Detection of the Northeastern African Rift Valley Fever Virus Lineage During the 2015 Outbreak in Mauritania

Ndée Sakho Bob,1 Hampté Bâ,2 Gamou Fall,1 Elkhali Ishagh,3 Mamadou Y. Diallo,4 Abdourahmane Sow,5 Papa Mbacké Sembene,6 Ousmane Faye,1 Brahiam El Kouri,2 Mohamed Lemine Sidi,7 and Amadou Alpha Sall1

1Pole of Virology, Arbovirus and Viral Hemorrhagic Fevers Unit, Pasteur Institute of Dakar, Senegal; 2Viral Hemorrhagic Fevers Diagnostics Unit, National Institute of Public Health Research, Nouakchott, Mauritania; 3Department of Epidemiological Surveillance, Ministry of Health, Islamic Republic of Mauritania, Nouakchott; 4Health Securities and Emergency, World Health Organization, Mauritania; 5West African Health Organization, Ouagadougou, Burkina Faso; 6Cheikh Anta Diop University of Dakar, Senegal; and 7Direction of Diseases Control, Ministry of Health, Islamic Republic of Mauritania, Nouakchott.

Background. Rift Valley fever (RVF) is an acute viral anthropozoonosis that causes epizootics and epidemics among livestock population and humans. Multiple emergences and reemergences of the virus have occurred in Mauritania over the last decade. This article describes the outbreak that occurred in 2015 in Mauritania and reports the results of serological and molecular investigations of blood samples collected from suspected RVF patients.

Methods. An RVF outbreak was reported from 14 September to 26 November 2015 in Mauritania. Overall, 184 suspected cases from different localities were identified by 26 health facilities. Blood samples were collected and tested by enzyme-linked immunoabsorbent assay (ELISA) and real-time reverse-transcription polymerase chain reaction (RT-PCR) at the Institut Pasteur de Dakar (IPD). Sequencing of partial genomes and phylogenetic analyses were performed on RT-PCR–positive samples. As part of routine surveillance at IPD, samples were also screened for dengue, yellow fever, West Nile, Crimean Congo hemorrhagic fever, Zika, and Chikungunya viruses by ELISA and RT-PCR.

Results. Of the 184 suspected cases, there were 57 confirmed cases and 12 deaths. Phylogenetic analysis of the sequences indicated an emergence of a virus that originated from Northeastern Africa. Our results show co-circulation of other arboviruses in Mauritania—dengue, Crimean Congo hemorrhagic fever, and West Nile viruses.

Conclusion. The Northeastern Africa lineage of RVF was responsible for the outbreak in Mauritania in 2015. Co-circulation of multiple arboviruses was detected. This calls for systematic differential diagnosis and highlights the need to strengthen arbovirus surveillance in Africa.

Keywords. Mauritania; Northeastern African lineage; phylogeny; Rift Valley fever; 2015.

Rift Valley fever virus (RVFV) is a mosquito-borne virus belonging to the Phlebovirus genus of the Bunyaviridae family. Rift Valley fever virus is a segmented virus with 3 single-stranded RNA genomes: L (large), M (medium), and S (small). The virus is primarily transmitted to animals and humans through infected mosquito bites. Other modes of transmission include exposure to body fluids, blood, and tissues of infected animals or contact with aerosols [1, 2]. In animals, Rift Valley fever (RVF) is characterized by abortions among pregnant females and high mortality rates for offspring [3]. In human, symptoms are generally mild but may evolve to severe disease, such as hemorrhagic, meningoencephalitis, and retinopathy with fatal outcomes [4]. Cases of miscarriage in pregnant women associated with RVFV infection have been shown recently in Sudan [5]. The disease causes a huge economic burden because of livestock-trade bans and restrictions, especially in the Arabian Peninsula.

Rift Valley fever virus is classified as a category A pathogen by the US Center for Diseases Control and Prevention and the Department of Agriculture. There is no commercially available vaccine for humans or animals, and effective antiviral drugs have not been identified. There is therefore an urgent need to develop countermeasures against propagation and introduction of RVFV into Europe and nonendemic countries. Rift Valley fever is one of the priority diseases identified by the World Health Organization (WHO) as a likely cause of a future outbreak [6]. Indeed, a new plan was developed after the Ebola epidemic for urgent research and development toward new diagnostic tools, vaccines, and medicines for RVFV.

