An Experimental Murine Model to Study Acquisition Dynamics of Tick-Borne Langat Virus in Ixodes Scapularis

Kundave V. Rajendran  
University of Tennessee at Knoxville

Waqas Ahmed  
University of Tennessee at Knoxville

Ashkan Roozitalab  
University of Tennessee at Knoxville

Girish Neelakanta  
University of Tennessee at Knoxville

Hameeda Sultana (✉ hsultana@utk.edu)  
University of Tennessee at Knoxville

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An experimental murine model to study acquisition dynamics of tick-borne Langat Virus in *Ixodes scapularis*

Kundave V. Rajendran 1, #, $, Waqas Ahmed 1, #, $, Ashkan Roozitalab 1, #, Girish Neelakanta 1, 2, #, and Hameeda Sultana 1, 2, #, *

1 Department of Biological Sciences, College of Sciences, 2 Center for Molecular Medicine, Old Dominion University, Norfolk, VA, USA.

# Present Address for all authors: Department of Biomedical and Diagnostic Sciences, College of Veterinary Medicine, University of Tennessee, Knoxville, TN, USA.

$ Equal Author Contribution: These authors contributed equally to this work.

Running title: Acquisition of murine LGTV by nymphal and larval ticks

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* Corresponding Author: Department of Biomedical and Diagnostic Sciences, College of Veterinary Medicine, University of Tennessee, Knoxville, TN 37996, USA, Email: hsultana@utk.edu, Phone: (865) 974-8217.
Abstract

*Ixodes scapularis* ticks acquire several pathogens from reservoir animals and transmit them to humans. Development of an animal model to study acquisition and transmission dynamics of these pathogens into and from ticks, respectively, is challenging due to the fact that in nature ticks feed for a longer duration and on multiple vertebrate hosts. To understand the complex nature of pathogen acquisition/transmission, it is essential to set up a successful tick blood feeding method on a suitable vertebrate host. In this study, we provide an evidence that murine model can be successfully used to study acquisition dynamics of Langat virus (LGTV), a member of tick-borne flaviviruses. Mice were inoculated intraperitoneally with LGTV that showed detectable viral loads in murine blood, skin and other tissues including brain. Both larval and nymphal ticks that were allowed to feed on murine host successfully acquired LGTV loads. Also, we found that after molting LGTV was transstadially transmitted from larval to nymphal stage. In addition, we noted that, LGTV down-regulated *IsSMase* expression in all group of ticks possibly for its survival in its vector host. Taken together, we provide an evidence on the use of murine model to not only study acquisition dynamics of LGTV but also to study changes in tick gene expression during acquisition of arboviruses into ticks.
Introduction

Tick-borne flaviviruses are recognized as one of the major global concerns to human health. These viruses are mostly transmitted through the bite of an infected *Ixodes* tick that results in meningitis, encephalitis, biphasic fever, febrile illness or hemorrhagic fever in humans \(^1^,\) \(^2\). Tick-borne flaviviruses including Tick-borne encephalitis virus (TBEV), Powassan virus (POWV), Langat virus (LGTV), Kyasanur forest disease virus (KFDV), Louping ill virus and Omsk hemorrhagic fever virus (OHFV) are both genetically and antigenically related to one another \(^1^,\) \(^3\)-\(^5\). Several studies, including our own, have primarily focused on studying tick-bacteria interactions \(^6^-^1^2\). In contrast, limited number of studies have focused on understanding interactions between tick-borne flaviviruses and their arthropod vectors at the molecular level \(^1^3^-^1^5\). Therefore, a detailed understanding of the connection between tick-borne flaviviruses and their vector host is essential to design preventive strategies and to disrupt the pathogen transmission.

LGTV is a naturally attenuated flavivirus with nonpathogenic/low-pathogenic concerns to human health. Therefore, our laboratory used it as a model pathogen to understand the interactions of virulent tick-borne flaviviral infections in ticks. In addition, work with LGTV can be safely performed at the Biosafety level 2 (BSL-2) \(^1^6\). Like other tick-borne flaviviruses, LGTV has a lipid-envelope, and a positive-sense single-stranded RNA genome of approximately 11 kb in length \(^1^7\). Arthropods play key role(s) in the acquisition and transmission of numerous viruses in different tropical and subtropical regions of the world \(^1^8^,\) \(^1^9\). Ticks have developed paramount strategies to indulge as successful vectors for pathogen acquisition and transmission.
They have an ability in efficiently taking a blood meal through biting host and allowing viral survival and replication within them for an extensive period of time \(^{20-23}\). Ixodid ticks are reported to be the most prominent vectors that play key role(s) in the flavivirus acquisition and transmission cycle \(^{20}\). Flaviviruses have the ability to infect Ixodid ticks at any developmental stage (larvae, nymphs or adults), and the infection can be maintained throughout the tick’s lifespan \(^{20,22,24}\). Tick-borne flaviviruses usually spend 95% of their life cycle in *Ixodes* tick vector than in the vertebrate host including accidental human host \(^{25}\).

