Bombali Virus in *Mops condylurus* Bat, Kenya

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Bombali virus (genus *Ebolavirus*) was identified in organs and excreta of an Angolan free-tailed bat (*Mops condylurus*) in Kenya. Complete genome analysis revealed 98% nucleotide sequence similarity to the prototype virus from Sierra Leone. No Ebola virus–specific RNA or antibodies were detected from febrile humans in the area who reported contact with bats.

The virus family *Filoviridae* is divided into 5 genera: *Cuevavirus*, *Marburgvirus*, *Ebolavirus*, *Striavirus*, and *Thammovirus* (https://talk.ictvonline.org/taxonomy). Six distinct members of *Ebolavirus* have been described; 4 are known to cause human disease (1,2). These include highly lethal pathogens capable of producing large outbreaks, namely Bundibugyo, Sudan, and Zaire Ebola viruses, the last responsible for the devastating 2013–2016 outbreak in West Africa and an ongoing extended outbreak in the Democratic Republic of the Congo (1,3,4). Although the natural reservoirs of Ebola viruses remain unconfirmed, considerable evidence supports a role for bat species, particularly fruit bats, analogous to findings implicating *Rousettus aegypticus* fruit bats as a reservoir for Marburg virus (1,5,6).

The most recent Ebola virus to be identified is named Bombali virus (BOMV) and was reported in August 2018 in mouth and fecal swabs collected from free-tailed insectivorous bat species (family Molossidae) *Mops condylurus* and *Chaerephon pumilus* in Sierra Leone (2). Although BOMV is not known to infect humans, its envelope glycoprotein shares the same NPC1 receptor as other filoviruses and is capable of mediating BOMV pseudotype virus entry into human cells (2). We describe the presence of BOMV in tissues and excreta of an Angolan free-tailed bat (*M. condylurus*) captured near the Taita Hills in southeastern Kenya, the easternmost distributional range of this bat species (7), >5,500 km from the original BOMV identification site in Sierra Leone (Figure 1). We also screened human serum samples collected from febrile patients in the Taita Hills area for markers of BOMV infection.

We identified BOMV in an adult female bat (B241) by reverse transcription PCR and next-generation sequencing. This bat was captured along with 15 others in mist nets in savannah habitat near a small river in May 2018; only this bat was BOMV positive (6% prevalence). Viral RNA was present in lung, spleen, liver, heart, intestine, mouth swab, and fecal samples but absent from the brain, kidney, urine, and a few fleas found on the bat; viral loads were especially high in the lung (Appendix, https://wwwnc.cdc.gov/EID/article/25/5/18-1666-App1.pdf). These tissue-positive findings confirm that BOMV can infect *M. condylurus* and is not an artifact of its insect diet, which could not be discounted from the previous analysis on the basis of mouth and fecal swabs (2). We also screened lung samples of sympatric *C. pumilus* bats (n = 13) and other bat species (Appendix Table 2) captured from the same area in February 2016 and May 2018; all were negative for BOMV RNA. Serologic analysis revealed antibodies against BOMV in the blood of the tissue-positive bat (Appendix Figure), but specific antibodies were not found in blood from the other bats (Appendix).

Our tissue-positive findings provide a strong host association between BOMV and *M. condylurus* bats; it is possible that BOMV–positive findings from other bat species result from local spillover or contamination. Moreover, phylogenetic analysis of the full BOMV genome from the bat lung revealed 98% nucleotide sequence similarity with the prototype reported in Sierra Leone (GenBank accession no. MK340750) (Figure 2). Considering the high sequence similarity between the 2 locations and that BOMV can infect *M. condylurus*, like most insectivorous bats, are believed to travel only short distances (8), BOMV is likely to be distributed throughout much of sub-Saharan Africa.

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However, further monitoring of *M. condylurus* and *C. pumilus* bats and other sympatric species across Africa is required to support this hypothesis.

