Durable sequence stability and bone marrow tropism in a macaque model of human pegivirus infection

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Human pegivirus (HPgV)—formerly known as GB virus C and hepatitis G virus—is a poorly characterized RNA virus that infects about one-sixth of the global human population and is transmitted frequently in the blood supply. We create an animal model of HPgV infection by infecting macaque monkeys with a new simian pegivirus (SPgV) discovered in wild baboons. Using this model, we provide a high-resolution, longitudinal picture of SPgV viremia where the dose, route, and timing of infection are known. We detail the highly variable acute phase of SPgV infection, showing that the viral load trajectory early in infection is dependent on the infecting dose, whereas the chronic-phase viremic set point is not. We also show that SPgV has an extremely low propensity for accumulating sequence variation, with no consensus-level variants detected during the acute phase of infection and an average of only 1.5 variants generated per 100 infection-days. Finally, we show that SPgV RNA is highly concentrated in only two tissues: spleen and bone marrow, with bone marrow likely producing most of the virus detected in plasma. Together, these results reconcile several paradoxical observations from cross-sectional analyses of HPgV in humans and provide an animal model for studying pegivirus biology.

INTRODUCTION

Few RNA viruses sustain high-titer viremia in humans. Those that do have evolved extensive mechanisms to evade host immune responses and induce pathological inflammation and disease over time. Human pegivirus (HPgV)—formerly known as GB virus C (GBV-C) and also as hepatitis G virus (HGV)—infects about one-sixth of the global human population and is transmitted frequently in the blood supply of the United States, yet little is known about the biology of this virus (1). HPgV belongs to the newly formed Pegivirus genus within the Flaviviridae family of positive-sense, single-stranded RNA viruses (2). Persistent, high-titer viremia is a hallmark of HPgV infection, but infection appears to be benign: it is asymptomatic and causes no overt pathology (3, 4).

Several epidemiological studies suggest that HPgV co-infection may be beneficial in the context of other viral infections that induce immune-mediated pathology (5–7). In particular, HIV–infected individuals co-infected with HPgV are protected from HIV-associated immune activation (8) and experience a 2.5-fold reduction in all-cause mortality compared to those infected with HIV only [see (9) for a meta-analysis and (1) for a review]. Although many plausible mechanisms as to how HPgV may mediate this protective effect have been proposed, compelling evidence in vivo data in support of a unifying mechanism are still lacking. Cross-sectional studies suggest that HPgV infection varies considerably between individuals and that HPgV may replicate within tissues that are difficult to access (for example, spleen and bone marrow). Furthermore, no animal model of HPgV infection currently exists. A macaque monkey (Macaca spp.) model of HPgV infection has long been sought, in part because the infection of macaques with simian immunodeficiency virus (SIV) is the most well-studied and established animal model of HIV infection (10). However, despite two reports documenting the infection of macaques with HPgV in the early 2000s (11, 12), several recent efforts to infect macaques with HPgV have been unsuccessful (J. Stapleton, personal communication). HPgV can persistently infect chimpanzees (Pan troglodytes) (13), but the use of chimpanzees in biomedical research has recently been reevaluated and is no longer ethically justifiable (14).

We recently described the first pegiviruses infecting Old World (that is, cercopithecoid) monkeys (15). These simian pegiviruses (SPgVs) were identified in red colobus monkeys (Procolobus tephrosceles; SPgVkr), red-tailed guenons (Cercopithecus ascanius; SPgVkr), and an olive baboon (Papio anubis; SPgVob) living in Kibale National Park, Uganda, and formed a distinct clade with respect to the Great Ape and New World monkey SPgVs. Here, we extend the geographic and host range of the Old World monkey SPgVs with the discovery of a new SPgV in samples collected from yellow baboons (Papio cynocephalus; SPgVmyb) in Mikumi National Park, Tanzania, and use baboon SPgVs to create a macaque model of HPgV infection.

