Emerging Filoviral Disease in Uganda: Proposed Explanations and Research Directions

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Abstract.
Outbreaks of Ebola and Marburg virus diseases have recently increased in frequency in Uganda. This increase is probably caused by a combination of improved surveillance and laboratory capacity, increased contact between humans and the natural reservoir of the viruses, and fluctuations in viral load and prevalence in this reservoir. The roles of these proposed explanations must be investigated to guide appropriate responses to the changing epidemiological profile. Other African settings in which multiple filoviral outbreaks have occurred could also benefit from such information.

INTRODUCTION

Marburgvirus and Ebolavirus, the two genera of the Filoviridae family of virus, cause human diseases known as Marburg virus disease (Marburg) and Ebola virus disease (Ebola), respectively. They are highly lethal diseases, with case fatality ratios typically between 20% and 90%, depending on the virus species.1–3 Evidence suggests that cave- and forest-dwelling fruit bats of the Pteropodidae family (suborder Megachiroptera) are the natural reservoir of both viruses.2–7

The frequency of filoviral disease outbreaks in Uganda has increased in recent years (Figure 1). Before 2007, there had only been one reported outbreak (of Ebola) in Uganda.8,9 Since 2007, nine filoviral disease outbreaks have occurred: six Ebola outbreaks and three Marburg outbreaks.10–16 One of these outbreaks in 2007 and 2008 involved Bundibugyo ebolavirus, which was the first new species of filovirus to be described in several years, an occurrence that has been ascribed to improved surveillance capacity.11,12

Similar patterns have been observed elsewhere: there were no reported outbreaks of Ebola globally between 1979 and 1994, but this time was followed by near simultaneous reemergence of the disease in Gabon, Cote D’Ivoire, and the Democratic Republic of the Congo (Figure 2).17,18 Similarly, there were no known cases of Marburg globally between 1987 and 1998,17 whereas three outbreaks have occurred in Uganda since 2007.

The reasons for the apparent increasing frequency of filoviral disease outbreaks in Uganda are unknown. We present the most likely explanations and suggest steps to further elucidate filoviral transmission dynamics and guide the healthcare response to the changing context.
POSSIBLE EXPLANATIONS OF CHANGING EPIDEMIOLOGICAL PROFILE

Changes to the surveillance system.

Improved sensitivity of filoviral disease surveillance in Uganda might account for the increased number of epidemics; the detection of isolated cases or small clusters would previously have gone undetected.

Awareness about filoviral diseases and their signs and symptoms is increasing among Ugandan health professionals, because experience grows with each subsequent outbreak. Investment in building surveillance capacity with the World Health Organization’s integrated disease surveillance and response strategy has also contributed. Political will and buy-in from the government of Uganda for strengthening surveillance combined with international assistance played a vital role in Uganda. International assistance included the establishment of a ready-to-use isolation ward by MSF in collaboration with the Ministry of Health, whereas the Centers for Disease Control and Prevention provided substantial technical and financial support for building capacity for the surveillance of viral hemorrhagic fevers in Uganda, particularly by improving capacity for laboratory diagnosis at the Uganda Virus Research Institute in Entebbe.

Educating communities and their health staff could result in additional improvements. Epidemics of filoviral disease frequently occur in settings with local beliefs in witchcraft and curses, which undermine surveillance and infection control at the peripheral level: affected individuals often seek healthcare from traditional healers first rather than at Ministry of Health-supervised facilities, during which time they remain undetected and can infect their contacts. This process also happens when formal healthcare providers do not suspect filoviral disease and when patients resist admittance to isolation wards during outbreaks because fear and stigmatization.

The improved sensitivity of surveillance activities raises the possibility that other outbreaks were not detected. Enhanced surveillance may, therefore, be providing a better understanding of filoviral transmission dynamics: low-level transmission, resulting in small, self-contained clusters among close contacts, may be more common than realized but are frequently missed by formal surveillance systems.

Changes in patterns of interaction between humans and bats.

There may be increasing interaction between humans and the natural reservoir of the filoviruses, perhaps through habitat encroachment or hunting activities. Commercial mining activity began at Kitaka cave in Ibanda district in January of 2007; within 9 months, four miners had contracted Marburg. A subsequent study reported that approximately 5% of the estimated 100,000 bats dwelling in Kitaka cave were infected by Marburgvirus. Two tourists visiting Uganda who developed Marburg disease on return to their countries both reported having visited the same cave inhabited by bats among which Marburgvirus circulates.