In West Africa, the first major RVF outbreak occurred on October 1987 at the end of the rainy season in Mauritania. It was particularly notable in the Trarza region along the Senegal River after the impoundment of the Diama and flooding around the lowlands [7]. During that outbreak, 220 human deaths were recorded [8–10], and high abortion rates among small
ruminants were observed [10, 11]. Subsequently, the country experienced several outbreaks, some of which had severe hemorrhagic manifestations [12–15]. In 1998, the Hodh El Gharbi region located in the southeast of Mauritania recorded >300 human RVF cases, including 6 deaths [10]. In 2003, several other regions of the south, southeast, and center experienced RVF outbreaks. Twenty-five confirmed human cases, including 16 with hemorrhagic signs and 4 deaths, were reported in the south (Trarza, Brakna, Gorgol), southeast (Assaba), and center (Tagant) [13]. Only the northern part was unaffected, until October 2010, when an RVF outbreak was identified in Adrar and Inchiri regions, with 70 human cases, including 13 deaths [14]. Two years later, another more extended RVF outbreak occurred in Assaba, Brakna, Hodh El Chargui, Hodh El Gharbi, Tagant, and Trarza regions and Nouakchott capital city of the country between September and October 2012. During this outbreak, 34 human cases, including 17 deaths, were reported [15]. The viruses isolated in 1989 formed a discrete West African lineage, whereas strains recovered in 2003 fall within the East/Central African lineage [13]. Indeed, the 2003 Mauritanian viruses were clustered into lineage C, together with strains isolated from Kenya in 2007, South Africa in 2008 and 2009, and Madagascar in 2008. Lineage C also includes strains isolated in Zimbabwe in 1976–1979 and 1998, suggesting that viruses originating from Zimbabwe might be responsible for outbreaks in Mauritania [16]. Phylogenetic analyses from the 2010 and 2012 outbreaks suggest a reemergence from local RVFV that clustered in the West African lineage [14, 15].

In the context of Ebola virus disease outbreak in West Africa, which was declared by WHO on August 8, 2014, to be a global public health emergency [17], the Mauritanian epidemiological surveillance office was alerted on September 14, 2015, of a suspected case of viral hemorrhagic fever. A sample was sent to the Institut Pasteur de Dakar (IPD) for Ebola diagnosis and arboviruses differential diagnosis in case of a negative Ebola result. Following this case, 10 more patients were hospitalized for hemorrhagic fever (Ebola was ruled out) in Mauritania and also tested for RVF, dengue, yellow fever, West Nile, CCHF, and Chikungunya viruses by RT-PCR and ELISA.

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After the RVF outbreak declaration, investigations were performed, and the following case definition was used: patient consulting with an axillary temperature >37.5°C lasting for 48 hours, associated with at least 1 of the following signs: exhaustion, back pain, myalgia, headache, nausea/vomiting, diarrhea and/or cutaneous bleeding, bleeding to bite sites, epistaxis, gingival bleeding, or other bleeding. Blood samples were collected from a total of 173 patients.

All of the blood samples collected before the outbreak declaration (n = 10) and during outbreak investigation (n = 173) were sent to the Institut national de la recherche en santé publique (INRSP) in Nouakchott and tested by ELISA for detection of immunoglobulin M (IgM) against RVF, yellow fever, and CCHF viruses after sample inactivation by heating at 56°C for 30 minutes. Following INRSP diagnostics, samples were sent to IPD for confirmation of serological tests, RNA detection by RT-PCR, and differential diagnosis. ELISA for IgM of RVFV and other arboviruses was performed in the WHO collaborating center for arboviruses and hemorrhagic fever viruses in IPD using in-house methods with antigens and immune ascites produced in mice.