RESULTS

Baboons naturally harbor distinct pegiviruses

As part of our ongoing effort to discover new viruses in wild nonhuman primates, we applied unbiased deep sequencing to plasma samples collected from 43 wild yellow baboons living in Mikumi National Park, Tanzania, during 1985 and 1986. In 17 of these animals (40%), we detected a new SPgV (SPgVmyb) and were able to assemble 14 coding-complete SPgVmyb genomes (Fig. 1A and table S1). In a phylogenetic analysis, these viruses grouped with other Old World monkey SPgVs...
and were most closely related to SPgV kob (Fig. 1B). However, plasma viral loads obtained by quantitative reverse transcription polymerase chain reaction (RT-qPCR) for SPgV myb-infected yellow baboons (average, $2.7 \times 10^5 \pm 7.3 \times 10^4$, SEM) were significantly less than the plasma viral load observed for the single SPgV kob-infected olive baboon identified previously ($2.4 \times 10^7$) (Fig. 2, A and B) (15).

**Baboon SPgVs infect macaques**

Among the monkey species in which we have discovered SPgVs, baboons are most closely related to macaques (Fig. 1C). Therefore, in an attempt to create an in vivo laboratory model of HPgV infection, we inoculated a Mauritian-origin cynomolgus macaque (cy0571) intravenously with 0.5 ml of plasma pooled from two Mikumi yellow baboons.
Fig. 2. Infection of macaques with baboon SPgVs. (A) Plasma SPgVmyb viral loads in the Mikumi yellow baboons (yellow) and the longitudinal viral loads from cy0571 infected with SPgVmyb. (B to E) Plasma SPgVkob viral load in the single Kibale olive baboon (green) and the longitudinal viral loads of macaques infected with SPgVkob from olive baboon plasma (green), cy0500 plasma collected at 8 dpi (D, black), or cy0575 plasma collected at 25 dpi (E, blue, male) or 36 dpi (C, pink, female). Y axis, log_{10} SPgV gc per milliliter of plasma; x axis, days post-infection unless otherwise stated. Specific colors and symbols refer to the same animal throughout the manuscript. Animal IDs surrounded by a box indicate SIVmac239 infection before SPgV inoculation, with dashed boxes signifying SIVmac239Δnef infection/vaccination. Additional information about each animal can be found in table S2. S, whole genome sequencing; X, splenectomy.
(baboons M23 and M77) containing a total of $2.0 \times 10^5$ genome copies (gc) of SPgVmb (Fig. 2A). A second macaque (cy0572) was inoculated intravenously with 0.5 ml of Kibale olive baboon plasma (baboon K23) containing $1.2 \times 10^7$ gc of SPgVkob (Fig. 2B). Before inoculation of these macaques, baboon plasma samples were screened for additional microbes (for example, simian arteriviruses) using a combination of unbiased deep sequencing, RT-PCR, RT-qPCR, and culture-based techniques (16–18). Blood was drawn from each macaque and assayed biweekly for the presence of SPgV using RT-qPCR assays specific for each virus.

SPgVmb was first detected in the plasma of cy0571 at $1.1 \times 10^4$ gc/ml 10 days post-inoculation (dpi) and slowly increased to a peak of $4.8 \times 10^5$ gc/ml at 56 dpi before gradually declining to a “set point” of $3.0 \times 10^4$ gc/ml, which was maintained until the animal was euthanized at 219 dpi (Fig. 2A). (Set point was defined throughout this study as the average of at least three viral load measurements, taken over a period of at least 25 days, that did not differ by more than 1 log$_{10}$.) In contrast, SPgVkob was first detected in the plasma of cy0572 at $4.1 \times 10^6$ gc/ml 7 dpi and rapidly increased to a peak of $3.1 \times 10^5$ gc/ml by 10 dpi (Fig. 2B). By 21 dpi, viremia had reached a relative nadir of $2.7 \times 10^6$ gc/ml but then gradually increased to a set point of $4.5 \times 10^6$ gc/ml, which was maintained for 31 days before rapidly declining to undetectable by 98 dpi.