Because *M. condylurus* bats commonly roost in human structures, such as house roofs (8,9), human exposure to this species is more likely than for many other bat species. Therefore, we screened for markers of human infection with BOMV by studying serum samples collected from febrile patients who sought treatment at clinics in the Taita Hills area during April–August 2016. Clinics are located in the surrounding areas, all within 15 km of the BOMV–infected bat collection site (Figure 1). We screened patients for filovirus RNA (n = 81) and Ebola virus–specific IgG (n = 250) by an immunofluorescence assay using Zaire Ebola virus VP40–transfected VeroE6 cells as antigen (Appendix). Many samples, including

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**Figure 1.** Locations of Bombali Ebola virus infection in Sierra Leone (gray shading at left; Bombali district in red) and Kenya (gray shading at right; Taita Hills area in green). Inset map shows collection site of the Bombali virus–positive bat (red dot) in Kenya, clinics in which human serum samples were collected (white squares), and the closest towns (black squares).

**Figure 2.** Phylogenetic tree of complete filovirus genomes (18,795–19,115 nt), including Bombali Ebola virus in Sierra Leone and now Kenya (19,026 nt; black dot). Representative sequences were retrieved from the Virus Pathogen Database and Analysis Resource and aligned with a MAFFT online server (http://mafft.cbrc.jp/alignment/software). The tree was built using the Bayesian Markov Chain Monte Carlo method, using a general time-reversible model of substitution with gamma-distributed rate variation among sites allowing the presence of invariable sites. Posterior probabilities are shown at the nodes. Scale bar indicates genetic distance.
all those screened for filovirus RNA, were from patients who reported contact with bats in the home or workplace. We found no evidence of filovirus infection by either screening method, providing no support that BOMV easily infects humans or is a common cause of febrile illness in the area. Ongoing surveillance is nonetheless necessary, and we cannot exclude the possibility that Bombali virus was a recent introduction to the Taita Hills area.

Our results markedly expand the distributional range of this new Ebola virus to eastern Africa and confirm the \textit{M. condylurus} bat as a competent host. Like Goldstein et al. (2), we stress that the virus is not known to infect humans, a premise supported by our screening of febrile patients in the Taita Hills area. Potential efforts to eradicate bats are unwarranted and may jeopardize their crucial ecosystem roles and human health (10,11).

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\textbf{About the Author}

Dr. Forbes is a disease ecologist and assistant professor at the Department of Biological Sciences, University of Arkansas. His research interests include the maintenance and transmission of rodentborne and batborne zoonotic pathogens in nature and the effects of anthropogenic environmental changes on these processes.

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Appendix

**Methods**

Bats were captured in February 2016 and May 2018 as part of an ongoing virus screening project in the Taita Hills area of rural Kenya; all *M. condylurus* were captured in 2018. We employed mist and hand netting, and structured trapping site selection to focus on habitat and species diversity and minimize the number of individuals collected from any 1 species or site. Captured bats were placed into individual cotton bags, and processed at the University of Helsinki Taita Research Station. Species identifications were made in the field using keys (1). Non-conservation priority bat species (classified as least concern by the IUCN) were euthanized via cervical dislocation to collect blood, lung, liver, spleen, kidney, intestine and brain samples, as well as urine, feces, and ectoparasites when possible. Dissections were performed in a sheltered outside area, using personal protective equipment, including FFP3 facemasks, latex gloves, and safety gowns. Bat tissues were placed into separate marked tubes with RNAlater (Sigma, https://www.sigmaaldrich.com), stored at −20°C, and later sent on dry ice to Helsinki, Finland.

At the University of Helsinki, under enhanced BSL-3 conditions, bat tissue samples were treated with Tripure (Roche, http://www.roche.com) to inactivate any potential hazardous agents before RNA extractions (Tripure method) and screening by a pan-filovirus RT-qPCR (2). Filovirus screening was initially conducted as a precaution, to facilitate screening for other viruses under less-strict biosafety conditions. The pan-filo RT-qPCR has been tested to detect Zaire EBOV, Bundibugyo, Sudan, Taï Forest, and Reston ebolavirus, in addition to Marburg virus (MARV) and Ravn Virus RNA. EBOV and MARV RNA were used as positive controls (in vitro RNAs) (3). Following the identification of a positive individual, with particularly high viral loads in the lung, lung samples from all bats were also screened with a Bombali virus–specific
real-time RT-PCR (4). All tissue, excreta, and ectoparasite samples were screened from the positive individual (Appendix Table 1), and viral loads determined by RT-qPCR with an in vitro transcribed RNA serving as the quantification standard. A full list of each bat species captured and screened is provided in Appendix Table 2.

