A Recently Isolated Lassa Virus From Mali Demonstrates Atypical Clinical Disease Manifestations and Decreased Virulence in Cynomolgus Macaques

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The virulence of Soromba-R, a Lassa virus strain recently isolated from southern Mali, was assessed in 2 animal models of Lassa fever: inbred strain 13 guinea pigs and cynomolgus macaques. In both models, the Malian isolate demonstrated tissue tropism and viral titers similar to those of historical Lassa virus isolates from Sierra Leone (Josiah) and Liberia (Z-132); however, the Soromba-R isolate was found to be less pathogenic, as determined by decreased mortality and prolonged time to euthanasia in macaques. Interestingly, in addition to the classic indicators of Lassa fever, Soromba-R infection presented with moderate to severe pulmonary manifestations in the macaque model. Analysis of host responses demonstrated increased immune activation in Soromba-R–infected macaques, particularly in neutrophil-activating or -potentiating proinflammatory cytokines or growth factors, including tumor necrosis factor α, macrophage inflammatory protein 1α, interleukin 1β, and granulocyte colony-stimulating factor, as well as interleukin 5, which may be responsible for the decreased lethality and uncharacteristic clinical presentation. These results suggest that the strain of Lassa virus circulating in Mali might be less pathogenic than strains circulating in the historical region of endemicity and may result in an atypical presentation for Lassa fever, which could complicate clinical diagnosis.

Keywords. Lassa fever; pathogenesis; disease modeling; guinea pigs; non-human primates; West Africa.

Lassa fever (LF) is an acute viral illness associated with Lassa virus (LASV; family Arenaviridae, genus Arenavirus) infection. LF is endemic in the West African countries of Nigeria, Liberia, Guinea, and Sierra Leone, where it represents a major burden on healthcare systems, with up to 500,000 combined infections identified per annum [1–4]. Sporadic cases of LF have also been diagnosed from countries in close proximity to the LF-endemic regions, including a confirmed case suspected to have originated from Ghana, Cote d’Ivoire, or Burkina Faso and, more recently, a confirmed case from Mali, where LASV was shown to be enzootic [5–7].

In humans, exposure to LASV can result in a range of manifestations, from apparently asymptomatic infection to severe hemorrhagic fever disease with multiorgan failure [1]. In the LF-endemic region, LASV is
associated with approximately 5000 deaths annually, representing an overall mortality rate of 1%–2% among infected individuals [4]. However, the mortality rate of LF among hospitalized cases increases to around 20%, and mortality rates exceeding 50% have been documented in outbreak scenarios [8, 9]. The remote locations where LASV is considered endemic, coupled with the risk of secondary transmission of LASV, has hindered clinical and pathological investigations into LF. As such, the pathogenesis of LF remains poorly understood [10].

Currently, 2 animal models have been described for studying LF: inbred (strain 13) guinea pigs and nonhuman primates (NHPs) [11]. While the inbred guinea pig model is useful as a screening model, the disease manifestations in this model do not recapitulate those seen in human cases; therefore, the preferred model for pathogenesis and vaccine or therapeutic studies is NHPs, most commonly rhesus or cynomolgus macaques. To date, the majority of studies conducted in NHPs have focused on infection with LASV strain Josiah [12–18]. The purpose of this study was to use the strain 13 guinea pig and cynomolgus macaque models of LF for infection with a recent LASV isolate from Mali and to compare potential differences in virulence with LASV strains from Liberia and Sierra Leone.

METHODS

Animal Work and Biosafety
Animal experiments were approved by the Institutional Animal Care and Use Committee of the Rocky Mountain Laboratories (RML) and were performed following the guidelines of the Association for Assessment and Accreditation of Laboratory Animal Care, International (AAALAC), by certified staff in an AAALAC-approved facility. Experiments were conducted in the biosafety level 4 (BSL-4) facility at the RML.