Regardless of the significance of Ixodid ticks as a key vector for bacterial acquisition and transmission, little is known regarding the flavivirus-vector host interactions. An understanding of the molecular mechanism(s) is essential to gain in-depth knowledge on the acquisition of the virus from the vertebrate host to ticks, pathogen replication in the tick midgut, crossing of the mid-gut barrier, persistent survival in the salivary glands, and then transmission to the vertebrate host. Technical challenges faced in a research laboratory have limited the number of studies conducted on the flavivirus-tick host interactions. The two common methods used for infecting ticks in a laboratory condition are parenteral inoculation by microinjection \(^{26}\) and synchronous infection of ticks by immersion method \(^{20}\). There are several limitations for these two methods, as majority of the flaviviruses need high biocontainment level facilities (such as BSL-3 or BSL-4), which add to the technical issues on working with these pathogens. Microinjection method to generate LGTV-infected ticks is an artificial system. The success in generating LGTV-infected ticks by microinjection method greatly depends on the ability of LGTV to cross the midgut barrier, reach salivary glands, then replicate and persist as a colony. This factor is a critical determinant to assess vector competency in these artificially-infected ticks \(^{26,27}\). For synchronous
LGTV-infection of ticks, large volumes (1-2 ml) of concentrated virus stocks (such as $10^7$ or $10^8$ pfu/ml) and longer incubations (17 days) are required for generation of small batch of infected ticks. Also, only 60-70% of ticks were capable of acquiring pathogen loads by body absorbance or bathing in the virus stocks $^8,^{10}$.

Our previous studies have demonstrated the role of arthropod-derived Extracellular Vesicles (EVs or exosomes) in the transmission of flaviviruses such as LGTV, and DENV2/3, which perhaps is mediated by the CD63 ortholog Tsp29Fb $^{16,28,29}$. In addition, we noted that LGTV infection, suppress *Ixodes scapularis* *Ismase* gene expression, a molecule involved in exosome biogenesis $^{10}$. *Ismase* (Sphingomyelinase D or SMase D, a venomous protein ortholog of spiders) is a tick saliva component that regulates the cytokine expression to modulate the programming of immune response and perhaps via arthropod exosomes $^{10,30}$. *Ismase* neutralizes the Th1 cytokine response towards a Th2-induced cytokine response and modulates CD4+ T cells in order to express interleukin 4 (IL-4) $^{10,30}$. Our recent studies focused on understanding the molecular mechanisms during pathogen-vector-host interactions have revealed different survival strategies used by microbes $^{6-10,16,28,29}$. In the current study, utilizing LGTV as a model pathogen, we established a method to infect naïve *I. scapularis* by feeding these ticks on LGTV-infected blood from murine host. Viral loads were detected both in the murine tissues and in nymphal/larval ticks fed on LGTV-infected mice suggesting viral-replication in mice blood and dissemination into tissues including skin and furthermore acquisition into the nymphal/larval tick’s body from the infected murine host. We have also provided evidence for the transstadial transmission of LGTV from the fed larval ticks to molted nymphs. In summary, our study provides a method not only to generate LGTV-infected ticks but also provides a tool to study
acquisition and transmission dynamics of this virus and perhaps other high priority tick-borne flaviviruses, such as TBEV and POWV.
Results

