Identification and pathological characterization of persistent asymptomatic Ebola virus infection in rhesus monkeys


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Abstract

Ebola virus (EBOV) persistence in asymptomatic humans and Ebola virus disease (EVD) sequelae have emerged as significant public health concerns since the 2013–2016 EVD outbreak in Western Africa. Until now, studying how EBOV disseminates into and persists in immune-privileged sites was impossible due to the absence of a suitable animal model. Here we detect persistent EBOV replication with possibly coinciding systematic inflammatory responses in otherwise asymptomatic rhesus monkeys that had survived infection in the absence of or after treatment with candidate medical countermeasures. We document progressive EBOV dissemination into eyes, brain, and
tests through vascular structures similar to observations in humans. We identify CD68+ cells (macrophages/monocytes) as the cryptic EBOV reservoir cells in the vitreous humor and its immediately adjacent tissue, in the tubular lumina of the epididymides, and in foci of apparently histiocytic inflammation in the brain but not in organs typically affected during acute infection. In conclusion, our data suggest that human persistent EBOV infection could be modelled in rhesus monkeys, and demonstrate that promising candidate medical countermeasures may not completely clear EBOV infection. A rhesus monkey model may lay the foundation to study EVD sequelae and to develop therapies to abolish EBOV persistence.

Introduction

Ebola virus disease (EVD) is a severe human viral hemorrhagic fever caused by infection with either Bundibugyo virus (BDBV), Ebola virus (EBOV), Sudan virus (SUDV), or Taï Forest virus (TAFV).

From 2013 to 2016, an EBOV-caused EVD outbreak in Western Africa led to a total of 28,646 cases and 11,323 deaths worldwide (case-fatality rate of 40%)\(^1\). The large number of EVD survivors revealed previously underestimated sequelae of EBOV infection or “post-Ebola (virus disease) syndrome.” This syndrome includes arthralgia, cognitive impairment, headaches, hearing loss, and myalgia\(^2\)\(^-\)\(^5\). About 50% of surveyed survivors of this outbreak complained of various ocular complications such as blurry vision, pain, and discomfort. One fifth of these survivors were diagnosed with uveitis\(^3\)\(^-\)\(^12\). In one uveitis case, EBOV was isolated from the eye\(^11\). Another EVD survivor developed
meningoencephalitis more than 9 months after convalescence, and EBOV could be isolated from the patient’s cerebrospinal fluid.

EBOV RNA was detected in seminal fluid of survivors up to 18 months after the onset of symptoms, and in one case, EBOV was successfully isolated from seminal fluid at least several weeks after convalescence. Most importantly, male-to-female sexual transmission from two persistently EBOV-infected EVD survivors, 6 and 15 months after convalescence, respectively, was confirmed by genetic sequence comparison of EBOV in the blood of virus donors and recipients. One of these transmissions became the epidemiological basis for a new cluster of EVD cases. Several other EVD transmission chains are likely to have originated from the reintroduction of EBOV from a persistently infected source.

Together, these reports indicate the need for a high priority effort to understand EBOV persistence and relapse in vivo. Persistent EBOV might not only threaten the long-term survival of immediate EVD survivors, but also can be transmitted and start novel flare-ups. However, until now, no animal model for EBOV persistence and relapse has been developed, and the molecular pathology of persistent EBOV infection in immune-privileged sites has not been examined. Here we report for the first time persistent EBOV infection in eyes, epididymides, and brain tissues of experimentally EBOV-infected rhesus monkey survivors. We illustrate the anatomical location and cellular targets of EBOV in these tissues. We demonstrate that EBOV spreads into specific locations at immune-privileged sites through vascular structures. In addition, we provide evidence that EBOV persistence is associated with ongoing replication in the likely presence of an inflammatory host response.
Results

Ebola virus persists in rhesus monkey survivors. Nonhuman primates (NHPs), in particular macaques, are broadly used as experimental animal models of EVD\textsuperscript{19}. Yet, to our knowledge, persistent EBOV infection has not been reported in NHP survivors of experimental infection. Using \textit{in situ} hybridization (ISH), we screened a collection of archived tissue (eye, testicle, brain, lymph node, liver, spleen) samples from 112 rhesus monkeys that had survived experimental EBOV infection through day 43 post-exposure. Only a portion of survivors, 11 of 112 (9.82%), had detectable EBOV genomic RNA in eye, testicle, or brain tissues. Specifically, 9 of 78 tested eyes (11.54%), 1 of 76 testes (1.32%) and 1 of 80 brains (1.25%) contained EBOV genomic RNA. In contrast, EBOV genomic RNA was undetectable in the common acute EBOV infection target tissues (liver, lymph node, and spleen). Next to rhesus monkeys, crab-eating macaques are frequently used for experimental EBOV infections\textsuperscript{19}. We therefore also screened a collection of archived tissues samples from 48 crab-eating macaques that had survived EBOV infection. Importantly, the rhesus monkey tissues were derived from animals that either received non-vaccine MCMs or were untreated, whereas the only tissues from crab-eating macaques available for this study originated from animals that had received candidate vaccines. There was no evidence of EBOV persistence in the examined crab-eating macaque eye, testis, and brain tissues using ISH (data not shown).