Viruses
Low-passage (defined as ≤4 passages) LASV isolates Josiah [19], Z-132 (which genetically is similar to LASV Z-148) [7, 20], and Soromba-R [7] were cultured in Vero cells and titered using standard median tissue culture infective dose (TCID50) methods.

In Vivo Infection (Strain 13 Guinea Pigs)
Twenty-one strain 13 guinea pigs (Cavia porcellus; age, 2–4 months) were infected with a TCID50 of 1 × 10^4 LASV Josiah, Z-132, or Soromba-R by intraperitoneal injection and were monitored daily for signs of illness (ie, abnormal activity, posture, and respiration; recumbency; and weight loss). Animals in the terminal stage of disease (weight loss of >15%, inability to move, and/or respiratory distress) were euthanized by exsanguination, and blood, liver, spleen, and lung samples were collected for pathological and virological analysis from 3 animals per group.

In Vivo Infection (Cynomolgus Macaques)
Nine cynomolgus macaques (Macaca fascicularis; age, 6–7 years; sex, 7 males and 2 females) were infected with 1 × 10^4 TCID50 of LASV Josiah, Z-132, or Soromba-R (3 animals per group) by intramuscular injection. Animals were monitored twice daily and assigned a numerical score based on clinical signs of disease (ie, fever; abnormal posture, respiration, feces/urine, food intake, attitude, and skin turgor; and recumbency), using an approved end point scoring sheet. On days −7, 0, 1, 3, 7, 10, and 14, animals were examined under anesthesia, at which point chest radiography was performed; pulse rate, blood pressure, temperature, and respiration rates were measured; and animals were bled for analysis of blood chemistry, coagulation parameters, differential blood count, virologic characteristics, and cytokine profiling. Animals were euthanized (by exsanguination while under deep anesthesia) when clinical signs indicated terminal disease (ie, a score of ≥30) [21], and a complete necropsy was performed, with collection of clinical specimens from nasal mucosa, oral mucosa, conjunctiva, tonsils, salivary gland, trachea, bronchi, right and left lung (upper, middle, and caudal lobes), lymph nodes (cervical, axillary, inguinal, and mesenteric) heart, liver, spleen, pancreas, jejunum, transverse colon, kidney, adrenal gland, testes/ovary, femoral bone marrow, urinary bladder, cervical spinal cord, and brain (stem, frontal, and cerebellum).

Hematological, Serum Biochemical, and Coagulation Parameters
Hematological, serum biochemical, and coagulation analyses were accomplished as previously described [18, 22, 23].

Serum Cytokine/Chemokine Analysis
Concentrations of granulocyte colony-stimulating factor (G-CSF), granulocyte macrophage colony-stimulating factor, interferon-γ (IFN-γ), interleukin 1β (IL-1β), interleukin 1Ra, interleukin 2 (IL-2), interleukin 4, interleukin 5 (IL-5), interleukin 6 (IL-6), interleukin 8 (IL-8), interleukin 12/23 (IL-12/23), interleukin 13, interleukin 15, interleukin 17, interleukin 18, monocyte chemotactic protein 1 (MCP-1), macrophage inflammatory protein 1α (MIP-1α), macrophage inflammatory protein 1β, sCD-40L, tumor necrosis factor α (TNF-α), transforming growth factor α, and vascular endothelial growth factor were determined in serum samples, using a NHP cytokine kit (Millipore). Data were analyzed with 2-way analysis of variance (ANOVA) and a Bonferroni posttest comparing Soromba-R and Z-132 infection groups to Josiah.

Virus Detection/Titration
RNA was extracted from NHP blood samples by use of QIAamp viral RNA extraction kits (Qiagen) and was screened...
for the presence of LASV RNA as previously described [24]. Infectious LASV titers were determined in blood and tissue samples by a standard TCID50 assay on Vero cells (limit of detection, approximately 10^3 infectious particles/g). Viral titers were analyzed with 1-way ANOVA and the Tukey-Kramer posttest.

