and possible early warning of BRBV transmission risk. However, whether any of these mammalian species are competent amplifier hosts for BRBV remains to be determined.

Acknowledgments
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About the Author
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References

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Fatal Case of Lassa Fever, Bangolo District, Côte d’Ivoire, 2015

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**Table.** PRNT_{70} results for mammals tested for Bourbon virus–neutralizing antibodies, Missouri, USA, 2012–2013*

<table>
<thead>
<tr>
<th>Common name</th>
<th>Species name</th>
<th>No. positive/no. tested</th>
<th>Titer</th>
<th>Proportion positive (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Domestic cat</td>
<td>Felis catus</td>
<td>0/2</td>
<td>&lt;10</td>
<td>0 (0.0–0.66)</td>
</tr>
<tr>
<td>Domestic dog</td>
<td>Canis lupus familiaris</td>
<td>2/13</td>
<td>10–320</td>
<td>0.15 (0.04–0.42)</td>
</tr>
<tr>
<td>Eastern cottontail</td>
<td>Sylvilagus floridanus</td>
<td>2/9</td>
<td>&gt;320</td>
<td>0.22 (0.06–0.55)</td>
</tr>
<tr>
<td>Fox squirrel</td>
<td>Sciurus niger</td>
<td>0/4</td>
<td>&lt;10</td>
<td>0 (0.0–0.49)</td>
</tr>
<tr>
<td>Horse</td>
<td>Equis caballus</td>
<td>1/24</td>
<td>20</td>
<td>0.04 (0.007–0.20)</td>
</tr>
<tr>
<td>Raccoon</td>
<td>Procyon lotor</td>
<td>31/62</td>
<td>10–320</td>
<td>0.50 (0.38–0.62)</td>
</tr>
<tr>
<td>Virginia opossum</td>
<td>Didelphis virginiana</td>
<td>0/28</td>
<td>&lt;10</td>
<td>0 (0.0–0.12)</td>
</tr>
<tr>
<td>White-tailed deer</td>
<td>Odocolites virginianus</td>
<td>12/14</td>
<td>10–320</td>
<td>0.86 (0.60–0.96)</td>
</tr>
</tbody>
</table>

*PRNT_{70}, 70% plaque reduction neutralization titer.
Lassa fever has not been reported in Côte d’Ivoire. We performed a retrospective analysis of human serum samples collected in Côte d’Ivoire in the dry seasons (January–April) during 2015–2018. We identified a fatal human case of Lassa fever in the Bangolo District of western Côte d’Ivoire during 2015.

Lassa fever is endemic to western Africa. Nigeria, Guinea, Sierra Leone, and Liberia regularly have outbreaks of Lassa fever, mostly during the first few months of the year, corresponding to the dry season (January–May), when the Mastomys natalensis rodent reservoir of Lassa fever virus (LASV) has more contact with the human population in rural areas to access food. The epidemic zone of Lassa fever has recently been extended into Benin, and sporadic cases have been documented in Burkina Faso, Mali, Ghana, and Togo.

Côte d’Ivoire appears to be an exception; no Lassa fever cases have been reported in this country. A tourist from Germany traveling through Côte d’Ivoire, Burkina, and Ghana died from Lassa fever upon her return to Germany but it was not possible to determine in which country she contracted the disease (1). In 2013, LASV RNA was identified in M. natalensis rodents captured in northern Côte d’Ivoire, near Korhogo (2). Virus RNA corresponded to the same AV strain of LASV as that isolated from the tourist, suggesting that she might have been infected in Côte d’Ivoire. Seroprevalence among forest workers in western Côte d’Ivoire also suggests that LASV might currently circulate in this country (3).

We performed a retrospective analysis of 268 human serum samples received at the National Reference Center for Yellow Fever (Institut Pasteur de Côte d’Ivoire, Abidjan, Côte d’Ivoire) for diagnosis of arbovirus infection. We selected yellow fever–negative samples from the western region of Côte d’Ivoire (Biankouma, Danané, Duékoué, Guiglo, Man, Odienné, Touba, Toulépleu, and Zouan Hounien Districts), near the borders with Liberia and Guinea collected during January–April 2015–2018.