Tissues from 8 of the 11 rhesus monkey survivors that tested positive for EBOV both by ISH and immunofluorescence were more closely examined (Supplementary Table 1). Of note, 2 of the 8 (25%) natural survivors that had not been treated with...
Ebola virus persistently infects the vitreous humor and its adjacent structures.

Genomic EBOV RNA was detected in the ocular tissue (vitreous humor and its adjacent structures were predominant) of 6 of 8 rhesus monkeys by ISH using EBOV genomic nucleoprotein (NP) gene-specific probes (Fig. 1). In contrast, genomic EBOV RNA could not be detected in brains, livers, lymph nodes, ovaries/uteri, spleens, or prostates/testicles from these animals (Supplementary Table 1).

Ebola virus spreads into sites of persistence through blood vessels in the eye.

Intramuscular infection of rhesus monkeys with 1,000 pfu of EBOV generally results in 100% fatality with death occurring 5–11 days after exposure\textsuperscript{20-22}, here called “acute course of disease death” (ACDD). Previous studies revealed the presence of EBOV antigen in the scleras, corneal epithelia, retinas, and optic nerve leptomeninges of EBOV-infected rhesus monkeys that perished prior to the study endpoint but survived longer than ACDD (i.e., 16–24 days post-exposure, here referred to as “delayed time of death [DTOD]”)\textsuperscript{21,23}. We further analyzed sections of eyes from 24 rhesus monkeys with ACDD (Supplementary Table 2) and 14 rhesus monkeys with DTOD using ISH (Supplementary Table 3). Strikingly, in animals with ACDD, EBOV genomic RNA was not detected in the parenchymal eye tissues (Fig. 1f), but was consistently detected in blood vessels of the choroids (17 of 24, 70.83 %; Supplementary Fig. 1a), ciliary processes (18 of 24, 75%; Supplementary Fig. 1b) and/or in the optic nerve.
leptomeninges (6 of 24, 25%; Supplementary Fig. 1c). Consistently, EBOV glycoprotein (GP)₁,₂ antigen was specifically detected within CD31 antibody-labeled blood vessels by immunofluorescence staining (Supplementary Fig. 1d-d'').

Among animals with DTOD (Fig. 1g), EBOV genomic RNA was predominantly detected in the optic nerve leptomeninges (12 of 14, 85.71%; Supplementary Fig. 1e), scleral connective tissues (7 of 14, 50%; Supplementary Fig. 1f), cilias and irises (3 of 14, 21.43%; Supplementary Fig. 1g), and choroids (2 of 14, 14.29%; Supplementary Fig. 1h). Interestingly, in 1 infected animal with DTOD, EBOV genomic RNA was present in cells in the vitreous humor in addition to optic nerve leptomeninges, and scleras (1 of 14, 7.14%; Supplementary Fig. 1i-i').

Persistent EBOV infection in survivors was confined to the inner vitreous humor and immediately adjacent structures (Fig. 1h). These data suggest the possibility that ocular EBOV infection originates from EBOV in blood vessels of the choroid and cilias, or optic nerve leptomeninges with subsequent spread to adjacent structures. The infection of outer scleral and uveal layers and optic nerve leptomeninges may be cleared after recovery while EBOV persists in the inner vitreous humor and its adjacent structures.

**CD68⁺ cells are cellular targets of Ebola virus persistence in the eye.** We examined survivor eye sections using immunofluorescence staining with anti-EBOV GP₁,₂ antibodies to determine whether EBOV RNA-positive tissues actually contain EBOV GP₁,₂ antigen. EBOV GP₁,₂ antigen was detectable in the same group of infected cells from survivors that tested positive by ISH (Fig. 2a-b). Dendritic cells (DCs) and macrophages are believed to be primary targets of acute EBOV infection in primates.²⁴,²⁵
Hematoxylin and eosin (H&E) staining suggested numerous inflammatory cells, including cells that had the morphologic appearance of macrophages, in the vitreous humor and the retinal inner limiting membranes of survivors (Fig. 2c-d). To identify the cellular targets of persistent EBOV infection, we stained survivor eye sections by immunofluorescence using anti-EBOV GP$_{1,2}$ antibody and antibody against a macrophage/monocyte marker, CD68. Most cells that tested positive for EBOV GP$_{1,2}$ antigen were also labeled by the anti-CD68 antibody (Fig. 2e–h’).

**Uveitis, retinitis, and vitritis are accompanied by reactive gliosis with Ebola virus persistence.** Six of 8 survivors indeed revealed moderate-to-severe uveitis. Infiltration of cells, including macrophages, was found within the ciliary bodies, irises, and scleras near the corneoirideal angles (Fig. 3a-b, Supplementary Fig. 2a–d). Infiltration occasionally extended along cyclitic membranes. CD68 staining suggested that some of the uvea-infiltrating cells were macrophages or monocytes (Supplementary Fig. 2e).