**Histopathological and Immunohistochemical (IHC) Analysis**

Tissues were inactivated and fixed with 10% formalin and processed according to standard methods and BSL-4 standard operating procedures. Thin sections were stained with hematoxylin and eosin or tested for viral antigen by IHC, using a monoclonal antibody against LASV GP2 (1:200 dilution, anti-tissue in addition to data not shown). Kindly provided by Dr. Lisa Hensley, US Army Medical Research Institute of Infectious Diseases [18] on a Discovery XT instrument (Ventana Medical Systems).

**RESULTS AND DISCUSSION**

**LASV Infection in Inbred Guinea Pigs**

LASV Josiah and Z-132 were uniformly lethal in inbred guinea pigs, with mean times to death of 20.7 days (range, 18–26 days) and 19.9 days (range, 17–23 days), respectively. In contrast, infection with LASV Soromba-R was lethal in 4 of 7 animals (57%), with a mean time to death of 20.8 days (range, 19–22 days; Figure 1A). The 3 survivors in the Soromba-R group were monitored until 55 days after infection, and although minor weight loss was noted, no other signs of disease were observed. Infection was confirmed in the survivors through identification of anti-LASV antibodies via recombinant nucleoprotein enzyme-linked immunosorbent assay [25].

Viral titers in 3 organs (lung, liver, and spleen) and blood collected from terminally ill animals revealed similar, high levels of infectious LASV (Figure 1B), and IHC staining of tissues demonstrated little difference between the 3 infection groups. Macrophages and pneumocytes stained positive in lung samples, with minimal positivity noted in the endothelium. In the liver, the mesothelium frequently stained positive for LASV antigen and, to a lesser extent, periductal monocytes. Histological changes in lung, liver, and spleen samples were also similar and consistent across all 3 groups. Changes in the spleen were characterized by mild to moderate red pulp histiocytosis (hyperplasia of histiocytes/macrophages), with variable but fewer numbers of heterophils. Liver samples exhibited hepatocellular lipid degeneration, as well as mild to moderate lymphohistiocytic portal hepatitis, which appeared to be centered primarily on bile ductules. Moderate, multifocal, bronchointerstitial pneumonia, primarily histiocytic with fewer heterophils, was noted in lung samples. In addition, there was multifocal alveolar septae expanded by macrophages with abundant, vacuolated cytoplasm. Multifocal, mild, perivascular aggregates of lymphocytes were also noted, and the pleural mesothelium was hypertrophic. Representative lung and liver samples from a Soromba-R–infected guinea pig are shown in Supplementary Figure 1.

**Clinical Progression of LASV Infection in NHPs**

Physical signs of infection were unremarkable in cynomolgus macaques until days 5–7 after infection, at which point animals demonstrated reduced food intake, decreased activity, mild depression, and piloerection (Figure 2A). Fever was noted in 5 animals infected with Josiah and Z-132 isolates and peaked between days 7 and 10 (peak temperature, 40.3°C; average, 39.75°C). A single animal infected with Z-132 and all 3 animals infected with Soromba-R had no detectable fever during infection. Facial edema was apparent in half of the terminally ill animals. Similar to the results of the guinea pig study, infections with Josiah and Z-132 were 100% lethal in macaques, with mean times to death of 11.3 and 11.7 days, respectively, whereas infection with Soromba-R was lethal in 2 animals, with a prolonged time to death (average time to death, 15.5 days; Figure 2B).