**Infected dose dictates acute-phase SPgV viral load trajectory but not chronic-phase viremic set point**

To examine viremia in a larger number of individuals, two additional cynomolgus macaques (cy0575 and cy0500) were infected with olive baboon plasma containing SPgVkob as done previously (Fig. 2B), and plasma from these animals was passaged into 11 additional cynomolgus macaques (Fig. 2, C to E). These macaques were repurposed from several other infection studies, and several were SIV-positive. However, heterogeneity in the infecting strain of SIV, the duration of SIV infection, and host genetic factors precluded us from examining the effect of SPgV co-infection on SIV pathogenesis (table S2 and fig. S1). We found that the acute phase of SPgV viremia was highly dependent on the infecting dose, with high-dose infections ($1.2 \times 10^7$ gc to $2.3 \times 10^7$ gc) reproducibly achieving a high-titer peak of viremia within the first 8 to 12 dpi, whereas low-dose infections ($6.4 \times 10^3$ gc to $6.0 \times 10^4$ gc) had a highly variable acute-phase viral load trajectory, with several animals not reaching their lifetime maximum of viremia until after 100 dpi (Fig. 3, A to C). Longitudinal viral load analysis in SPgVkob-infected macaques revealed that the inoculating dose did not affect the chronic-phase viremic set point (Fig. 3D). Durable SPgV infection could not be established in one macaque (cy0649) despite re-inoculation with $1.1 \times 10^5$ gc of SPgVkob from animal cy0575 at 43 dpi. Additionally, SPgV in plasma declined to undetectable levels after about 100 days of detectable viremia in four macaques, including the original two animals inoculated with olive baboon plasma, highlighting individual differences in the course of SPgV infection (Fig. 2).

**Baboon SPgVs elicit little to no disease in macaques**

At no point after SPgV infection did macaques develop overt signs of disease, nor were gross hematologic or postmortem abnormalities observed in SIV-negative macaques infected with SPgV (fig. S2A). In 9 of 12 animals chronically infected with SIV and SPgV, liver sections revealed minimal to mild periportal and random lymphohcytic hepatitis (fig. S2, B and C). The severity of hepatitis in these animals was not greater than typically noted in SIV-infected cynomolgus macaques.
SPgVs exhibit a high degree of sequence stability
We performed deep sequencing of SPgVkob from several infected macaques at multiple time points (Fig. 2 and tables S3 to S5). Nucleotide variants that rose to greater than 50% of the within-host viral population were rare, with each intrahost viral population accumulating an average of 1.5 consensus-level variants per 100 days of infection, although no consensus-level variants were detected in some individuals despite persistent viremia (Fig. 4, A to C). Of the 11 nonsynonymous variants that were detected across all animals, 7 were due to one of two identical changes observed in the putative NS2 protein (Fig. 4, A, B, and D). One of these variants (T746M) was present at 0.72% in the inoculum administered to cy0565, cy0567, and cy0568 but was not detectable in the inoculum administered to cy0651 or cy0562, yet rose to near fixation in these five animals (Fig. 4D). A different amino acid substitution (A→N) was found at the same codon position in SPgVmyb at 219 dpi of cy0571 (table S5).

Sequencing analysis of SPgVmyb in wild yellow baboons revealed a degree of sequence stability similar to what we observed for SPgV in macaques, with dN/dS greater than 2.0 at four sites throughout the genome (fig. S3). Subconsensus-level variation in SPgV-infected macaques was also low, with longitudinal intrahost analyses revealing a viral population that remained highly homogeneous during the acute-phase burst of viral replication and remained stable with low diversity throughout the chronic phase of infection (Fig. 5).

SPgV is primarily bone marrow–tropic
Studies investigating the tissue tropism of HPgV have been limited (19–21). Therefore, we sought to comprehensively and quantitatively define the tissue distribution of SPgV at each stage of infection. We collected a large panel of tissues from four macaques (cy0500, cy0493, cy0388, and cy0571) at 8, 13, 21, and 219 dpi, respectively. RNA was extracted from each tissue sample, quantified by RT-qPCR, and normalized to obtain a tissue-specific viral load per milligram of tissue. To identify tissues containing SPgV RNA at concentrations greater than plasma, tissue viral loads were normalized to the plasma viral load (in microliters) of the respective individual at the time of tissue collection. This approach revealed that SPgV RNA is highly concentrated in only two tissues: the spleen and bone marrow (Fig. 6A). To investigate the relative contribution of each of these tissue compartments to systemic SPgV viral load (that is, virus production), we surgically removed the spleen from six macaques via laparotomy (that is, splenectomy) and closely monitored plasma viral loads for an extended period of time after splenectomy (Fig. 6B). With the exception of one macaque (cy0572), splenectomy did not have a significant effect on plasma viral load (Fig. 6, B and C).