Retinitis occurred in 4 survivors, typically characterized by perivascular accumulation of likely inflammatory cells (Fig. 3c–d, Supplementary Fig. 2g–h). CD68$^+$ cell infiltration of the retinas was commonly observed (Fig. 3e–f). Infiltrating cells, including CD68$^+$ cells, and/or neutrophils, were also found in the vitreous humor adjacent to the cillas and retinas of the 4 survivors (vitritis, Fig. 3b-c, g-h, Supplementary Fig. 2f–h).

Using an antibody against glial fibrillary acidic protein (GFAP) to outline retinal structures of the six survivors, we unexpectedly found dramatically upregulated GFAP immunoreactivity in Müller cell processes that extended into the inner and outer nuclear...
layers (Fig. 3j–l, Supplementary Fig. 2i–j). In contrast, very limited GFAP immunoreactivity was observed in the ganglion cell layer of a normal eye of an uninfected control monkey (Fig. 3i). Both upregulation of GFAP and cell proliferation are hallmarks of reactive gliosis subsequent to acute retinal injury or chronic neuronal stress. Some retinal cells were labeled with the cell proliferation marker Ki67 (Fig. 3m–p), suggesting that cell proliferation was activated to maintain retinal homeostasis during EBOV persistence.

**Ebola virus persistence in the brain of a rhesus monkey survivor.** Persistence and relapse of EBOV in the central nervous system in an EVD survivor have been reported. However, how EBOV is able to bypass the blood-brain barrier to access immune-privileged sites and which cells are being targeted by EBOV to establish persistence in the brain remain unknown. We identified 1 of 8 rhesus monkey survivors that survived to the end of study (28–43 days) on day 36 and tested positive for EBOV RNA in the corpus striatum of the brain (Supplementary Table 1). EBOV RNA was specifically detected in foci of histiocytic encephalitis (glial nodules) in which we detected activated microglia (Fig. 4a and Supplementary Fig. 3a). Additional evidence of encephalitis included intense perivascular infiltration of mononuclear cells that filled the Virchow-Robbins space in the corpus striatum of the brain (Supplementary Fig. 3b, c).

Immunofluorescence staining revealed most of the cells in the glial nodule to be CD68+ cells, some of which were associated with EBOV GP1,2 antigen (Fig. 4b–b′′′). Interestingly, EBOV genomic RNA was absent from parenchymal brain tissues but was consistently detected in endothelial cells of blood vessels of animals with ACDD (19 of
24, 79.16%; Fig. 4c, Supplementary Table 2). However, in animals with DTOD, EBOV genomic RNA was mainly found in foci of histiocytic encephalitis most often associated with perivascular glial nodules (14 of 15, 93.33%; Fig. 4d, (Supplementary Table 3). EBOV GP\textsubscript{1,2} antigen was specifically located in CD31 antibody-labeled vascular (Fig. 4e–e’) endothelial cells of the blood brain barrier in animals with ACDD. Additionally, some of the blood endothelial cells infected by EBOV e underwent apoptosis (Fig. 4f). In contrast, GP\textsubscript{1,2} antigen was mainly detected in CD68 antibody-labeled microglial clusters, which were predominantly observed in the vicinity of CD31 antibody-labeled blood vessels in the brains of animals with DTOD (Fig. 4g–i). Our observations suggest that initial EBOV infection of the vascular endothelium in the brain may lead to breakdown of the blood-brain barrier and may result in subsequent histiocytic inflammation and perivascular EBOV infection in microglia. EBOV may replicate and persist in these inflammatory microglial cells after recovery (Fig. 4j).

**Ebola virus persistence in the epididymis of a rhesus monkey survivor.** We identified 1 of 8 rhesus monkey survivors, euthanized on day 43, with EBOV persistence in testicular tissue using ISH (Supplemental Table 1 and Fig. 5a-a’). Strikingly, EBOV RNA was only found in the tubular lumen of one epididymis but not in the testes. The tubular lumen of this immature animal was congested by cells, including CD68\textsuperscript{+} cells, and necrotic cellular debris, composed of degenerate neutrophils, macrophages, and spermatogonia (Fig. 5b). EBOV GP\textsubscript{1,2} antigen was detected in some CD68\textsuperscript{+} cells or extracellularly within the tubular lumen of the epididymis (Fig. 5c–c’’). In addition,
inflammatory cells, including macrophages, eosinophils, and lymphocytes, were found in interstitial connective tissue (Fig. 5b), suggesting epididymitis.

To further evaluate EBOV persistence in the testes and epididymides, we screened testicular tissues from 24 rhesus monkeys with ACDD (Supplementary Table 2) and 8 rhesus monkeys with DTOD (Supplementary Table 3). Twenty two of 24 (91.67%) animals with ACDD had detectable EBOV genomic RNA mostly in the vascular structure including in endothelial cells (Fig. 5d–d’) in both testes and epididymides.