Prior to whole-genome sequencing, RT-PCR positive samples were treated with DNase I (Thermo Fisher, http://www.thermofisher.com), and purified with Agencourt RNA Clean XP magnetic beads (Beckman Life Sciences, https://www.beckman.com). Ribosomal RNA was removed using a NEBNext rRNA depletion kit (New England BioLabs, https://www.neb.com), according to the manufacturer’s protocol. The sequencing library was prepared using a NEBNext Ultra II RNA library prep kit (New England BioLabs). Libraries were quantified using a NEBNext Library Quant kit for Illumina (New England BioLabs). Pooled libraries were then sequenced on a MiSeq platform (Illumina, https://www.illumina.com) using a MiSeq v3 reagent kit with 300 bp paired-end reads. Raw sequence reads were trimmed and low-quality (quality score <15) and short (<36 nt) sequences were removed using Trimmomatic (5). Thereafter, de novo assembly was conducted using MegaHit (6). Open reading frames were sought using MetaGeneAnnotator (7), followed by taxonomic annotation using SANSparallel (8). We confirmed bat species identity of the positive individual by retrieving cytochrome-b sequences from the NGS reads (GenBank accession no. MK330941).

The phylogenetic tree was constructed using the Bayesian Markov chain Monte Carlo (MCMC) method, implemented in Mr Bayes version 3.2 (9) using a GTR-G-I model of substitution with 2 independent runs and 4 chains per run. The analysis was run for 5 million states and sampled every 5,000 steps. The average standard deviation of split frequencies was 0.000732.

Febrile patients seeking care at 3 health facilities in the Taita Hills (Wundanyi, Mwatate, and Voi) were recruited into the study by clinicians. A questionnaire was used to capture socio-demographic data and pertinent history, including a tickbox question regarding contact with bats at home or work. Based on the criterion of exposure to bats, a total of 81 patients (2.9–83.4 years of age; average, 38.8 years) were selected for analysis of filovirus RNA. Samples were collected within 5 days of the onset of fever. No patients reported bleeding. Reported symptoms included, in addition to fever; myalgia (54/81), joint pain (45/81), rash (9/81), diarrhea (8/81), vomiting
(7/81), headache (6/81) and cough (4/81). Serum samples were stored at the University of Helsinki Taita Research station at −20°C for ≤3 weeks, and then transported on ice to a central laboratory at the University of Nairobi where they were stored at −80°C and later shipped on dry ice to Helsinki. Nucleic acids were extracted from 100µL of serum and eluted to 50µL using the QIAamp Viral RNA Mini Kit (QIAGEN, https://www.qiagen.com) according to manufacturer’s instructions. Pan-filovirus RT-qPCR was then conducted as described above, as well as Bombali virus–specific RT-PCR (4).

Human serum samples were analyzed for Ebola virus–specific IgG antibodies using an immunofluorescence assay (IFA) based on a recombinant Zaire ebolavirus VP-40 with a similar IFA protocol as described before (10), and demonstrated within the EbolaMoDRAED EU-IMI project to react with Zaire ebolavirus patient serum. Bombali virus VP40 protein is 75%–78% similar to that of other ebolaviruses, which have been demonstrated to cross-react within the genus (11). As antigen, we used acetone-fixed Vero E6 cells transfected with the pCAGGS-Ebola VP40 construct (Zaire ebolavirus, isolate Ebola virus/ H.sapiens-wt/SLE/2014/Makona-G3856.1 sequence, GenBank KM233113.1), and as controls, cells transfected with the empty vector. Patient serum samples were diluted 1:60 in PBS and incubated for 1 h at 37°C. Fluorescein isothiocyanate–conjugated anti-human IgG (Jackson ImmunoResearch, https://www.jacksonimmuno.com) was diluted 1:30 in PBS, and incubated for 30 min at 37°C. Unbound antibodies and anti-human IgG were washed 3 times with PBS and then once with distilled water. The slides were covered with mounting medium and coverslips, and read using a ×20 objective of fluorescence microscope Olympus IX71 (Olympus Corporation, www.olympus-global.com).