LGTV infection is detected in murine blood collected from mice that allowed the feeding of during and post fed nymphal/larval ticks. This study was conducted to demonstrate the ability of I. scapularis ticks to acquire LGTV loads from the infected murine host. So far, we believe that no tick acquisition studies have been recorded and shown in the literature for tick-borne LGTV. Our study provides evidence for the acquisition of tick-borne LGTV by both nymphal and larval ticks upon feeding on an infected murine host. Studies that determine tropism of LGTV in rodents are minimal. Therefore, we determined viral loads in murine blood and other murine tissues (such as liver, spleen, skin, and brain). At days 3-5 p.i., LGTV loads were highly detectable in blood samples of all mice inoculated intraperitoneal via needle, and at a dose of 500,000 (5 x 10^5) pfu/ml. Mice blood was collected at both the time points (of 48 h DF or 120 h PF, and these time points correspond to 3 or 5 days p.i., respectively). The schematics show the experimental plans for tick acquisition (nymphal or larval acquisition of LGTV loads) carried out in this study (Figs. 1A and 1B). QRT-PCR analysis showed that LGTV readily infected mice with increased viral loads detected at an early stage of infection (3 days p.i.,) in comparison to the later stage of infection (5 days p.i.,) (Fig. 2A). QRT-PCR amplification showed LGTV (143 bp) product on 1.5 % agarose gel electrophoresis in both the infected groups of mice (3 days p.i. or 5 days, p.i.,), while as expected the uninfected control group (both 3 or 5 days p.i.) showed no amplification products (Fig. 2B). A similar experiment was carried out to feed larval ticks on LGTV-infected mice for collection of ticks and murine blood and tissues at 120 h PF time point (or 5 days p.i.,). This subsequent experiment also showed successful LGTV infection in mice via needle inoculation (Fig. 2C). The gel images showed amplicons of the LGTV product (143 bp)
in the infected group, while no product was found in the uninfected mice (Fig. 2D). These results indicated a successful infection of mice with detectable LGTV loads in the murine blood.

**Detection of LGTV loads in murine tissues.** Tissues such as spleen, liver, skin and brain were collected at different time points (of 48 h DF or 120 h PF, that correspond to 3 or 5 days p.i., respectively) and analyzed for detection of LGTV loads by QRT-PCR analysis. At 3 days p.i., LGTV loads were detected to be higher in all tested tissues compared to the levels noted at day 5 p.i. (Fig. 3A and 3B). At 5 days p.i., the tissues remained positive for LGTV infection with detectable viral loads. Among the tested tissues and at both time points of days 3 or 5 p.i., LGTV showed higher detectable loads in liver, followed by skin, spleen and brain (Fig. 3A and 3B). The agarose gel images confirmed the presence of amplicons (product of 143 bp) in the LGTV-infected mice tissues and no amplification products were detected in the uninfected controls as expected (Fig. 3C, 3D and 3E). Tissues were also collected from the mice that were used for larval tick feeding. LGTV loads in these mice showed similar trend as noted in samples generated from mice used for nymphal tick feeding (Fig. 3B and Fig. 4A). All mice tissues remained positive for LGTV loads in these group of murine hosts that allowed larval tick feeding. However, liver showed higher viral loads followed by skin, spleen, and brain (Fig. 4A) a trend similar to samples generated from mice used for nymphal tick feeding. Gel electrophoresis analysis confirmed the presence of LGTV amplicons in all infected mice tissues, and as expected no amplification product was noted in the uninfected control group of mice (Fig. 4B and 4C).

**Acquisition of LGTV loads by nymphal ticks.** Viral loads were highly detectable in all tested ticks that included both the partially fed group (of 48 h DF ticks, collected at 3 days p.i., of mice)
or the fully fed/engorged repleted group of ticks (120 h PF, collected at 5 days p.i. of mice) (Fig. 5A). There were no detectable LGTV loads in the respective uninfected groups of ticks collected from either 48 h DF or 120 h PF groups of uninfected control mice (Fig. 5A). No significant difference in viral loads were noted between ticks collected during 48 h DF or 120 h PF group (Fig. 5A). Amplification of LGTV product (143 bp) was recorded as an enhanced signal in both DF, and PF group of ticks (Fig. 5B and 5C). These results elucidate that naïve nymphs fed on LGTV-infected mice are capable of acquiring the virus from the murine host.

**Acquisition of LGTV loads by larval ticks.** Larval ticks were allowed to feed completely on LGTV-infected or uninfected control mice. These ticks were collected after repletion from mice and were considered as 120 h PF group (5 days p.i. of mice). QRT-PCR analysis revealed the presence of viral loads in all individually processed larval ticks that were fully fed to repletion (Fig. 6A). Agarose gel electrophoresis of QRT-PCR products confirmed the presence of amplicons corresponding to 143 bp of LGTV fragment (Fig. 6B). Except for two fed larval ticks, all other ticks showed the presence of amplicons and were positive for LGTV loads (Fig. 6A and 6B). As expected, no PCR amplified products were observed in larvae that fully fed on uninfected control mice group (Fig. 6B). These results indicate that larvae fed on LGTV-infected mice could also acquire virus from the murine host.