Immunofluorescence staining illustrated that EBOV was detected mainly in vascular structures including blood vessels labeled by CD31 antibody and CD31-negative lymphatic vessels (Fig. 5e–e’). In contrast, 5 of 8 (62.5%) animals with DTOD had broad detectable EBOV genomic RNA in the interstitia of testes and/or epididymides (Fig. 5f–f’). Consistently, immunofluorescence showed wide EBOV antigen distribution in interstitia (Fig. 5g). EBOV genomic RNA was also found in the seminiferous tubules in 2 of those 5 animals with DTOD but not in any of the animals with ACDD (Fig. 5h–h’).

EBOV GP1,2 antigen was detected in germ cells in seminiferous tubules labeled by DDX4/Vasa antibody (Fig. 5i–i”). Consistently, it has been reported that EBOV was detected in seminiferous tubules in addition to interstitial infection in a fatal human case of Ebola virus infection.

Ebola virus replicates in immune-privileged sites. Our study was based on archived fixed tissues and live titration of EBOV could therefore not be performed. Thus, to determine whether EBOV actively replicates in immune-privileged sites associated with persistence, we developed a multiplex fluorescence in situ hybridization (FISH) assay.
that differentiates between genomic and antigenomic EBOV RNA in tissues.

Surprisingly, we detected not only the EBOV genome, but also the EBOV antigenome, in the eye, epididymis, and brain tissues of survivors, suggesting ongoing EBOV replication at the time of euthanasia (Fig. 6a–f′′′). Interestingly, EBOV replicated at a lower level in the eyes of two survivors compared to that observed in two others (Fig. 6a–d′′′ and g). This higher level of viral replication coincided with more severe inflammation-like responses in the eyes.

**Discussion**

EBOV persistence and post-EVD syndrome have become significant public health concerns amid reports of sexual EBOV transmission and EVD outbreak flare-ups originating from persistently infected survivors since the 2013–2016 EVD outbreak in Western Africa. Our data suggest that EBOV may progressively disseminate into eyes, brain, and testes through vascular structures, and likely persists in CD68⁺ (macrophage/monocyte) cells in rhesus monkey survivors of EBOV infection. We observed a likely inflammatory host immune response in immune-privileged tissues possibly due to active viral replication. In NHPs, acute EBOV infection primarily affects areas connected directly or adjacent to blood vessels with replication mainly occurring in the liver, kidney, lung, and lymphatic organs with possible later transition to infection of blood vessel-adjacent structures in the brain, eyes, and testes. After that, EBOV eventually spreads to sites of persistence from which the virus cannot be cleared. The animals here called survivors would still have been at risk of “relapse.” However, as
rhesus monkey survivors of EBOV infection typically are not maintained beyond ≈40 days after initial EBOV exposure, the possibility of relapse could not be assessed. The large and unprecedented 2013–2016 EVD outbreak in Western Africa highlighted diverse clinical sequelae of EVD in thousands of survivors⁴⁻⁵. While the molecular etiology of these sequelae is unclear, EBOV persistence accompanied by local inflammatory reactions in apparently healthy survivors could at least partially explain the persistence of pain at various body sites and/or visual, acoustic, and cognitive impairments. Indeed, EBOV persistence in the eyes, testicles, and brains of a few human survivors, substantiated by virus isolation⁸⁻¹¹,¹²,²⁹, possibly caused frequently reported EVD sequelae such as vision loss, orchitis, and confusion, respectively. In at least once case, EBOV persistence led to repeated disease recurrence in a survivor several weeks after clinical recovery⁸, suggesting that survivors should not necessarily be considered completely cleared of the virus. Future studies aimed at understanding the molecular basis of EBOV persistence could include genomic characterization of EBOV in infected cells at immune-privileged sites compared to actively circulating EBOV in acute disease. Genomic sequencing of tissue-specific EBOV populations may help explain the extraordinarily low mutation rate of EBOV associated with persistence in humans¹⁶,³⁰. Several instances of new EVD case clusters were linked directly to persistently EBOV-infected, apparently healthy survivors of the 2013–2016 EVD outbreak¹³⁻¹⁷. Although rare, such transmissions could prove to be catastrophic. The majority of EVD outbreaks could be traced back to single virus introductions from a natural reservoir into the human population, with subsequent person-to-person transmission³¹. In the 2013–2016 EVD outbreak, specifically, a single EBOV introduction ultimately led to 28,646
human infections\textsuperscript{16,32}. Understanding the molecular-virological mechanism of EBOV persistence is of paramount importance, including the conditions that favor persistence and reactivation and the time frame in which they may occur. Ultimately, the goal must be the development of MCMs that prevent or abolish EBOV persistence in survivors.