**Serum Biochemical Findings**

A summary of the serum biochemical findings can be found in Table 1. Liver indices showed an overall trend toward increase levels of transaminases (alanine aminotransferase [ALT] and aspartate aminotransferase [AST]) during infection. On average, AST values in all groups were higher than ALT values. ALT levels began to spike at day 10 after infection, peaked around day 13 after infection, and, only in the Soromba-R group, began to normalize. Levels of albumin and total protein had a downward trend across all infection groups. This is likely a result of redistribution into the extracellular space, secondary to capillary leak syndrome, as can occur in sepsis. These decreases could also be a component of decreased hepatic synthesis; however, the slow trend downward is suggestive of inflammation-mediated capillary leakage rather than poor hepatic synthesis, because liver failure tends to occur more abruptly. Alkaline phosphatase (ALP) levels increased up to the time of euthanasia in all but 1 case of Soromba-R infection, also suggesting liver injury in most animals. Usually, the ALP level is elevated in cholestatic, infiltrative, and inflammatory liver diseases, and any or all of these processes can be occurring with arenavirus infection. The γ-glutamyltransferase values also increased, which signifies that the process was likely cholestatic (ie, due to intra- or extrahepatic bile duct blockage). Blood urea nitrogen values were increased in several animals in the terminal stage of disease,
which, in the face of more normal creatinine values, is suggestive of a prerenal insult (ie, compromised blood flow to kidney probably due to decreased circulating volume, decreased cardiac output, or renal artery disruption). The remaining biochemical markers were relatively unchanged or demonstrated no trend during infection.

**Hematological Findings**

Hematological findings (Table 1) demonstrated a transient lymphopenia, neutropenia, monocytopenia, and eosinopenia, as well as anemia, in all infected animals 4–10 days after infection. The trends in hematocrit and hemoglobin level were nearly identical, which implies that the animals were losing whole blood from bleeding as opposed to hemoconcentrating. This was followed by generalized leukocytosis on days 10–13 after infection, which was mainly due to neutrophilia. There was a superimposed trend of lymphopenia beginning around days 4–10 after infection, but this had little effect on overall white blood cell counts, as neutrophilia predominated. Platelets showed an overall downward trend during infection, although the lowest counts were rarely below the threshold of thrombocytopenia (defined as a platelet count of <150 000 platelets/µL) in most animals. The timing of the drop coincided with the coagulation disorders noted below. The trend of the mean platelet volume increased during infection, which implies that the bone marrow was able to produce new (larger) platelets, thereby confirming consumptive platelet loss as opposed to decreased production.
Coagulation Parameters

Coagulation profiles (Table 1) showed general increases in prothrombin time, activated partial thromboplastin time, and thrombin time, beginning 7–10 days after infection. Corresponding to this, an increase and precipitous fall in fibrinogen levels was observed, with the decline occurring in parallel with an increased detection of fibrinogen degradation products (ie, D-dimers). There was also a trend toward decreased protein S activity and a pronounced, transient decrease in protein C activity 7–10 days after infection. Disseminated intravascular coagulation (DIC) is often stated not to be a part of LF in humans; however, combined, our results suggest DIC involvement in the macaque model of LF. In general, the changes in coagulation profiles were slightly delayed in the Soromba-R group. Interestingly, despite similar trends in coagulation parameters, hemorrhage was exclusively noted in the 3 animals.

Figure 2. Disease progression of Lassa virus (LASV)–infected cynomolgus macaques. Groups of 3 macaques were infected with a median tissue culture infective dose of $1 \times 10^4$ LASV strain Josiah (green), Z-132 (red), or Soromba-R (blue) and scored daily for clinical signs of disease, as outlined in Materials and Methods. Animals infected with the Malian isolate (Soromba-R) demonstrated prolonged time to death (A) and decreased lethality (B), compared with animals infected with traditional LASV strains from Sierra Leone (Josiah) or Liberia (Z-132). Euthanasia occurred when animals reached a clinical score of 30 (see Materials and Methods). The bar with the dot above it indicates the clinical score for the macaque that did not progress to the point of euthanasia.
infected with LASV strain Z-132 and presented as mild to moderate multifocal petechiation of the epididymis and mucosal linings of the bladder and colon, as well as dermal petechia.