To assess the concentration of SPgV RNA on a per-cell basis, we performed SPgV-specific RT-qPCR on leukocytes purified from spleen...
Fig. 5. SPgV maintains low intrahost diversity over the course of infection. Deep sequencing reads from time points throughout the infection of cy0572 were mapped to the SPgVkob_K23 reference genome, and variants >1% were identified (see table S4 for sequencing statistics and coverage details). A schematic of the SPgVkob genome with putative protein products is shown at the top. Blue bars represent synonymous variants present in the viral population, and red bars represent nonsynonymous variants.
and bone marrow. This approach also allowed us to measure SPgV viral loads in peripheral blood mononuclear cells (PBMCs), a lymphoid compartment in which HPgV RNA has been detected (22–24). No significant differences in the quantity of cell-associated SPgV RNA were observed between spleen, bone marrow, and PBMCs (Fig. 6D). However, stratification of these data by the phase of infection during which tissues were harvested suggests that RNA may be most concentrated in PBMCs during the acute phase and most concentrated in bone marrow during the chronic phase (fig. S4).

**DISCUSSION**

HPgV infection is highly prevalent among humans, yet fundamental aspects of HPgV biology remain poorly understood. Our discovery of prevalent SPgV infection in yellow baboons from Tanzania extends the known host range and geographic range of these viruses and suggests that species-specific variants of SPgV may be widespread among primates. Infection of macaques—a “non-natural” host for these baboon SPgVs—did not elicit any overt signs of disease including histologic evidence of hepatitis during acute infection. In a fraction of animals evaluated in the chronic phase of SPgV infection, we observed histologic changes indicative of minimal to mild hepatitis. Although it seems likely that concurrent SIV infection contributed to the development of hepatitis in these animals, the severity of hepatitis we observed was still significantly less than reported previously in macaques inoculated with HPgV (11). The apparent nonpathogenicity of SPgVs from African Old World monkeys in macaques contrasts starkly with other blood-borne RNA viruses naturally found in African Old World monkeys, namely, the SIVs and simian arteriviruses (also known as simian hemorrhagic fever viruses), which respectively cause AIDS-like disease and viral hemorrhagic fever in macaques (10, 25). This raises the intriguing possibility that PgV infection in primates may be universally nonpathogenic, although detailed long-term studies are needed to further examine PgV pathogenesis (or lack thereof).

This study provided the opportunity to examine the replication kinetics of SPgV during the acute phase of infection. We found that the acute phase of SPgV viremia was highly dependent on the infecting dose, with high-dose infections (1.2 × 10^7 gc to 2.3 × 10^7 gc) reproducibly achieving a high-titer peak of viremia within the first 8 to 12 dpi, whereas low-dose infections (6.4 × 10^3 gc to 6.0 × 10^4 gc) had a highly variable acute-phase viral load trajectory, with several animals not reaching their lifelong maximum of viremia until after 100 dpi. The high-titer viremia we observed within 2 weeks of SPgV inoculation in macaques is consistent with the kinetics of viremia seen in patients infected with HPgV via blood transfusion (26) but contrasts with experimental high-dose inoculation of HPgV in chimpanzees in which viremia was not detectable until 10 weeks after inoculation (13), possibly suggesting that the species barriers to PgV replication may be lower in baboons and macaques than in humans and chimpanzees. Additionally, the relationship between
infecting dose and viral load trajectory is in contrast to HIV and SIV, which reproducibly replicate to a high-titer peak during the first few weeks of infection independent of infecting dose (27, 28). The apparent connection between dose and viral load trajectory could explain the wide range of HPgV viral loads observed cross-sectionally in HPgV-infected humans (29, 30), because people exposed via sexual contact may be infected by a relatively low dose, whereas those exposed via the administration of HPgV-positive blood products may be infected with a much greater dose (31–36). The eventual clearance of SPgV RNA is caused by a primary site of SPgV replication in a macaque with very little active bone marrow (cy0493), which seems likely that SPgV RNA is concentrated in only the primary site of SPgV replication. However, considering that we observed robust acute-phase SPgV replication in a macaque with very little active bone marrow (cy0493), it seems likely that SPgV production can occur in other tissues. One possible explanation for this observation may be that SPgV tropism shifts throughout infection, with the PBMCs and spleen supporting most of the viral replication early during infection and the bone marrow sustaining chronic, persistent infection.