Currently, no established animal models of persistent EBOV infection are available. All frequently used animal models for EVD typically result in 100\% fatality\textsuperscript{33}. Our study clarifies that a robust rhesus monkey model for EBOV persistence could be developed based on partial clearance of EBOV from experimentally infected animals and used to evaluate various candidate MCMs. Additionally, two natural rhesus monkey survivors with persistent EBOV reported here had not been treated with MCMs, which suggests that a robust rhesus monkey model of EBOV persistence might be developed by increasing the survival rate of infected macaques. We found persistent EBOV in three key immune-privileged sites (i.e., eyes, testicles, brains) only in rhesus monkeys, but not in crab-eating monkeys. However, all the crab-eating monkeys examined in this study had been exposed to EBOV after receiving candidate vaccinations, whereas all the examined rhesus monkey survivors had been treated with experimental MCMs post-EBOV exposure or had not been treated at all. Vaccine-related EBOV suppression possibly prevented EBOV dissemination to and replication in immune-privileged sites. Consequently, our findings in rhesus monkeys and crab-eating macaques ought not to be compared directly. Instead, follow-up studies should investigate whether crab-eating macaques that survived EBOV infection after treatment with non-vaccine MCMs also develop persistent EBOV infections. In addition, follow-up studies ought to clarify

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whether our observations regarding EBOV persistence in brain and testis tissues, both based on single animals, can be generalized.

Methods

Tissues. Formalin-fixed paraffin-embedded (FFPE) brain, eye, testis, and epididymis tissues of rhesus monkeys (*Macaca mulatta*) or crab-eating (aka cynomolgus) monkeys (*Macaca fascicularis*) that had been experimentally infected with Ebola virus/H.sapiens-tc/COD/1995/Kikwit-9510621 (EBOV) were retrieved from the US Army Medical Research Institute of Infectious Diseases (USAMRIID) Pathology Division tissue archives. For the purpose of this manuscript, “survivor” refers to monkeys that survived EBOV infection after exposure until study endpoints (28–43 days after EBOV exposure) with or without treatment with distinct MCMs, including cocktails of monoclonal antibodies (mAbs), small interfering RNAs (siRNAs), antisense phosphorodiamidate morpholino oligomers (PMOs), or vaccination with Venezuelan equine encephalitis virus replicon particles expressing EBOV GP$_{1,2}$ (VRP-GP). For the purposes of comparing the type of cells and anatomical sites infected during acute and persistent infection, we also collected the same tissues from rhesus monkeys that did not survive EBOV exposure. For analysis, we further divided this group into animals that died within the typical post-exposure time frame [5–11 days post-exposure, here referred as “acute course of disease death (ACDD)” in this study] and animals that perished prior to the study endpoint but survived longer than the typical post-exposure time frame [usually 16–24 days post-exposure, here referred as “delayed time of death (DTOD)” in this study]. Sections were
stained with hematoxylin and eosin (H&E) after deparaffinization. Sections of different tissues from uninfected rhesus monkeys were used as controls.

RNA in situ hybridization. EBOV has a linear, single-stranded, negative-sense non-segmented RNA genome (≈19 kb) that encodes seven structural proteins\(^{35}\). EBOV genome replication involves synthesis of the (positive-sense) RNA antigenome, which in turn serves as a template to generate progeny (negative-sense) genomes; and synthesis of (positive-sense) mRNAs from individual genes that are translated into structural proteins\(^{36}\). To detect EBOV RNA in FFPE tissues, *in situ* hybridization (ISH) was performed using the RNAscope® 2.5 HD RED kit\(^{37}\) (Advanced Cell Diagnostics, Newark, CA) according to the manufacturer’s instructions. Briefly, 20 ZZ probe pairs targeting the genomic EBOV *NP* gene were designed and synthesized by Advanced Cell Diagnostics (Cat# 448581). After deparaffinization with xylene, a series of ethanol washes, and peroxidase blocking, sections were heated in antigen retrieval buffer and then digested by proteinase. Sections were exposed to ISH target probe and incubated at 40°C in a hybridization oven for 2 h. After rinsing, ISH signal was amplified using company-provided Pre-amplifier and Amplifier conjugated to horseradish peroxidase (HRP), and incubated with a red substrate-chromogen solution for 10 min at room temperature. Sections were then stained with hematoxylin, air-dried, and mounted.

Multiplex fluorescence in situ hybridization. Multiplex fluorescence *in situ* hybridization (FISH) was performed using the RNAscope® Fluorescent Multiplex Kit (Advanced Cell Diagnostics) according to the manufacturer’s instructions with minor
modifications. Twenty ZZ probe pairs with C1 channel (red, Cat# 448581) targeting EBOV genomic NP gene and 20 ZZ probe pairs with C3 Channel (Cat# 451691), green) targeting the EBOV antigenomic VP35 gene were synthesized by Advanced Cell Diagnostics. FFPE-tissue sections underwent deparaffinization with xylene and a series of ethanol washes. These tissue sections were treated with 0.1% Sudan Black B (Sigma-Aldrich, St. Louis, MO, USA) to reduce autofluorescence, heated in antigen retrieval buffer, and digested by proteinase. Sections were exposed to ISH target probes and incubated at 40°C in a hybridization oven for 2 h. After rinsing, ISH signal was amplified using company-provided Pre-amplifier and Amplifier conjugated to fluorescent dye. Sections were counterstained with 4',6-diamidino-2-phenylindole (DAPI, Thermo Fisher Scientific, Waltham, MA, USA), mounted, and stored at 4°C until image analysis. FISH images were captured on an LSM 780 Confocal Microscope (Zeiss, Oberkochen, Germany) and processed using open-source ImageJ software (National Institutes of Health, Bethesda, MD, USA).