**Detection of transtadial transmission of LGTV from infected larval ticks to molted nymphs.** Fully fed larvae (120 h PF) repleted from LGTV-infected or uninfected control group of mice were collected and allowed to molt into nymphs. QRT-PCR analysis revealed the presence of viral loads in all individually molted nymphs (Fig. 6C). PCR products run on agarose
gel electrophoresis also confirmed the presence of amplicons corresponding to LGTV fragment of 143 bp (Fig. 6D). No PCR amplified products were observed in uninfected control group of molted ticks (Fig. 6D). These results showed that LGTV acquisition from murine host by larval ticks allows transstadial transmission of the pathogen to the molted nymphs.

**IsSMase expression is reduced in ticks that acquired LGTV loads from mice.** Our recent study showed that LGTV suppress *IsSMase* transcripts for its survival and replication in unfed and partially fed (24 h DF) nymphaal ticks. To support the use of this murine model to study changes in the arthropod gene expression during pathogen acquisition into ticks, we determined the *IsSMase* levels in partial of fully fed nymphaal/larval ticks and in freshly molted nymphs. QRT-PCR analysis showed that *IsSMase* transcript levels were significantly reduced in LGTV-infected nymphaal ticks that were collected either during feeding (48 h DF) or post-feeding (120 h PF) in comparison to the levels noted in the respective group of ticks fed on uninfected control group of mice (Fig. 7A). Also, *IsSMase* levels were significantly downregulated in larvae (120 h PF) that fed on LGTV-infected mice when compared to the levels noted in larvae fed on uninfected mice (Fig. 7B). In addition, *IsSMase* transcript levels in molted nymphs were significantly reduced in LGTV-infected larval ticks when compared to the nymphs that fed on uninfected control group of mice (Fig. 7C). Collectively, these data suggest that LGTV downregulates *IsSMase* in partial or fully fed nymphaal/larval ticks during its acquisition from murine host into ticks or in molted nymphs that acquired the pathogen loads by transstadial transmission to successfully replicate and to enhance its transport via exosome biogenesis.
Discussion

The developmental cycle of an arthropod vector from an egg through larva and/or nymph to the adult stage could take years \(^{20,31}\). Arthropod vectors such as *I. scapularis* ticks feed on the vertebrate host for about 3-5 days to acquire a complete blood meal (or full engorgement) and this extended event allows successful acquisition and/or transmission of pathogens \(^{31-33}\). The ability of ticks to transstadially transmit flaviviruses provide greater possibilities for the evolutionary changes in the virus at the phenotype and/or genotype level in the vector \(^1\).

Consequently, thoughtful investigations to study the interactions between an arthropod vector and the tick-borne flaviviruses are essential to understand an in-depth mechanism of viral pathogenesis.

To combat vector-borne viral diseases, novel methods need to be developed to address prevention and controls strategies. Development of simple approaches to infect ticks would rapidly advance investigations in learning vector-host interactions. Several studies have reported the acquisition and subsequent transmission of bacterial pathogens from the vertebrate host to ticks such as *I. scapularis*, the vector of *Borrelia burgdorferi* \(^{34,35}\), human granulocytic ehrlichiosis \(^36\), and transmission of *Anaplasma phagocytophilum* \(^37\). Michael Levin and colleagues infected the mice with *A. phagocytophilum*, and upon feeding of larval ticks they demonstrated the pathogen loads in ticks. The infected larval ticks were allowed to molt, and the nymphs were found to have maintained the bacterial loads \(^37\). In the current study, we fed the nymphal and larval ticks on LGTV-infected mice and observed that ticks acquired a significant level of viral burden that was detectable in both nymphal (partial or fully fed) and larval ticks.
Furthermore, arthropod vector such as, *Amblyomma tigrinum*, *A. ovale*, and *A. tonelliae* larvae effectively acquired the flaviviruses after feeding on viremia chicks. The technical difficulties of the previously used methods for infecting ticks, such as parenteral inoculation by microinjection and synchronous infection of ticks by immersion method, restrict key studies of the tick-virus interactions at the higher containment levels. Our previous studies have shown that synchronous infection of ticks with LGTV or *A. phagocytophilum* is possible in 60-70% of ticks; however, it would take 17 days of incubation. Also, the immersion method is arduous to assess with high priority pathogens such as POWV and TBEV due to the requirement of higher containments/facilities and the use of larger volumes of concentrated viruses (at the dose of $10^7$-$10^9$ pfu or above).