Serum Cytokine and Chemokine Analysis
Analysis of serum cytokine/chemokine levels revealed interesting differences in immune activation between the 3 groups; however, several general trends were noted in our study that are consistent with previous work. A transient spike in IFN-γ levels 7–10 days after infection and moderate increases in IL-6 and IL-12/23 levels at the time of death were noted in all groups. These patterns have previously been associated with poor prognosis in humans [26] and NHPs [18, 27]. Similarly, a decrease in IL-8 levels, which has been suggested as an indicator of fatal infection in humans [26], was noted across all groups. Increases in IL-2, IL-18, and MCP-1 levels were also observed during the course of infection in all groups. Perhaps most interesting, infection with Soromba-R resulted in the most immune activation, specifically, neutrophil-activating or -potentiating proinflammatory cytokines or growth factors, including TNF-α, MIP 1α, IL-1β, and G-CSF, as well as IL-5 (Figure 3). This is in sharp contrast to the Josiah infection group, which, consistent with previous studies, demonstrated little to no activation of MIP-1α, IL-1β, or TNF-α [26, 28]. Importantly, increased IL-1β has previously been associated with nonfatal cases of LF and may play a role in the decreased lethality and increased time to death observed in Soromba-R–infected NHPs [26].

Pathological and Histopathological Findings of LASV Infection
At necropsy, several gross pathological findings were common among the 3 LASV strains used in these studies. Mild to moderate hepatomegaly and splenomegaly were observed in all cynomolgus macaques at the time of death. All lymph nodes examined (inguinal, axillary, submandibular, mesenteric, and bronchial) were mildly to moderately enlarged and hyperplastic in all 3 groups. Radiography revealed an increased cardiothoracic size in the majority of animals, which could represent pericardial effusion, cardiomyopathy, or multichamber enlargement. However, to differentiate these would require a functional test (ie, echocardiography).

The most striking observation in these studies was the extent of lung pathology in the Soromba-R infection group (Figures 4 and 5). Radiographically, extensive pulmonary infiltrates were apparent in the 2 terminally ill NHPs infected with Soromba-R. At necropsy, pulmonary lesions were most pronounced in these animals. Specifically, the lungs were severely wet and heavy, with diffuse areas of reddening affecting 50%–100% of all lobes. In contrast, 1 of 3 animals infected with Z-132 and 2 of 3 animals infected with Josiah demonstrated multifocal areas of discoloration of varying degrees (on average affecting 25%–50% of individual lobes); however, only those from the single Z-132–infected animal were obviously edematous. Histologically,
LASV infection resulted in mild to severe subacute interstitial pneumonia characterized by multifocal to coalescing thickening of alveolar septa by edema, fibrin deposition, and varying numbers of immune cells (plasma cells, lymphocytes, etc.).

Figure 3. Serum cytokine and chemokine responses in Lassa virus (LASV)-infected cynomolgus macaques. Serum cytokine/chemokine concentrations were determined in serial serum samples collected from macaques infected with LASV isolates Josiah (green), Z-132 (red), or Soromba-R (blue), using a nonhuman primate cytokine kit (Millipore) as described by the manufacturer. Several predictors of severe Lassa fever were confirmed in the 3 infection groups, including a transient spike in interferon γ (IFN-γ) level, moderate increases in interleukin 6 (IL-6) and interleukin 12/23 (IL-12/23) levels, and a decrease in interleukin 8 (IL-8) level (A). Interestingly, LASV strain Soromba-R and, to a lesser extent, strain Z-132 resulted in increased activation of several cytokines/chemokines, compared with LASV strain Josiah, including tumor necrosis factor α (TNF-α), macrophage inflammatory protein 1α (MIP-1α), interleukin 1β (IL-1β), granulocyte colony-stimulating factor (G-CSF), and interleukin 5 (IL-5; B). Data were analyzed with 2-way analysis of variance, using a Bonferroni posttest comparing Soromba-R and Z-132 infection groups to the Josiah group. Error bars represent the standard error of the mean. *P < .05, **P < .01, and ***P < .001.
macrophages, and neutrophils) and multifocal type 2 pneumocyte hyperplasia. Consistent with the radiographic and gross pathological observations, LASV strain Soromba-R induced the most severe changes (average severity score, 4), with coalescing to diffuse interstitial pneumonia in every lung lobe (Figure 5). The pneumonia caused by LASV strains Josiah and Z-132 was considerably less diffuse, with decreased edema and fewer inflammatory cells (average severity score, 1.7 for both strains), which is in agreement with previous studies [18].

