequenced data was done with ONT Albacore Sequencing Pipeline software version 2.1.10. Adapter removal and demultiplexing were achieved with Porechop 0.2.3.

Filovirus genus references were retrieved from the Hemorrhagic Fever Viruses (HFV) database at the Los Alamos National Laboratory. Multireference prealignment analysis of quality and error profiles using reads with basecall quality ≥11 was done with the software package NanoOK 1.10. Ebola virus reads were corrected with Canu 1.6, and consensus sequences were produced by alignment of reads to the corresponding Ebola virus reference genome following signal-level analysis using Nanopolish 0.9.

One sequence per outbreak was retrieved from the HFV database and used for phylogenetic inference together with the nanopore consensus sequences. We also included a previously obtained VP35 sequence from an unpublished strain from the outbreak in Likati in 2017. Phylogenetic analyses were done using maximum likelihood methods with approximate likelihood ratio test nonparametric branch support based on a Shimodaira-Hasegawa–like procedure using PhyML version 3.1 software. The HKY85 model plus a discrete gamma model were used as nucleotide substitution models.

RESULTS

The 2 positive control samples from Boende 2014 were successfully amplified and no PCR product was obtained for the negative sample. The 2 samples from Bikoro/Iboko 2018 that tested initially positive with the commercial tests were also positive in the VP35 PCR, 2 other samples had a weak PCR product, and the remaining sample resulted in no amplification. We attempted sequencing on the 6 samples with PCR amplification. After 1 hour, between 400 and >4000 reads were obtained depending on the sample. NanoOK report indicated that >95% of the reads from the 2 known positive samples from Bikoro/Iboko and the 2 positive control samples from Boende mapped to the EBOV species. The remaining reads and the reads from the samples with weak positive PCR product did not align to any filovirus genera and were queried in BLAST (Basic Local Alignment Search Tool), and the best hits pointed to human DNA. Consensus sequences were produced for the new EBOV strains and aligned with reference sequences from previous Ebola outbreaks including the 5 Ebola virus species and other filoviruses such as Marburg. The phylogenetic tree shown in Figure 1 underscores that this highly conserved region is discriminatory between Ebola virus species and illustrates that the new strains from the ninth Ebola outbreak in DRC belong to the Ebola Zaire species (EBOV).

The laboratory results of the 5 suspected cases and the clinical signs and history for each patient are summarized in Supplementary Table 1. Overall, all 5 patients had diarrhea, severe fatigue, and abdominal pain. Additional symptoms were muscle or joint pain and anorexia for 4 of them. Three patients had also gum and/or nose bleeding among symptoms, but only 1 of them was confirmed as infected with Ebola virus by sequence and PCR analysis.

DISCUSSION

After only 10 days of the ninth Ebola outbreak declaration, the virus strain was confirmed as belonging to the Zaire species (EBOV). As in the outbreak in West Africa, nanopore sequencing using the MinION device proved useful for quick pathogen confirmation and strain identification [8, 9]. The device provided speed (library preparation, basecalling of preliminary read data and multireference prealignment took <6 hours) and ease of use besides portability advantages over other sequencing machines. The rapid viral characterization of the strain backed up the rVSV-ZEBOV candidate vaccine for the ring vaccination strategy recommended by WHO [2]. Today >1000 people have been vaccinated, mainly health professionals and people exposed to patients with confirmed EBOV and their contacts [1]. Nevertheless, further efforts to increase adequate hygiene practices and safe burials must also be continued.

Previous outbreaks in DRC were in very remote locations or small towns, but this new outbreak affected simultaneously at least 3 different locations in the Equateur Province. It included 2 remote sites, Bikoro and Iboko/Itipo health areas, and cases in Mbandaka, a city of >1 million inhabitants and with transport and port links to the rest of the country and neighboring countries. The current outbreak is thus complex with the potential for urban and larger geographic spread. It was thus urgent to start rapidly the ring vaccination to stop the spread of the virus.

DRC had already experienced 8 other Ebola outbreaks since 1976 [6]. The country has thus implemented a strategy at the national level to rapidly identify a potential new outbreak. This includes rapid reporting of clinical suspected cases and shipment of samples to the reference laboratory at INRB in Kinshasa for confirmation. In the current situation, the health service from Equateur Province gave the alert in late April. Nevertheless, the clinical presentations of the 5 suspected cases illustrate how difficult it is to distinguish Ebola from some other diseases without laboratory confirmation. A similar alert was issued in February in the same area with 15 cases, among which 8 individuals died, but no laboratory confirmation occurred (unpublished data). This observation underscores that the outbreak may have started earlier and emphasizes the potential of Ebola outbreaks to go unrecognized in remote areas. However, any relation between the 2 alerts remains to be examined.
Each EVD outbreak is the result of a cross-species transmission event from a reservoir or an intermediate/amplifying animal host species [6]. This new outbreak, together with the large outbreak in West Africa, illustrates the major consequences of recurrent zoonotic spills. Studies on the ecology and animal reservoir of Ebola viruses are thus urgently needed so exposure to infection can be anticipated or avoided. Finally, the advantages provided by novel diagnostic technologies such as nanopore sequencing using the MinION device highlight the importance of local capacity-building programs for rapid pathogen confirmation and corresponding public health interventions.

Supplementary Data
Supplementary materials are available at Clinical Infectious Diseases online. Consisting of data provided by the authors to benefit the reader, the posted materials are not copyedited and are the sole responsibility of the authors, so questions or comments should be addressed to the corresponding author.

Notes
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