In the current study, we reported a simple method of acquisition of LGTV infection by a large number of larval or nymphal ticks feeding on murine blood. Mice administered intraperitoneal with $5 \times 10^5$ pfu of LGTV dose showed replicative viral loads on day 3 post infection, however, the viral infection persisted until day 5 p.i., in blood and peripheral tissues such as spleen, liver and skin (Figure 2). Also, we found that LGTV disseminated to the mice brain tissue, suggesting compromised blood-brain barrier (BBB) and neuroinvasion of the virus. This data suggested that inoculation with viral dose such as $5 \times 10^5$ pfu is required for LGTV replication and dissemination into mice tissues. Also, detection of viral loads in mice blood and skin indicated the LGTV spread in the murine host. The detection of higher viral loads in the peripheral tissues (such as spleen and liver) suggested enhanced replication of tick-borne viruses at these sites. The dissemination of viral loads into the brain further suggests that high viremia in blood and peripheral tissues could lead to the breach of the BBB and neuroinvasion. In addition,
detection of LGTV in partially fed nymphal ticks (collected at 48 h during feeding that acquired
the LGTV load from mice at day 3 p.i.) also demonstrated higher viral loads suggesting an early
and successful acquisition of virus into ticks. The detection of LGTV loads at 120 h post-feeding
(or day 5 p.i. for mice) of nymphal or larval ticks further suggested successful acquisition,
replication, and persistence of virus in blood-fed ticks collected at later time point of mice
infection. Transstadial transmission of LGTV from infected larval ticks to molted nymphs
further suggested that acquisition of pathogen loads by larval ticks is transmitted to the next tick
stage and is perhaps maintained in the tick body. Furthermore, down-regulation of $I_s$SMase
expression in LGTV-infected nymphs (partial or fully fed), larvae and molted nymphs suggested
pathogen influence of gene expression in all stages of ticks. Our previous study has shown that
$I_s$SMase reduced levels correlates with down regulation of its enzymatic activity and
accumulation of sphingomyelin (SM) lipid levels that may support membrane associated viral
replication and exosome biogenesis upon LGTV infection in tick cells. Taken together, we report
an appropriate, fast and efficient method to generate LGTV infected blood fed ticks (as large
batches). Compared to the other approaches such as microinjections or synchronous infections,
we believe this method is a natural way to generate large batches of LGTV-infected ticks. In
summary, this study would considerably boost investigation not only in understanding the viral
acquisition and transmission and possibly other vector-host interactions but could accelerate anti-
vector/transmission-blocking vaccine research in the field of tick-borne viral diseases.
Materials and methods

Mice and ticks. Laboratory reared *I. scapularis* ticks (larvae and nymphs) obtained from a continuously maintained colony from BEI resources/Center for Disease Control and Prevention (CDC) were used in this entire study. C57BL/6 mice (females, 6 weeks old, Charles River Laboratories, USA) were used in all animal experiments. All experiments were carried out in strict accordance with the recommendations in the Guide for the care and use of Laboratory Animals of the NIH, USA. Mice studies were performed based on animal protocol (# 18-011, PI; HS) approved by the Old Dominion University Institutional Animal Care and Use Committee (IACUC) as reported previously. Animal husbandry and administration of tranquilizer during animal experiments was performed as reported previously. We also confirm that this study is reported in accordance with ARRIVE guidelines.

Langat Virus infection and replication in mice. For the nymphal tick feeding/acquisition experiment, a total of twelve mice were clustered into four groups. Three mice were considered in each of the four groups; 1) uninfected group of mice were used to collect during feeding (DF)-group of ticks (ticks were pulled off from mice at 48 h during feeding or at 3 days post-infection, p.i. of mice,) or 2) uninfected group of mice used to collect post-feeding (PF)- group of ticks (ticks were allowed to fully engorge and replete from mice after a complete blood meal or at 5 days p.i. of mice,) or 3) LGTV-infected mice were used to collect infected-DF ticks or 4) independent LGTV-infected mice group were used to collect infected-PF ticks. For the larval tick feeding/acquisition experiment, six mice were grouped into two sub-groups (uninfected or LGTV-infected), with three mice in each sub-group. Ticks were collected at only one time point
as post feeding -PF group (these ticks were allowed to fully engorge and replete after completing a blood meal). Virus dilution was prepared from the laboratory virus stocks of 1 x 10⁹ plaques forming units (pfu/ml). Mice in the LGTV-infected group were injected intraperitoneal with 0.1 ml (500,000 pfu/ml) of diluted virus suspension in 1X PBS containing 1% gelatin (SIGMA Aldrich). After 3 or 5 days p.i., with LGTV, mice were euthanized at the given time point of tick collection as (DF)- or (PF)- groups. Murine blood and tissues (such as spleen, liver, skin and brain) were harvested for downstream processing such as RNA isolation, cDNA synthesis and Quantitative Real-Time PCR (QRT-PCR) analysis to detect viral loads.