Histologically, the remaining organs analyzed showed no significant differences between the 3 infection groups and concurred with findings of other studies (Supplementary Figure 2) [18, 27]. Most animals demonstrated a mild, multifocal portal hepatitis with portal vessels bounded by small numbers of plasma cells, lymphocytes, and few macrophages and neutrophils. Several animals also had multifocal, random necrosis of hepatocytes, with loss of hepatocytes and replacement by karyorrhectic and cellular debris, along with small numbers of neutrophils, macrophages, and plasma cells. All 3 LASV strains induced meningoencephalitis in the frontal lobe, cerebellum, and brainstem, with lesions distinguished by small numbers of lymphocytes and plasma cells that multifocally expanded throughout Virchow-Robin spaces and by multifocal gliosis. Every animal had evidence of a normal immunologic response. All lymph nodes examined had follicular hyperplasia and thickening of medullary cords by abundant plasma cells. The spleen displayed similar changes, with minimal to mild follicular hyperplasia of the white pulp. Histological changes in the remaining tissues analyzed were largely unremarkable.

Results of IHC staining for LASV antigen did not vary significantly between the 3 groups. LASV antigen was detectable in endothelial cells in all tissues examined (lung, mediastinal lymph nodes, liver, spleen, kidney, and brain), as well as in hepatocytes, dendritic cells, and macrophages in spleen, bronchial epithelial cells, type II pneumocytes, lymph node follicular centers, and foci of gliosis.

Viremia and Viral Load in Tissues
LASV produced a systemic infection in macaques, with all 3 isolates replicating to similarly high titers in all organs analyzed (Figure 6A). Viremia was detected in all animals by days 7–10 after infection, with high viral titers in blood maintained until euthanasia (Figure 6B). There was no significant difference in LASV titers from the blood of the surviving animal as compared to those with lethal infections. Viremia was detected for approximately 1 month after challenge in the survivor. These findings suggest the potential for increased risk of person-to-person transmission of LASV by individuals who recover from LF.

CONCLUSIONS

The apparently low incidence of clinically significant cases of LF in Mali (and other countries between Sierra Leone, Liberia, Guinea, and Nigeria) led us to speculate that the Malian LASV isolate may be less virulent. Recently, we found that the LASV prevalence among peridomestic Mastomys natalensis collected in southern Mali was 25%–50% [7, 28], suggesting
that there is a high probability that people living in close contact with these animals are being exposed to LASV. However, even with the increased awareness of LF in Mali, only the imported case in early 2009 has been identified to date [6]. Differences in the virulence of Liberian LASV isolates have been previously demonstrated in inbred guinea pigs but not in a more stringent NHP model [20]. Here, we provide evidence in both animal models of LF that Soromba-R is less pathogenic, with decreased lethality and increased time to death in macaques, compared with LASV isolates from Sierra Leone and Liberia. In support of these findings, LASV AV, which was also isolated from outside the historical region of LASV endemicity [5] and is phylogenetically most closely related to Soromba-R [6, 7], was recently shown to be less pathogenic in cynomolgus macaques [27].