The robust replication of baboon SPgVs in macaques appears to recapitulate key features of HPgV infection and is an advance for HPgV research. However, this study has several limitations. Most notably, the macaques infected with SPgV in this study were heterogeneous in several important respects including SIV status, SIV disease progression, age, sex, infecting SPgV strain, and infecting SPgV dose. Future studies that control for these factors are necessary to more clearly define the impact of these variables on SPgV infection. Controlling for variability in SIV pathogenesis will be particularly challenging but will be essential for the study of SPgV/SIV co-infection as a model of HPgV-mediated protection from HIV pathogenesis. Additional studies are also needed to determine whether or not observations regarding SPgV tissue tropism and the effect of SPgV infecting dose on acute-phase viremia faithfully recapitulate HPgV infection in humans.

The cross-species transmission of an RNA virus that replicates to high titers and persists without accumulating genetic variation or causing disease violates conventional wisdom and could have far-reaching implications for understanding the interplay between viral persistence, inflammation, immune tolerance, and disease. Future studies should focus on identifying the cell type permissive for PgV replication and the mechanisms by which PgVs maintain persistence. Studies of macaques co-infected with PgV and pathogenic SIV should also be prioritized, because this could shed light on the significant but poorly understood phenomenon of HPgV-mediated protection from immune activation and AIDS.

MATERIALS AND METHODS

Study design

This was a proof-of-concept study designed to establish a macaque model of HPgV infection. All protocols were approved by the University of Wisconsin–Madison Institutional Animal Care and Use Committee conforming to National Institutes of Health (NIH) animal care guidelines. A total of 14 macaques were infected with SPgV/kob from an olive baboon (1.2 × 10^5 gc to 2.3 × 10^5 gc, n = 4; 6.0 × 10^4 gc, n = 5; 5.6 × 10^3 gc, n = 5) and used to study viral transmission, tissue tropism, persistence, and intrahost evolution. One macaque was infected with 2.0 × 10^5 gc of SPgVmyb from two yellow baboons.

Collection of samples from wild baboons

Blood samples from Kibale olive baboons (P. anubis) were collected in 2010 to 2014 as described previously (15, 42), and approved by the Uganda Wildlife Authority (permit UWA/TDO/33/02), the Uganda National Council for Science and Technology (permit HS 364), and the University of Wisconsin Animal Care and Use Committee (protocol V01409-0-02-09). Samples were imported under CITES permit #002290 (Uganda). Samples from Mikumi yellow baboons (P. cynocephalus) were collected by Rogers and colleagues in 1985 and 1986 using standard methods for field studies of baboons as described previously (43). All procedures were approved by the appropriate Tanzanian government.
Care and use of macaques at the Wisconsin National Primate Research Center

All macaque monkeys used in this study were cared for by the staff at the Wisconsin National Primate Research Center (WNPRC) in accordance with the regulations and guidelines outlined in the Animal Welfare Act and the Guide for the Care and Use of Laboratory Animals. Details of this study (University of Wisconsin–Madison Animal Care and Use Protocol No. G00707) were approved by the University of Wisconsin Institutional Animal Care and Use Committee, in accordance with recommendations of the Weatherall report. Animals used in this study were repurposed from several different studies, the details of which can be found in table S1.

Inoculations. Plasma samples were thawed at room temperature, and the appropriate volume (200 to 500 μl) was loaded into a 1-ml syringe and kept on ice until administration. Animals were sedated, and inocula were administered via an intravenous catheter followed by a sterile saline flush. At the conclusion of the procedure, anesthesia was reversed and animals were closely monitored by veterinary staff for adverse reactions and signs of disease.

Deep sequencing. Samples were processed for sequencing in a biosafety level 3 laboratory as described previously (15, 42) with slight modifications. Briefly, for each animal, viral RNA was isolated from about 200 μl of plasma using the QIAamp MinElute virus spin kit (Qiagen), omitting carrier RNA. Samples were then treated with deoxyribonuclease (DNase), and complementary DNA (cDNA) synthesis was primed using random hexamers (Double-Stranded cDNA Synthesis Kit, Invitrogen). Samples were fragmented and sequencing adaptors were added using the Nextera DNA Sample Preparation Kit (Illumina). Deep sequencing was performed on the Illumina MiSeq. Sequence data were analyzed using CLC Genomics Workbench 6.5 (CLC bio) and Geneious R5 (Biomatters). Low quality (<Q30, Phred quality score) and short reads (<100 base pairs (bp)) were removed, and coding-complete (44) genome sequences for each virus were acquired using the de novo assembly algorithm in CLC Genomics Workbench version 6.5. Viral genomes were annotated in CLC Genomics Workbench version 6.5, and putative open reading frames were confirmed by querying the National Center for Biotechnology Information (NCBI) GenBank database (45).