**Feeding of nymphal ticks on LGTV-infected mice.** Naïve or uninfected nymphal ticks were fed on uninfected or LGTV-infected mice (injected intraperitoneal with 500,000 pfu/mouse) to demonstrate the virus acquisition into the tick body from the murine host. Nymphal ticks were collected at two given time points of 48 h during feeding (DF) or 120 h post-feeding (PF) for further analysis. These time points of tick collection correspond to day 3 or 5-post LGTV-infection of mice. Partially fed nymphal ticks attached to the body of the mice (from three LGTV-infected mice or respective uninfected control mice group) were pulled off with forceps during blood feeding (these ticks are referred as 48 h DF ticks). Fully fed nymphal ticks (that were allowed for completing a blood meal from the murine host are referred as 120 h PF ticks) were collected from the uninfected or LGTV-infected groups. Uninfected ticks partially or fully fed on naïve/uninfected C57BL/6 mice were used as control for both DF and PF groups of ticks.

**Feeding of larval ticks on LGTV-infected mice.** Uninfected larval ticks were fed on either naïve or LGTV-infected mice. Fed larval ticks (referred as 120 h PF) were collected after
repletion from the naïve or LGTV-infected mice for further processing. Uninfected larval ticks
fully fed on naïve mice were used as control group. After larval tick repletion (referred as 120 h
PF group), mice were euthanized, and blood/tissues (such as spleen, liver, skin and brain) were
harvested at day 5 p.i. and processed for RNA isolation, cDNA synthesis, and QRT-PCR
analysis to detect LGTV loads. Also, a large batch of larval ticks (80-100 of uninfected or
LGTV-infected ticks) were independently fed on uninfected or LGTV-infected mice (as
described previously) and fed larvae were allowed to molt into nymphal stage for 4-5 weeks.
Molting nymphs were collected and stored at -80 °C until needed for further analysis.

Isolation of total RNA, cDNA synthesis and QRT-PCR analysis in mice tissues and ticks fed
on mice. For QRT-PCR analysis, RNA was converted to complementary DNA (cDNA), which
was used to detect the presence or absence of LGTV loads and IsSMase transcript levels. Total
RNA was generated from naïve or LGTV-infected mice blood, harvested tissues (such as spleen,
liver, skin and brain), partially fed (48 h DF) or fully fed (120 h PF) nymphal or larval ticks (120
h PF) or molting nymphs by using the Aurum Total RNA Mini kit (Bio-Rad, USA) and follow
ing the manufacturer’s instructions. During RNA extractions, on-column DNaseI digestion was
performed as per the manufacturer’s recommendations. The eluted RNA was then converted to
cDNA using a cDNA synthesis kit (Bio-Rad, USA). The generated cDNA was used as a
template for the QRT-PCR reactions to analyze the viral loads or the IsSMase transcript levels.
QRT-PCR was performed using CFX96 or CFX-Opus QRT-PCR system (Bio-Rad, USA), and
iQ-SYBR Green Supermix (Bio-Rad, USA) and the isolated cDNA samples. In QRT-PCR
reactions, the quantity of mice or tick beta-actin transcripts was used to normalize the amount of
template in each reaction. To determine viral loads, levels of LGTV RNA was quantified in the
cDNA samples. The standard curve for each gene fragment was generated using ten-fold serial
dilutions starting from 1 ng to $10^{-5}$ ng of known quantities of respective fragments. For standard
preparation, initial RNA concentration was measured by taking optical density readings using a
TECAN plate reader (TECAN, USA). After measurement of concentrations, ten-fold serial
dilutions were made to prepare various standards. Oligonucleotides for mice/tick actin used in
QRT-PCR analysis are published in our previous studies 8,16. In addition, oligonucleotides used
to detect LGTV loads and IsSMase transcript levels are also published in previous studies 10,16
After completion of QRT-PCR cycles, products were analyzed on 1.2-1.5 % agarose gels
containing ethidium bromide. Internal quality control included parallel PCR amplifications of no
template control (NTC) and positive control (sequenced standard fragments) along with the
QRT-PCR reaction samples. QRT-PCR products obtained from LGTV specific oligonucleotides
from every experimental group including murine blood, tissues, fed nymphs (DF and PF) and
larvae (PF) (from both acquisition experiments) were gel purified and sequenced at the Eurofins
Genomics facility (USA).