Additional Results

Serologic analysis revealed antibodies against ebolavirus in the blood of the tissue-positive bat (Appendix Figure), but antibodies were not present in blood from the other bats. Note that bat blood samples (from RNA-negative individuals) were first heat inactivated under enhanced BSL-3 conditions. To minimize exposure risk, the blood sample from the positive bat was sent to the Public Health Agency of Sweden and screened under BSL-4 conditions. To detect bat antibodies in blood samples, Vero E6 cells transfected as above to produce ZEBOV
VP40, or at Public Health Agency of Sweden, infected with Zaire ebolavirus, were used in IFA according to a previously described protocol (12). Blood samples were diluted to 1:20 in PBS before incubation. Detection was done with goat anti–bat antibody Ig (Bethyl Laboratories, https://www.bethyl.com) at 1:1,000, followed by donkey anti–goat cyanin 2 (Cy2)-labeled Ig (Dianova, https://www.dianova.com) at 1:100. Slide staining and analysis were conducted as described above.

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**Appendix Table 1.** Viral loads from *Mops condylurus* bat that tested positive for Bombali Ebola virus.

| Sample      | Ct value | Copy number/500 ng total RNA |
|-------------|----------|------------------------------|
| Mouth swab† | 24.00    | Not applicable               |
| Spleen      | 32.76    | 414                          |
| Liver       | 33.95    | 181                          |
| Intestine   | 32.76    | 413                          |
| Heart       | 29.82    | 3,173                        |
| Feces       | 29.14    | 5,121                        |
| Lung        | 16.74    | 27,950,000                   |
| Kidney      | Negative | 0                            |
| Urine       | Negative | 0                            |
| Fleas       | Negative | 0                            |

*Viral loads for each sample type were estimated using a standard curve based on in vitro transcribed and quantified RNA.
†Mouth swab has no copy number because it was screened in a BSL-4 laboratory in Sweden using a different protocol and without the standard curve.

**Appendix Table 2.** Bat species screened for filoviruses, Kenya*

| Species                      | 2016 | 2018 |
|------------------------------|------|------|
| Mops condylurus              | 0    | 16   |
| Chaerephon pumilus           | 4    | 7    |
| Cardioderma cor              | 36   | 20   |
| Chaerephon chapini           | 1    | 0    |
| Epomophorus wahlbergi        | 19   | 23   |
| Glauconycteris argentata     | 1    | 0    |
| Hipposideros caffer          | 2    | 1    |
| Lavia trons                  | 0    | 1    |
| Lissonycteris angolensis     | 10   | 0    |
| Myotis tricolor              | 1    | 0    |
| Neoromicia nana              | 0    | 3    |
| Nycticeinops schlieffeni     | 3    | 1    |
| Nycteris thebaica            | 0    | 1    |
| Rhinolophus clivosus         | 0    | 2    |
| Rhinolophus landeri          | 1    | 2    |
| Species                     | 2016 | 2018 |
|----------------------------|------|------|
| *Rousettus aegyptiacus*    | 17   | 13   |
| *Scotoecus hirundo*       | 34   | 4    |
| *Scotophilus dinganii*    | 4    | 12   |
| *Rousettus lanosus*       | 0    | 1    |
| *Pipistrellus sp.*        | 20   | 0    |
| *Neoromicia sp.*          | 0    | 1    |
| *Miniopterus sp.*         | 6    | 0    |
| *Hypsugo sp.*             | 4    | 0    |

*Bats were captured from the Taita Hills area in 2016 and 2018. All bat lung samples were screened for filovirus RNA via a new pan-filovirus reverse transcription qualitative PCR (2) and a Bombali virus–specific real-time reverse transcription PCR (4).

**Appendix Figure.** Detection of Ebola virus–specific antibodies in Bat B241 (the BOMV RNA positive individual) using an immunofluorescence assay based on Zaire ebolavirus (ZEBOV)–infected, acetone-fixed Vero E6 cells. The slides contain ZEBOV-infected and noninfected control cells. A) 4′,6-diamidino-2-phenylindole (DAPI) staining for cell nuclei. B) Staining with rabbit anti–ZEBOV-GP showing ZEBOV-infected cells. C) Staining with bat B241 serum at a dilution of ≈1:200, demonstrating specific granular staining of ZEBOV-infected cells. D) A merge of stains demonstrating that the antibody response of bat B241 is Ebola virus genus cross-reactive, but targeting other viral proteins than the ZEBOV GP.