Phylogenetic analysis. Complete coding sequences representing the known diversity of pegiviruses were aligned using a codon-based version of open source software multiple alignment using fast Fourier transform (MAFFT) (46) and cleaned using Gblocks, implemented in TranslatorX (47). Phylogenetic history was inferred from aligned nucleotide sequences using the maximum likelihood method (1000 bootstrap replicates) via Molecular Evolutionary Genetics Analysis version 6 (MEGA6) open source software (48). The best nucleotide substitution model, general time reversible model coupled with a Γ distribution for rate variation (GTR+Γ), was estimated using MEGA6. All positions containing gaps and missing data were eliminated, resulting in a final data set of 8118 positions. The initial tree for the heuristic search was obtained by applying the neighbor-joining method to a matrix of pairwise distances, estimated using the maximum composite likelihood approach.

Evolutionary analysis. Complete coding sequences for all 14 SpgVmyb genomes were acquired using the de novo assembly algorithm in CLC Genomics Workbench version 6.5. dN and dS were calculated using the HIV Synonymous Non-synonymous Analysis Program (SNAP) version 2.1.1 (49, 50) found at www.hiv.lanl.gov.

Quantitative reverse transcription PCR

A TaqMan RT-qPCR assay was developed to quantify viral RNA for both SpgVkob (forward primer: 5′-CGGTTGTTGTCGGCTGAT-3′; reverse primer: 5′-CAAGTGTTAGCAGGCGAT-3′; probe: 5′-6FAM-ATGACGCCTGATGTAAGCTGGGCAABHQ1-3′) and SpgVmyb (forward primer: 5′-GTTGGAGCCGACCTAGCA-3′; reverse primer: 5′-CTAACCTCAGGGCAGGATT-3′; probe: 5′-6FAM-CGGCGCTGAACTGGGCTTAC-BHQ1-3′). A standard curve was created by cloning a 381-bp fragment of the SpgV kob_K23 genome (forward primer: 5′-CCCCCTGGTGTGGTTAGA-3′; reverse primer: 5′-CACTGATCGCCTCAGGTA-3′) into the zero blunt PCR vector (Invitrogen), followed by linearization (Hind III, New England Biolabs) in vitro transcription for 6 hours (MEGAscript 17 Transcription Kit, Invitrogen), cleanup (MEGAclear Transcription Cleanup Kit, Invitrogen), quantification (Qubit RNA High-Sensitivity Assay Kit, Invitrogen), and dilution to a concentration of 1 × 10^10 transcription copies per microliter. Tenfold dilutions of this transcript were used as a standard curve for both SpgVkob and SpgVmyb. RNA was reverse-transcribed and amplified using the SuperScript III One-Step qRT-PCR system (Invitrogen) on a LightCycler 480 (Roche). Reverse transcription was carried out at 37°C for 15 min and then 50°C for 30 min followed by 2 min at 95°C, and then 50 cycles of amplification as follows: 95°C for 15 s and 60°C for 1 min. The reaction mixture contained MgSO4 at a final concentration of 3.0 mM, with the two amplification primers at a concentration of 500 nM and probe at a concentration of 100 nM. The standard curve was linear over (at least) eight orders of magnitude and was sensitive down to 10 copies of RNA transcript per reaction.

Purification of RNA from plasma. Animals were sedated and blood was drawn from the femoral vein into an EDTA-treated vacutainer tube and centrifuged at 1400g for 7 min. Plasma was collected and centrifuged for an additional 5 min at 670g to remove residual cells. RNA was extracted from 300 μl of plasma using the Viral Total Nucleic Acid Purification Kit (Promega) on a Maxwell 16 MDx instrument.