**Immunofluorescence staining.** After deparaffinization and treatment with 0.1% Sudan Black B to reduce autofluorescence, tissues were heated in citrate buffer (Sigma-Aldrich), pH 6.0, for 15 min to reverse formaldehyde cross-links. After rinses with phosphate-buffered saline, pH 7.4 (PBS, Thermo Fisher Scientific), sections were blocked overnight with PBS containing 5% normal goat serum (Sigma-Aldrich) at 4°C. Sections were then incubated with primary antibodies (USAMRIID rabbit polyclonal antibody against EBOV GP\(_{1,2}\) at a dilution of 1:2,000; mouse anti-human CD68 antibody at a dilution of 1:200 [Clone KP1, Dako Agilent Pathology Solutions, Carpinteria, CA],
USA]; rabbit polyclonal antibody against glial fibrillary acidic protein [GFAP] at a
dilution of 1:100 [ab7260, Abcam; Cambridge, MA, USA]; rabbit polyclonal antibody
against Ki67 [ab15580, Abcam] at a dilution of 1:400; rabbit polyclonal antibody against
DDX4/Vasa [ab13840, Abcam] at a dilution of 1:100; rabbit polyclonal antibody against
CD31 at a dilution of 1:100 [ab28364, Abcam; Cambridge, MA, USA]; rabbit
monoclonal antibody against Cleaved Caspase-3 at a dilution of 1:100 [9664, Cell
Signaling; Danvers, MA, USA]; USAMRIID mouse IgG2A anti-EBOV GP
monoclonal 6D8-1-2\textsuperscript{38} at a dilution of 1:400) for 2 h at room temperature. After rinsing in
PBS, sections were incubated with secondary goat IgG Alexa Fluor 488-conjugated anti-
rabbit or anti-mouse and goat IgG Alexa Fluor 561-conjugated anti-mouse or anti-rabbit
antibodies (Life Technologies, Carlsbad, CA) for 1 h at room temperature. Sections were
cover-slipped using VECTASHIELD Antifade Mounting Medium with DAPI (Vector
Laboratories, Burlingame, CA, USA). Images were captured and processed as describe
above.
References


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All USAMRIID studies described in this manuscript involved protocols approved by the Institute Animal Care and Use Committee and were conducted under an IACUC approved protocol in compliance with the Animal Welfare Act, PHS Policy, and other Federal statutes and regulations relating to animals and experiments involving animals. The facility where this research was conducted is accredited by the Association for Assessment and Accreditation of Laboratory Animal Care, International and adheres to principles stated in the Guide for the Care and Use of Laboratory Animals, National Research Council, 2011.

**Author contributions**

X.Z. conceived and designed the experiments. X.Z. and C.B.D performed the in situ hybridization. X.Z. performed the immunofluorescence staining and confocal imaging. K.A.K., C.W.S., J.J.B., S.P.H., and T.B.C. performed histopathology. T.K.W., J.W.F., K.A.C., J.M.D., and S.B. provided experimental materials. X.Z., S.R.R., G.P., and J.H.K. interpreted the data and wrote the manuscript. S.B. and M.G.S coordinated and oversaw the study.

**Additional Information**

Supplementary information is available for this paper. Reprints and permissions information is available at www.nature.com/reprints. Correspondence and requests for materials should be addressed to X.Z.
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523 Competing interests

524 The authors declare no competing financial interests.
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Figure 1 | Detection of genomic EBOV RNA in eyes of rhesus monkey survivors by *in situ* hybridization. Genomic EBOV RNA (red) detected by *in situ* hybridization (ISH) using an EBOV *NP* gene-specific probe in hematoxylin-stained (blue) FFPE sections. Single prime figures a′–e′ are insets of a–e at high magnification. a–a′. Representative images demonstrating genomic EBOV RNA in cells within vitreous humor. a″, Schematic of EBOV-infected regions (red stars) including cells in the vitreous humor and cells attached to the retinal inner limiting membrane. b–b′, Genomic EBOV RNA detected by ISH in a cluster of cells in vitreous humor. b″, Schematic of EBOV-infected regions (red stars) in vitreous humor. c–c′, Representative images demonstrating EBOV RNA in cells at the interface between vitreous humor and ciliary bodies. c″, Schematic of EBOV-infected regions (red stars) in vitreous humor and at the interface between vitreous humor and its adjacent structures, including ciliary body and retina. d–d′, Representative image demonstrating EBOV RNA in cells of the retinal inner limiting membrane adjacent to vitreous humor. d″, Schematic of EBOV-infected regions (red stars) in vitreous humor and the retinal inner limiting membrane. e–e′, Representative image demonstrating genomic EBOV RNA in cells at the retinal inner limiting membrane. e″, Schematic of EBOV-infected regions (red dots) in the retinal inner limiting membrane. f–h, Compare ocular EBOV infection pattern of survivors h to animals with ACDD f and animals with DTOD g.
Figure 2 | Ocular macrophages of survivors express EBOV GP1,2 antigen. 