Changes in filoviral seroprevalence in the host species.

Cyclic population changes of the natural reservoir could result in changes in filoviral seroprevalence and an increased frequency of zoonotic transmission events. The seroprevalence of both Ebola virus and Marburgvirus in various bat species in Gabon and the Republic of Congo has been reported to fluctuate over time. Seasonal variation in Marburgvirus seroprevalence among Rousettus aegyptiacus has also been
shown, with peaks coinciding with the two times per year birthing seasons. The same study found a strong temporal correlation between historical epidemics of Marburg among humans and these seasonal peaks.

POSSIBLE SCENARIOS

Three scenarios are suggested by the above explanations.

1. The increase in outbreak frequency is perceived and not real. This scenario would result from successful strengthening of the national surveillance system. We would expect to see similar increases in other filoviral-endemic countries in which capacity building has been or will be implemented.
2. The increase is real but linked to factors specific to Uganda. In this scenario, the public health concern is limited in geographical scope, and we would not expect to observe an increase in outbreak frequency outside of Uganda.
3. The increase is real and caused by factors that are not limited to Uganda. In this scenario, an increase in outbreak frequency outside of Uganda would be expected.

The most likely of these scenarios is the first scenario, because strengthening of the Ugandan national surveillance system has demonstrably taken place in recent years. However, scenarios 2 and 3 are both plausible given the possible explanations described above, and therefore, they necessitate additional investigation as to their roles in the observed increased frequency of outbreaks of filoviral disease.

DISCUSSION

These scenarios imply different interpretations and would require different responses in those countries that experience more frequent filoviral disease outbreaks. However, the available evidence in support of any of these possible scenarios is mostly circumstantial; therefore, the roles that any, all, or none of these scenarios play in the increased frequency of filoviral disease outbreaks in Uganda must be investigated. Activities should focus on the following seven topics.

1. Reinforcing targeted surveillance activities, particularly around areas that have experienced multiple outbreaks and areas inhabited by bat colonies proven to be seropositive for filoviruses. Establishing in-country capacity for ongoing, sustained surveillance (with concomitant reduced reliance on external intervention) is essential for a variety of diseases, including filoviral disease, in the (rural) African context.
2. Establishment of additional ready to use isolation wards at hospitals near the most affected areas; one such ward has been established at Mbarara Hospital, which is close to Ibanda district. Such wards can be established and maintained with minimal investment—all that is required is a suitably sized building constructed of material that is easy to disinfect (e.g., concrete) and easy to isolate from the rest of the health facility—and a minimum of supplies (particularly personal protective equipment) to be kept in storage on site.
3. Improved in-country or regional capacity for accurate laboratory and differential diagnosis, which is underway in Uganda. The introduction of rapid testing for these diseases would also be a welcome, albeit secondary, measure.
4. Accelerated genotyping of circulating filovirus strains to determine the frequency of their introduction into the human population.
5. Seroprevalence surveys of filoviral infection in the general population of the most affected areas to estimate the prevalence of undetected disease in the at-risk population.
6. Additional studies to elucidate the ecology of filoviruses in bats are required. These studies include investigations of the mechanisms of natural infection and the processes of virus shedding to understand the pathways of transmission. In addition, studies of the prevalence of species-specific Ebola virus antibodies among bat populations, similar to those studies done for Marburg virus, are necessary to identify the most likely natural reservoirs, thereby defining the geographical scope of risk.
7. Ecological and serological comparisons between areas that have experienced multiple epidemics and areas that have experienced single epidemics, which may reveal factors in susceptibility to and transmission of filoviral disease.

These activities could help healthcare providers to adapt and respond appropriately to the new paradigm. This information is relevant not only for Uganda
but also, other African countries that have experienced outbreaks of filovirus disease during the past 45 years, particularly areas that have experienced multiple outbreaks, including the Western Equatoria state in South Sudan, Durba and Kampungu in the Democratic Republic of the Congo, and the Ogooué-Ivindo region of northeastern Gabon (Figure 2) (where Marburg virus has been found to be enzootic). Lessons from the Ugandan experience, whatever they may be, must be investigated, learned, and shared.

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REFERENCES


**FIGURE 1.** Timeline of outbreaks of filoviral disease known to have occurred in Uganda. This figure appears in color at www.ajtmh.org.

**FIGURE 2.** Map of the known outbreaks of filovirus disease in Central and East Africa. This figure appears in color at www.ajtmh.org.