Regarding the RVFV RT-PCR, the primers (forward TGCCACGAGTGYAGGCCA, reverse TTGAACAGTGGGT CCGAGA) and probe 6FAM-TCCCTCTCCCCAGTCAGCCCCCACA C-BHQ1 were used [19]. For other arboviruses, RT-PCR for Chikungunya [20], dengue [21], West Nile [22], yellow fever [23], Zika [24], and CCHF [25] viruses was performed as previously described. The RNA was amplified using ABI Prism 7000 SDS Real-Time apparatus (Applied Biosystems) with the QuantiTect kit (Qiagen).

Samples that were RT-PCR–positive for RVFV were used for sequencing of partial small (S), medium (M), and large (L) segments using specific primers (Supplementary Table 1). Sequences were analyzed using online tools revseq and merger emboss (http://www.bioinformatics.nl/cgi-bin/emboss/merger; http://emboss.bioinformatics.nl/cgi-bin/emboss/revseq), and phylogenetic studies were conducted using maximum likelihood method and MEGA software (MEGA 6.06-mac) [26].
RESULTS

From September 14 to November 26, 2015, the IPD laboratory received in total 184 blood samples collected in 26 health facilities from patients with suspected hemorrhagic fever in Mauritania (Table 1). The mean age of patients was 25 years (range = 0.6–90), and the sex ratio (male/female) was 3.38. Among the 184 samples, 57 were positive for RVF by either RT-PCR and/or IgM (16 RT-PCR+/IgM+, 40 RT-PCR+/IgM−, and 1 RT-PCR−/IgM−). One sample was found coinfected with both RVF and CCHF viruses by RT-PCR, and 5 samples were positive only for CCHF IgM by ELISA. Twenty-seven samples were positive for dengue (8 RT-PCR+/IgM+, 19 RT-PCR−/IgM−), with 8 samples positive for both RVF and dengue by RT-PCR. One sample was positive for West Nile IgM by ELISA and RVFV by RT-PCR (Table 1).

The overall infection rate by RVFV was 30.97% (n = 57/184), of which 78.94% were males (n = 45/57) and 21.05% were females (n = 12/57). The age groups 0–20 years and 21–40 years were the most affected with 38.59% (n = 22/57) and 45.61% (n = 26/57) of confirmed cases, respectively. With respect to age, the fatality rate was 33.3% (n = 1/3) in patients aged >61 years and 13.6% (n = 3/22) in patients aged 0–20 years (Table 2).

Among the 5 health facilities that reported at least 1 death, lethality varied between 15% for the National Hospital Center (n = 3/20) and 100% for Health Center Moudjeria (n = 1/1). Kiffa’s hospital center recorded the highest number of deaths with a lethality rate of 25% (n = 4/16). Of the 26 health facilities that reported at least 1 suspected case, 30.76% (n = 8/26) were in the southeast, 23.07% (n = 6/26) were in the south, and 30.76% (8/26) were in Nouakchott (Figure 1). The health facilities in the center and north each represented 7.69% (n = 2/26) of cases. Thus, 50.87% (n = 29/57) of the confirmed RVF patients were from the south and southeast parts of the country, and 47.36% (n = 27/57) originated from Nouakchott capital city, where 24% of the national population lives.

The operating notifications records showed that the first confirmed case was hospitalized on September 14, 2015, at Health Center Maghta Lahjar, and the date of disease onset was September 9, 2015. The last positive case was recorded at Hospital Center Aïoun on November 3 and confirmed by November 13, 2015. The outbreak began at epidemiological week 38 and stopped at week 46, with 2 peaks at weeks 41 and 45. The analysis showed 2 epidemic periods; the first covered weeks 38–43, and all deaths occurred during this period, with the lethality being highest at week 40 (Figure 2).

Phylogenetic analysis based on RVFV partial small (709 bp), medium (718 bp), and large RNA (4764 bp) sequences showed that sequences from different areas clustered together with a
low level of variation between isolates. The strains were different from the other RVF strains isolated in Mauritania in 2010 and 2012 and more closely related to those from Senegal in 2013 [27] and clustered in the Egyptian lineage (Figure 3).