We inactivated serum samples by using AVL buffer (QIAGEN, https://www.qiagen.com) and ethanol and isolated RNA by using the QIAamp Viral RNA Extraction Kit (QIAGEN). We analyzed RNA by using a reverse transcription PCR (RT-PCR) and pan–Old World arenavirus primers specific for the large RNA segment (OW RT-PCR).

We identified 1 positive serum sample (001/15) by OW RT-PCR; we then determined that this sample was LASV positive by using a LASV-specific RT-PCR specific for the glycoprotein complex (GPC) gene (Figure, panel A). We sequenced amplicons for the GPC and L genes and aligned partial sequences of this new strain, Bangolo-CIV-2015, with the corresponding regions of a set of representative published LASV strains (4).

We generated phylogenetic trees by using a general time reversible plus gamma plus proportion of invariable sites model and parallel maximum-likelihood with PhyML Smart Model Selection (5). The GPC (MK978784) and large RNA (MK978785) fragments of the Bangolo-CIV-2015 strain (Figure, panel B) were genetically similar to strain BA-366 isolated from an M. natalensis rodent captured in the Bantou District of central Guinea during 2003 (6). On the basis of these phylogenetic trees, we concluded that Bangolo-CIV-2015 belongs to the IV clade, along with the highly pathogenic LASV strain Josiah, and diverges from the clade V AV strain of LASV (Figure, panel B) found in rodents and a patient from Germany (1,2).

The LASV-positive serum sample originated from a 30-year-old man from the Bangolo District of Côte d’Ivoire who was admitted to Duékoué Hospital in January 2015 because of fever, asthenia, and gingivorrhagia. His health rapidly deteriorated after admission; he had hypotension and a consciousness disorder and died 4 days later. The sample was collected at the time of death. Further investigations by doctors at Institut Pasteur de Côte d’Ivoire were unable to obtain more information about this patient. Without the travel history of the patient during the 3 weeks preceding his hospital admission, we could not determine whether this case of Lassa fever was endemic or imported. Exported cases are common (7) because many workers travel to Côte d’Ivoire from Guinea and Sierra Leone.

Next-generation sequencing in an outbreak setting, combined with phylogenetic analyses, has recently showed that many strains of LASV have been responsible for cases of Lassa fever in Nigeria, suggesting independent transmission events from the reservoir, rather than the emergence of an epidemic strain (8). The case we report remains isolated because no other suspected cases were reported during this period. None of the healthcare workers who had been in contact with the patient showed any signs of Lassa fever. An additional set of 35 human serum samples collected during October 2014–April 2015 in the Biankouma, Duékoué, Guiglo, Issia, Man, Minignan, Odienné, Soubre, and Tengrela Districts were LASV negative by RT-PCR.

In 2015, Kouadio et al. highlighted possible underreporting of Lassa fever cases in Côte d’Ivoire because of lack of diagnoses (2). We provide evidence of a fatal case of human Lassa fever in the Bangolo District of western Côte d’Ivoire. Thus, measures should be taken to reinforce the diagnosis of Lassa fever and arenavirus surveillance in general in this country. Human serologic surveys should help in identifying the area of LASV circulation in Côte d’Ivoire. RNA from novel arenaviruses has recently been identified in rodents captured in Côte d’Ivoire, but their pathogenic potential for humans remains unknown (9).
Acknowledgments

We thank the Ministère de la Santé et de l’Hygiène Publique de Côte d’Ivoire and personnel from Duékoué Hospital for providing information about the patient; all members of Département des Virus Epidémiques, Plateforme de Biologie Moléculaire, and Département Technique et Technologie at IPCI and West African Biobank for assistance and support during the visit of M.M. to the IPCI; Ghislain Tiemoko Gueu for technical assistance in managing human samples; N’cho Sonan for helpful discussions; Marc Jouan and Marianne Lucas-Hourani for administrative and logistical assistance; and Frédéric Lemoine for assistance with phylogenetic analyses.