Purification of RNA from cells. Macaques were appropriately sedated, and tissues were excised, chopped into ~1-cm cubes, and placed in 30 ml of ice-cold RPMI + 10% fetal calf serum (FCS) (Thermo Fisher Scientific). To achieve single-cell suspensions, tissues were manually disrupted with a scalpel and the butt of a sterile 60-ml syringe and then passed through a 100-μm cell strainer. Blood was overlayed on Ficoll at a 1:1 ratio and centrifuged at 1860g for 30 min to obtain PBMCs. Cells from each sample were washed with 40 ml of RPMI + 10% FCS, treated with ACK lysis buffer to remove red blood cells, and washed twice more with 40 ml of RPMI + 10% FCS. Centrifugations were performed at 670g for 5 min at 4°C, and samples were kept on ice throughout processing. Cells from each tissue were then counted (Z2 particle counter, Beckman Coulter). For each sample, 3.0 × 10^6 cells were disrupted using a QIAshredder column (Qiagen) and RNA was isolated using the Cells LEV Total RNA Purification Kit (Promega) on a Maxwell 16 MDx instrument. All extractions were performed in duplicate.
Purification of RNA from tissues. Immediately before euthanasia, macaques were sedated with ketamine and sodium pentobarbital was administered. Transcardiac perfusion of the entire body with 5 liters of sterile saline was performed to remove blood from the cardiovascular tree and vessels within tissues. Tissues were collected for routine histologic analysis, and three sections of each tissue (about 3 cm × 1 cm × 0.4 cm) were excised using autoclave-sterilized instruments and placed immediately into 12 ml of RNAlater (Sigma-Aldrich). Tissues were allowed to sit in individual tubes at −80°C. Tissues were homogenized using the TissueLyser II (Qiagen) according to the manufacturer’s instructions, and RNA was extracted from each tissue using the Tissue LEV Total RNA Purification Kit (Promega) on a Maxwell 16 MDx instrument.

Splenectomy. Animals were sedated and intubated. The spleen was surgically removed, and splenic vessels were ligated to prevent blood loss. At the conclusion of the procedure, animals were extubated and closely monitored by veterinary staff for adverse reactions and signs of disease.

Statistical analysis
Phylogenetic analyses were performed using a maximum likelihood method (1000 bootstrap replications) via MEGA6 open source software. Statistical analyses were performed using GraphPad Prism 5.0 software. Peak viremia and time-to-peak viremia in high-dose versus low-dose groups were compared using a two-tailed paired t test (α = 0.05). Comparison of set-point viremia (as defined by the average of at least three viral load measurements taken over a period of at least 25 days that did not differ by more than 1 log10) among groups was performed using a one-way ANOVA (α = 0.05). Synonymous and nonsynonymous mutation rates were analyzed using univariate linear regression. Viral loads from each tissue were compared using one-way ANOVA (α = 0.05). Plasma viral loads before and after splenectomy were compared using a two-tailed paired t test (α = 0.05). Viral loads from different cellular compartments were compared using one-way ANOVA.

SUPPLEMENTARY MATERIALS
www.sciencetranslationalmedicine.org/cgi/content/full/7/305/305ra144/DC1
Fig S1. Parameters of SW-induce d disease in study animals.
Fig S2. Lack of SPgV-induced disease in study animals.
Fig S3. SPgVs exhibit little sequence variability in natural hosts.
Fig S4. The tissue tropism of SPgV may vary depending on the duration of infection.
Table S1. Sequencing of SPgVmb in cy0571.
Table S2. Sequencing of macaques infected with SPgVkb.
Table S3. Demographic information for macaques used in this study.
Table S4. Sequencing of macaques infected with Tissue LEV Total RNA Purification Kit (Promega) on a Maxwell 16 MDx instrument.

REFERENCES AND NOTES


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Data and materials availability: Coding-complete genome sequences for the SPgV/mObv strains used in this study have been made available in GenBank under accession numbers KP890672-KP890673 and KP96142-KP96153. The coding-complete genome sequence for the SPgV/kob strain used in this study was published previously (15) and can be found in the GenBank database under accession number KF234530.

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Durable sequence stability and bone marrow tropism in a macaque model of human pegivirus infection
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Editor's Summary

An animal model for the "Good Boy Virus"

Human pegivirus (HPgV, formerly called GB virus C) can protect HIV-infected individuals from developing AIDS, a phenomenon that has earned it the nickname of the "Good Boy Virus." How HPgV imparts this protective effect remains a mystery, in part because no animal model of HPgV infection exists. Bailey and colleagues discovered viruses related to HPgV in wild baboons and showed that these viruses can infect laboratory macaque monkeys, providing unprecedented insights into the transmission, replication, and anatomical preference of HPgV with a view towards understanding HPgV-mediated protection from AIDS.

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