**DISCUSSION**

As in 2010 [10] and 2012 [11], Mauritania experienced a new RVF outbreak in 2015, which spread over 8 weeks, with 184 suspected cases, including 57 confirmed cases and 12 deaths. Similar to 2010 and 2012 outbreaks, RVF positive patients were predominantly males (78.95%) and younger individuals (aged <40 years); the fatality rate was greater in male patients. This can be explained by the high exposure risk of men to potentially infected mosquitoes during agricultural work or direct contact with viremic livestock, infected tissues, and aborted animals [15]. The outbreak affected the same areas as the 2012 RVF outbreak (Assaba, Brakna, Hodh El Chargui, Hodh El Gharbi, Tagant, Trarza, and Nouakchott), contrasting with the 2010 outbreak, which was confined to northern regions (Inchiri and Adrar).

Differential diagnosis in IPD revealed co-circulation of RVF and CCHF with 1 coinfection. This was similar to the 2012 outbreak where 1 coinfection with CCHF was also found in Moudjeria [15]. Co-circulation of RVFV with dengue virus was also identified in 8 samples during the 2015 outbreak. In addition to dengue and CCHF, the circulation of West Nile virus was identified for the first time in Mauritania. Taken together, these results indicate circulation of different arboviruses in the region during the RVF outbreak and call for systematic differential diagnosis or a syndromic approach for febrile and hemorrhagic fever surveillance to prevent a large-scale outbreak.

Phylogenetic analysis revealed that the same strain was circulating in the different affected areas. Rift Valley fever virus
isolates from the 2015 outbreak clustered in the Northeastern (Egyptian group) lineage with the strains (from mosquito: *Aedes ochraceus* and human) from Senegal in 2013 [27]. Emergence of RVF strains belonging to the Northeastern African lineage was already notified in West Africa, particularly in Senegal in 1983 (Figure 3), but no outbreak was reported. A reemergence or new introduction of this lineage was identified in Senegal in 2013 [27], and to our knowledge, this is the first evidence of the circulation in Mauritania of strains belonging to the Northeastern African lineage. These phylogenetic analyses suggest that this outbreak may result from an introduction of isolates from Barkedji, Senegal, or Egypt to Mauritania. Because Senegal and Mauritania shared a border and a previous study suggested that Barkedji functions as a hub broadcasting RVFV in the neighboring countries [28], we hypothesized that the outbreak in Mauritania might be due to an introduction of RVFV from Barkedji. This emphasized the need to implement surveillance and control measures after reported cases in Senegal to prevent propagation of RVFV isolates. However, further studies are needed to confirm the precise origin of Mauritania strains. These findings also confirm the existence of RVFV strain mixing between different geographic areas.

In pastoral countries like Mauritania, RVF epizootics are often responsible for serious economic losses and cross-border problems related to livestock transhumance. The reasons for the emergence or reemergence of RVF, the increasing frequency of outbreaks, and the reduced period between consecutive epidemics of RVF in Mauritania remain unknown [29]. However, there is a common observation that the occurrences of outbreaks in Mauritania are often preceded by large inflow of small ruminants when the traditional major Muslim holidays approach and are often associated with heavy rains and hydrographic changes, which result in the proliferation of RVF mosquito vectors.

The mechanism of RVFV spread likely depends on animal population transhumance, and migration routes between West Africa and other areas need to be identified to elucidate the different routes of introduction of the virus.

An RVF outbreak occurred in southern Mauritania in 2015 with 184 suspected cases, of which there were 57 confirmed cases and 12 deaths. Phylogenetic analysis of isolates indicated a virus origin from Northeastern Africa closely related to isolates detected in Senegal in 2013. Surveillance and control measures after reported cases in Senegal are needed to prevent propagation of RVFV isolates in neighboring countries. The identification of CCHV, West Nile, and dengue infections during the RVF outbreak indicates the co-circulation of various arboviruses and calls for systematic differential diagnostics or a syndromic approach for febrile and hemorrhagic fever surveillance to avoid a large-scale outbreak.

**Supplementary Data**

Supplementary materials are available at *Open Forum 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.

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