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References


Figure. Analysis of LASV strains, Bangolo District, Côte d’Ivoire, 2015. A) RT-PCR analysis of human serum samples 132/16 and 001/15 by using OW and LASV RT-PCRs. Lane 1, negative control; lane 2, positive control; lane 3, 132/16; lane 4, 001/15. B) Phylogenetic analysis of LASV strains. Trees were inferred by using the PhyML Smart Model Selection (5) general time-reversible plus gamma plus proportion of invariable sites model with 200 bootstrap replicates. Poorly supported branches with bootstrap values <0.50 are indicated by dotted lines. Lassa virus lineages are indicated by the Roman numerals on the right. The Bangolo-CIV-2015 strain (solid box), which was isolated in this study, appears to be related to clade IV, and the AV strain (dotted box) is related to clade V. GPC, glycoprotein complex gene; L, large RNA segment; LASV, Lassa virus; OW, Old World; RT-PCR, reverse transcription PCR.
Household Transmission of Human Adenovirus Type 55 in Case of Fatal Acute Respiratory Disease

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We identified a case of fatal acute respiratory disease from household transmission of human adenovirus type 55 (HAdV-55) in Anhui Province, China. Computed tomography showed severe pneumonia. Comparative genomic analysis of HAdV-55 indicated the virus possibly originated in Shanxi Province, China. More attention should be paid to highly contagious HAdV-55.

1These authors contributed equally to this article.

Human adenoviruses are associated with mild and acute respiratory infections, depending on the virus type and host immunity. Human adenovirus type 55 (HAdV-55) (1), formerly known as HAdV-11a (2), is a reemergent respiratory pathogen that has caused severe pneumonia outbreaks in military and civilian populations in Europe and Asia (2–7). However, household transmission of HAdV-55 is rarely reported. We report a case of household transmission of HAdV-55 involving 3 confirmed adult cases with 1 death. Epidemiologic, clinical, and laboratory investigations, along with whole-genome sequencing, elucidate the disease progression and the pathogen origin.

During April 1–May 5, 2012, 7 household members (5 males and 2 females; 3 children and 4 adults) in Anhui Province, China, sequentially experienced influenza-like symptoms, including fever, productive cough, fatigue, pharyngalgia, dyspnea, and other symptoms. The youngest patient was 4 months of age, the oldest, whom we refer to as AQ-1, was a 55-year-old man. The family lived together near a farm in a house with poor sanitary and ventilation conditions.

The first onset of acute respiratory disease (ARD) occurred on April 1, when the index case, a 4-year-old granddaughter of AQ-1, had a febrile respiratory infection with cough. Three days later, AQ-1’s grandson, 1 year of age, displayed similar symptoms. On April 9 and 11, AQ-1’s daughter, 28 years of age, and another grandson, 4 months of age, both had influenza-like symptoms. On April 14, AQ-1 had a fever, chills, and lumbago. He was admitted to the hospital on April 14 where clinicians diagnosed pneumonia. AQ-1 had close contact with his sick grandchildren and granddaughter and had not been out of the house during the month he cared for them.

While hospitalized, AQ-1 had bilateral pneumonia seen on chest computed tomography (CT), a temperature of 41.0°C, and low total leukocyte (3.63 × 10⁹/L) and platelet (42 × 10⁹/L) counts. AQ-1 sustained high fever and yellow phlegm despite antiinflammatory and antiviral treatment, including levofloxacin, piperacillin sodium, tazobactam sodium, and ribavirin.

On April 24, AQ-1 had indications of severe pneumonia, including respiratory failure, hypoxemia, double lung rales, and a mass of shadows visible on chest CT. In addition, he had indications of liver damage and multi-organ failure. Transverse chest CT images demonstrated increased areas of patchy shadows and consolidation in both lungs compared to CT images from April 22, indicative of disease progression (Appendix Figure 1, http://wwwnc.cdc.gov/EID/article/25/9/18-1937-App1.pdf).

AQ-1 died on April 27, 3 days after onset of respiratory failure, and 13 days after his illness began. On the same day, his 20-year-old son, AQ-2, and 31-year-old nephew, AQ-3, who had taken care of AQ-1 for 5 days, also exhibited...