a, EBOV GP1,2 antigen (red) staining in a cluster of likely inflammatory cells, including macrophages adjacent to the inner limiting membrane of the retina (labeled using a glial fibrillary acidic protein (GFAP) antibody, green). Blue 4′,6-diamidino-2-phenylindole (DAPI) stain identifies nuclei. 

b, EBOV GP1,2 antigen staining (red) in roughly the same location detected by immunofluorescence. 

c, Likely inflammatory cells (e.g., macrophages) and necrotic debris in vitreous humor adjacent to the retinal inner limiting membrane (H&E staining). 

d, Likely inflammatory cells, including macrophages, in vitreous humor adjacent to the inner limiting membrane. 

e–e‴, Immunofluorescence staining of a cross section of the retina showing EBOV GP1,2 antigen (green) in CD68⁺ cells (monocytes/macrophages, red) attached to the retinal inner limiting membrane (ILM). ONL, outer nuclear layer; INL, inner nuclear layer; GCL, ganglion cell layer. 

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560 EBOV GP$_{1,2}$ antigen (green) in CD68$^+$ cells (red, monocytes/macrophages) in the vitreous humor. 
561 g. EBOV GP$_{1,2}$ antigen (green) in CD68$^+$ cells (red, monocytes/macrophages) in the vitreous humor. (H–H’) EBOV GP$_{1,2}$ antigen (green) in CD68$^+$ macrophages (red) in the vitreous humor. Blue, nuclear stain by DAPI.
Figure 3

Figure 3 | Persistent EBOV infection causes uveitis, retinitis, and vitritis

accompanied by reactive gliosis. a–b, Uveitis (H&E). Likely inflammatory cells (red arrows), primarily macrophages, in the ciliary body and iris a and ciliary body and vitreous humor b, red arrowhead. c, Perivascular cuffs (red arrows) in the ganglion cell layer of the retina and likely inflammatory cells (red arrowheads) on the vitreous side of the inner limiting membrane. d, Perivascular inflammation-like response (red arrow) in the retina. e–f, CD68⁺ cell (red) infiltration in the ganglion cell layer of GFAP-labeled retina (green). g–h, Excessive CD68⁺ cells (red) in the anterior vitreous humor. i, In a
healthy retina, GFAP staining (green) is limited to the ganglion cell layer (GCL) and
absent from the inner nuclear layer (INL) and outer nuclear layer (ONL). j–l, GFAP
staining (green) is dramatically increased in Müller cell apical processes that extend into
the INL and ONL. m–o, Assessment of cell proliferation by Ki67 detection (green) in a
healthy retina m and in retina of survivors n-o. p, Quantification of Ki67+ cells in normal
eye and eyes of survivors. Data are represented as mean ± standard error of the mean,
n=9, 5, 5.
Figure 4
Figure 4 | Ebola virus persistence in the brain of a rhesus monkey survivor. a, EBOV genomic RNA detection by ISH in a glial nodule in the corpus striatum of brain (the inset is the image with high magnification). b–b'', Immunofluorescence staining reveals EBOV GP$_{1,2}$ antigen (green) in CD68$^+$ microglia (red) or extracellularly within a glial nodule. Blue, nuclear stain by 4',6-diamidino-2-phenylindole (DAPI). c, Vascular EBOV genomic RNA in the brain of animals with acute course of disease death (ACDD). d, EBOV genomic RNA in the glial nodule in the brain of animals with delayed time of death (DTOD). e–e', Immunofluorescence demonstrated EBOV GP$_{1,2}$ (green) in CD31 antibody (red) labeled vascular endothelial cells in the brain of animals with ACDD. f, EBOV (green) infected endothelia cell was cleaved caspase-3 positive suggestive of ongoing apoptotic cell death. g–i, Immunofluorescence staining of three serial brain sections of animals with DTOD demonstrated that CD68 antibody-labeled (green in f) microglia, in which EBOV GP$_{1,2}$ (green in g-h) was detected, aggregates in the vicinity of CD31 antibody-labeled (red in f-g) blood vessel. j, EBOV infection pattern in the brain of animals with ACDD, animals with DTOD, and survivors.
Figure 5 | EBOV persistently infects the epididymis of a rhesus monkey survivor. 