Soromba-R–infected NHPs demonstrated increased immune activation, which was possibly responsible for decreased virulence and increased pulmonary manifestations. An important virulence factor for LASV is the ability to evade the host’s immune responses, with several studies demonstrating

**Figure 5.** Histological analysis of lungs from Lassa virus (LASV)–infected cynomolgus macaques. Infection with a median tissue culture infective dose of $1 \times 10^4$ LASV isolates Josiah (A and B), Z-132 (C and D), and Soromba-R (E and F) revealed a similar degree of viral antigen staining in the pulmonary endothelium across the 3 infection groups. Despite this, the severity of pathological changes was most prominent in the lungs of Soromba-R–infected macaques, with coalescing to diffuse interstitial pneumonia and pulmonary edema observed in every lung lobe. By comparison, the pneumonia caused by Josiah and Z-132 was considerably less diffuse, with decreased edema and fewer inflammatory cells. Images are at 40× magnification with a 400× inset. Abbreviations: H&E, hematoxylin-eosin stain; IHC, immunohistochemical analysis.
that LASV is extremely effective at dampening innate responses [18, 26, 29–33]. Consistent with published reports, LASV Josiah resulted in little to no activation of host responses in macaques. In contrast, LASV Z-132 infection and, most notably, Soromba-R infection resulted in increased innate responses, compared with the Josiah infection group. Importantly, TNF-α levels were considerably increased in NHPs infected with Soromba-R, which may partially explain the increased pulmonary manifestations noted in these animals, since TNF-α is known to increase the permeability of endothelial cells [34]. A key finding in this study was the atypical presentation of LF associated with Soromba-R infection. While the hallmarks of LF were present in these animals, we also found that NHPs infected with Soromba-R presented with increased pulmonary manifestations. On the basis of these observations, it seems possible that LF patients seeking medical attention in Mali may receive a misdiagnosis of pulmonary syndrome. Although the degree of respiratory involvement in the Josiah and Z-132 infection groups was low in our study, it is also possible that pulmonary manifestations

Figure 6. Viremia and organ titers from Lassa virus (LASV)–infected cynomolgus macaques. Macaques infected with a median tissue culture infectious dose (TCID₅₀) of 1 x 10⁴ LASV isolates Josiah (green; n = 3), Z-132 (red; n = 3), or Soromba-R (blue; n = 2) were euthanized as indicated by individual clinical scores, underwent necropsy, and had tissues titered by standard methods. A, LASV titers in solid organs did not differ across the 3 infection groups. B, LASV Josiah demonstrated slightly increased levels of viremia when compared to Z-132 or Soromba-R–infected nonhuman primates. The single survivor in the Soromba-R group did not have significantly lower levels of viremia (data not shown). Error bars represent the standard error of the mean. Shown are results for the right lung.
associated with LASV infection in general are underrecognized throughout West Africa.

Genetically, LASV isolates have a large degree of diversity, which may result in differences in pathogenicity and disease manifestations [20, 35]. Combined, the findings of differences in clinical manifestations and immune suppression/activation observed here support this hypothesis and demonstrate the importance of studying geographically distinct LASV isolates in animal models of disease to gain a better understanding of LASV pathogenesis and to evaluate potential medical countermeasures against LF. Despite the high incidence of LF in Sierra Leone, Liberia, Guinea, and Nigeria, as well as the increasing frequency of imported cases of LF worldwide, currently there are no antivirals or vaccines licensed for treating or preventing LASV infection or LF [4, 36]. Ribavirin has been proven effective and is frequently used in an off-label manner to treat LF, although its efficacy is reliant on early administration [4, 13, 37, 38]. To date, the few therapeutic modalities tested in animal models have mainly been evaluated against the commonly used LASV Josiah [13, 17, 39–41]. The results of our study provide an important point of reference for evaluating therapeutic agents against these other strains of LASV.

**Supplementary Data**

Supplementary materials are available at The Journal of Infectious Diseases online (http://jid.oxfordjournals.org/). Supplementary materials consist of data provided by the author that are published to benefit the reader. The posted materials are not copyedited. The contents of all supplementary data are the sole responsibility of the authors. Questions or messages regarding errors should be addressed to the author.

**Notes**

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