**Statistics.** Statistical significance in the data sets was analyzed using GraphPad Prism6 software
(https://www.graphpad.com/) and Microsoft Excel 2010 (https://www.microsoft.com). For data
to compare two means, the non-paired student’s t-test was performed. P values of $< 0.05$ were
considered significant in all analyses. Statistical tests and P values used in the study are shown in
data sets.
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AUTHOR CONTRIBUTIONS

K.V.R., W.A., A.R., G.N. and H.S. performed experiments, discussed, analyzed, and interpreted the data in several settings. G.N. infected mice, and generated fed *I. scapularis* uninfected or LGTV-infected ticks. K.V.R., W.A. and A.R. performed all molecular analysis. K.V.R. and W.A. wrote paper draft, H.S. modified, revised and finalize the paper. All authors read and edited the manuscript. No conflict is declared by all authors. H.S. collected all required materials and reagents, designed and coordinated the entire study, organized all the data, compiled and supervised overall investigations.
Competing interest

No conflicts is declared by all authors.
Figure Legends

Figure 1. Schematic representation of LGTV infection in mice and acquisition of pathogen by ticks. Mice were injected intraperitoneal (i.p.,) with diluted virus containing a dose of 5x \(10^{5}\) (500,000) pfu/mouse followed by nymph or larval tick attachment and feeding. Uninfected or naïve nymphal or larval unfed ticks were allowed to attach and feed on uninfected or LGTV-infected mice. Three mice were used in each group (uninfected or LGTV-infected mice). (A) Partially fed nymphal ticks (48 h during Feeding; DF group) were pulled off from the body of mice. Mice were euthanized and blood or tissues (such as spleen, liver, skin and brain) were harvested at day 3 post-infection (p.i.,). (B) Nymphal or larval ticks were allowed to take a full blood meal or fully engorge and repleted ticks were collected as 120 h post feeding/PF group. To generate the PF group of nymphal or larval ticks we used two independent batches (one batch for nymphal and other batch for larval tick feeding) of uninfected or LGTV-infected mice (three mice in each group), respectively. Mice were euthanized at day 5 p.i. and blood or tissues (spleen, liver, skin, and brain) were harvested. Mice blood, tissues, and ticks were processed for RNA extractions, cDNA synthesis, followed by QRT-PCR analysis and DNA agarose gel electrophoresis.

Figure 2. Detection of LGTV loads in blood from mice used for nymphal/larval tick feeding

(A) QRT-PCR analysis showing LGTV loads in mice blood that were used for nymphal tick feeding. Levels of LGTV RNA were normalized to mice beta-actin levels. (B) Agarose gel electrophoresis image showing amplification of LGTV (143 bp) in cDNA samples generated from mice blood infected with LGTV. Standard used in the QRT-PCR analysis serves as a
positive control on gel. NTC is no template control, and M represents marker in bp. ns indicates no significance between the days 3 or 5 p.i. of mice. QRT-PCR analysis showing LGTV loads in blood from mice that were used for larval tick acquisition of the pathogen (C). Levels of LGTV RNA were normalized to mice beta-actin levels. (D) Agarose gel electrophoresis image showing amplification of LGTV (143 bp) in cDNA samples generated from mice blood infected with LGTV. Standard used in the QRT-PCR serves as a positive control on gel. NTC indicates no template control and M represents marker in bp or Kb.

Figure 3. LGTV disseminated from blood and replicated in mice tissues. QRT-PCR analysis showing LGTV loads in cDNA samples extracted from mice spleen, liver, skin and brain tissues collected at either day 3 p.i. (A) or at 5 days p.i. (B). Levels of LGTV RNA were normalized to mice beta-actin levels. (C-E) Agarose gel electrophoresis image showing amplification of LGTV (143 bp) in mice spleen, liver, skin and brain tissues harvested from mice at day 3 p.i. (group of mice used to generate 48 h DF ticks) or at 5 days p.i., (group of mice used to generate 120 h PF ticks). Standard 4 (C) or 3 (D) or 2 (E), used in the QRT-PCR reactions were considered as positive controls. In (E), Agarose gel electrophoresis image shows amplification of LGTV from mice tissues (spleen, liver, skin and brain) at either day 3 or 5 p.i. from mouse 3 (M3). NTC indicates no template control and M represents marker in either bp or Kb.