**a** and magnified **a’**, EBOV genomic RNA detection by ISH in the tubular lumen of the epididymis (red arrows). **b**, Epididymitis in the same location. Likely inflammatory cells, including macrophages, eosinophils, and lymphocytes, are located in the interstitial connective tissue (red arrowhead). The tubular lumen (red arrows) is congested with likely inflammatory cells and necrotic cellular debris composed of degenerate neutrophils, macrophages, spermatogonia, and sloughed epithelium. **c–c’**, Immunofluorescence staining reveals EBOV GP_{1,2} antigen (green) in CD68\(^{+}\) cells (macrophage/monocyte, red) or extracellularly in the tubular lumen of the epididymis. **d–**
d', Genomic EBOV RNA in the vascular structure in interstitial testicular tissue d and connective tissue surrounding the epididymal ducts d' of animals with acute course of disease death (ACDD). e–e', EBOV GP$_{1,2}$ antigen (green) in vascular structure including CD31-positive blood vessel (red, arrowhead) and CD31-negative lymphatic vessels (arrow). f–f', EBOV genomic RNA in most of interstitial tissues of testis f and in connective tissue surrounding the epididymal ducts f' in animals with delayed time of death (DTOD). g–g', EBOV genomic RNA in seminiferous tubules g and in cells surrounding epididymal ducts g' in animals with DTOD. h, EBOV GP$_{1,2}$ antigen (green) in interstitial tissue of testis in animals with DTOD. i–i'', EBOV GP$_{1,2}$ antigen in seminiferous tubules, in which germ cells were labeled by DDX4/Vasa antibody.
Figure 6 | EBOV replicates in the eyes, epididymides, and brains of rhesus monkey survivors. Multiplex fluorescence in situ hybridization (FISH) was used to detect EBOV genome (red) and antigenome/mRNA (green) in eyes a–d'', tubular lumen of the epididymis e–e'', and brain f–f''. Blue, nuclear stain by 4',6-diamidino-2-phenylindole.
g. Quantification of the FISH intensity ratio between EBOV antigenome/mRNA and genome signals.
Supplementary Tables and Figures

Supplementary Table 1. *In situ* hybridization (ISH) results obtained from eight rhesus monkey survivors of Ebola virus (EBOV) infection

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<th>Survivor</th>
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/; not tested; Neg, ISH-negative; Pos, ISH-positive; siRNA, small interfering RNA; PMO, phosphorodiamidate morpholino oligomer; VRP-GP, Venezuelan equine encephalitis replicon particles expressing EBOV glycoprotein.
**Supplementary Table 2.** *In situ* hybridization (ISH) results obtained using tissues from 24 rhesus monkeys with acute course of disease death (ACDD)

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M, Male; Pos, ISH-positive; Neg, ISH-negative;
* EBOV RNA was only detected in blood vessel and/or in vascular endothelial cells but not in parenchymal tissue of eyes or brain.
**Supplementary Table 3.** *In situ* hybridization (ISH) results obtained using tissues from 18 EBOV-infected rhesus monkeys with DTOD, delayed time of death; /, not tested; Neg, ISH-negative; Pos, ISH-positive; PMO, phosphorodiamidate morpholino oligomer; siRNA, small interfering RNA; VRP, Venezuelan equine encephalitis replicon particles

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**Supplementary Figure 1 | Detection of EBOV in the eyes of non-survivors.** Genomic EBOV RNA was detected by ISH using an EBOV NP gene-specific probe. 

- **a**, Genomic EBOV RNA in blood vessels of choroid in non-survivors. 
- **b**, Genomic EBOV RNA in ciliary process of non-survivors. 
- **c**, EBOV genomic RNA in optic nerve leptomeninges of animals with acute course of disease death (ACDD). 
- **d–d’**, Immunofluorescence demonstrated EBOV GP\(_{1,2}\) (green) within CD31-antibody (red) labeled blood vessel in.
choroids of animals with ACDD. e, Genomic EBOV RNA in fibrous connective tissue of the optic nerve leptomeninges of animals with DTOD. f, Genomic EBOV RNA in scleral connective tissues of animals with DTOD. g, Genomic EBOV RNA in ciliaries and irises of animals with DTOD. (H) Genomic EBOV RNA in choroids of animals with DTOD. i–i′, Genomic EBOV RNA staining in infiltrating cells in the anterior vitreous humor in addition to predominant infection in fibrous connective tissue of the optic nerve leptomeninges and scleras in animals with DTOD.
**Supplementary Figure 2**

**Inflammation-like response in the eyes of rhesus monkeys with persistent EBOV infection.**

- **a**–**b**, Uveitis in the ciliary bodies and irises of survivors (H&E).
- **c**, Uveitis in the corneoiroideal drainage angle (H&E).
- **d**, Uveitis in the
ciliary body and iris at the corneoirideal angle (H&E). e, Immunofluorescence stain reveals CD68$^+$ cell (macrophage/monocyte, red) infiltration in the ciliary body and iris. f, Immunofluorescence reveals most infiltrating cells to be CD68$^+$ cells (macrophage/monocyte, red) in the vitreous humor adjacent to the ciliary body. g-h, Perivascular inflammation-like response (red arrow) in the retina and likely inflammatory cells (red arrowheads) in the vitreous humor adjacent to the cell-limiting membrane (H&E). i-j, Glial fibrillary acidic protein (GFAP) immunoreactivity (green) was dramatically increased in Müller cell apical processes that extend into the inner nuclear layer (INL) and outer nuclear layer (ONL). Blue, nuclear stain by 4’,6-diamidino-2-phenylindole (DAPI).
Supplementary Figure 3

Encephalitis in a rhesus monkey survivor with persistent EBOV infection. Encephalitis is evidenced by glial nodule a-b and perivascular cuffs c. Activated microglia (red arrow) in glial nodule.
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