Figure 4. Detection of LGTV loads in tissues from mice used for larval tick feeding. (A) QRT-PCR analysis showing LGTV loads in cDNA samples from mice tissues such as spleen, liver, skin, and brain collected at 5 days p.i.. Levels of LGTV RNA were normalized to mice beta-actin levels. Agarose gel electrophoresis images showing amplification of LGTV levels
(143 bp) in mice spleen, liver, skin and brain tissues at 5 days p.i.. Data from mice 1 and 2 (M1 and M2) is shown in (B) and data from mouse 3 (M3) is shown in (C). Standard 3 or 4 used in QRT-PCR were used as positive controls on gels. NTC indicates no template control, and M represents marker in bp and Kb.

**Figure 5. Detection of LGTV loads in partial or fully fed nymphal ticks.** (A) QRT-PCR analysis showing LGTV loads in cDNA samples of nymphal ticks collected from mice at 3 days p.i., (48 h during feeding, DF or partial ticks) or at 5 days p.i., (120 h feeding, PF ticks). Levels of LGTV RNA were normalized to tick beta-actin levels. Agarose gel electrophoresis images shows amplification of LGTV (143 bp) in nymphal ticks collected at either (DF) (B) or (PF) time points (C). Standards 1 and 2 (B) or 3 and 4 (C) used in QRT-PCR served as positive controls on gels. NTC indicates no template control and M represents marker in bp and Kb.

**Figure 6. Detection of LGTV loads in larval ticks and molted nymphs.** (A) QRT-PCR analysis showing LGTV loads in cDNA samples of larval ticks collected at 5 days p.i., of mice (120 h PF ticks). (B) Agarose gel electrophoresis image showing amplification of LGTV (143 bp) in larval ticks collected at 120 h post-feeding. QRT-PCR analysis (C) or PCR amplification (D) is shown to reveal the LGTV levels in molted nymphs. Levels of LGTV RNA were normalized to tick beta-actin levels. Standards 3 and 4 (B) or 2 and 3 (D) used in QRT-PCR were used as positive controls on gels. NTC indicates no template control, and M represents marker in bp and Kb.
Figure 7. *IsSMase* expression is downregulated upon LGTV acquisition in during feeding or post-fed ticks and in molted nymphs. QRT-PCR analysis showing *IsSMase* gene expression levels in ticks collected during (48 h DF) or post-feeding (120 h PF) and freshly molted nymphs. *IsSMase* transcript loads are shown in nymphal ticks (A) (collected from 48 h DF or 120 h PF) or in larval ticks (B) (collected from 120 h PF ticks) or in molted nymphs (C). *IsSMase* mRNA levels were normalized to tick beta-actin levels. *P*-value determined by Student’s two-tailed t-test is shown.
Figures

Schematic Representation of the Experimental Plans

**A**
During Feeding (48 h DF) 
(Nymphal ticks)

- Uninfected
- Infected

500,000 viruses

Next day

Tick feeding on mice

After 48 h DF

- Partially Fed Nymphs
- Blood
- Spleen
- Liver
- Skin
- Brain

Collected ticks as 48 h DF and harvested mice tissues as 3 days p.i.

RNA extractions, cDNA synthesis, QRT-PCR and Gel Electrophoresis

**B**
Post Feeding (120 h PF) 
Nymphal and Larval ticks

- Uninfected
- Infected

500,000 viruses

Next day

Tick feeding on mice

After 120 h PF

- Fully Fed Nymphs
- Blood
- Spleen
- Liver
- Skin
- Brain
- Fully Fed Larvae

Collected ticks as 120 h PF and harvested mice tissues as 5 days p.i.

RNA extractions, cDNA synthesis, QRT-PCR and Gel Electrophoresis

Figure 1

"Please see the Manuscript PDF file for the complete figure caption".
LGTV loads in mice blood (partial or engorged nymphal and larval ticks fed on these mice)

Figure 2

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LGTV loads in mice tissues (Nymphal ticks from 48 h DF and 120 h PF fed on these mice)

Figure 3
Figure 3

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Figure 4

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Figure 5

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Figure 6

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IsSMase expression in Fed (Nymphs & larvae) or molted (Nymphs) ticks

Figure 7

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