Title: Everything you wanted to know about Mayaro virus but were afraid to ask: Characterization and lifecycle of Mayaro virus in vertebrate and invertebrate cellular backgrounds

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Abstract

Mayaro virus (MAYV) is an emerging new world alphavirus that causes acute multiphasic febrile illness, skin rash, polyarthritis, and occasional severe clinical phenotypes. It is a member of the genus Alphavirus and family Togaviridae. The virus lifecycle alternates between invertebrate and vertebrate hosts. In this present report the replication features, cell entry, life cycle and virus-related cell pathology of MAYV were characterized using vertebrate and invertebrate in vitro models. Electron dense clathrin-coated pits in infected cells, and reduced viral production in the presence of dynasore, ammonium chloride, and bafilomycin, indicates that viral entry occurs through pH-dependent endocytosis. Increase in FITC-dextran uptake (an indicator of macropinocytosis) in MAYV-infected cells, and dose-dependent infection inhibition by 5-(N-ethyl-N-isopropyl) amiloride (a macropinocytosis inhibitor), indicated that macropinocytosis was an additional entry mechanism of MAYV in vertebrate cells. Acutely infected vertebrate and invertebrate cells formed cytoplasmic or membrane-associated extracytoplasmic replication complexes. Mosquito cells showed modified hybrid cytoplasmic vesicles that supported virus replication, nucleocapsid production, and maturation. The mature virus particles were released from cells by both exocytosis and budding from the cell membrane. MAYV replication was cytopathic and associated with induction of apoptosis by the intrinsic pathway and later by the extrinsic pathway in infected vertebrate cells. In summary, this study provides a comprehensive overview of the characterization and lifecycle of the emerging pathogen MAYV, in both the vertebrate and invertebrate host cellular backgrounds.
Introduction

Mayaro virus (MAYV) is a neglected emerging arboviral pathogen. It was first isolated in 1954 from Trinidad and Tobago, and since then, several outbreaks have been reported in South and Central America. Many of the clinical features, including arthralgia, overlap with Dengue and Chikungunya; however, biphasic or intermittent hyperthermia can distinguish MAYV from other arboviral infections. MAYV can cause neurological complications, myocarditis, hemorrhagic manifestations, and even death. MAYV alternates between vertebrate and invertebrate hosts, and is primarily transmitted through the bite of female Haemagogus (in sylvatic cycle) and Aedes (urban and peri-urban cycle) mosquito species in South and Central America. Transmission by multiple Anopheline mosquito species has also been demonstrated through laboratory studies, indicating a potential risk of this emerging virus in other parts of the world.

MAYV is a positive-sense, single-stranded-RNA virus that belongs to the genus Alphavirus in the family Togaviridae. It is a member of the Semliki Forest virus antigenic complex that consists of eight other viruses: Semliki Forest, Chikungunya (CHIKV), Bebaru, Getah, Ross River (RRV), O’nyong-nyong (ONNV), Sagiyama and Una viruses. Its genome is approximately 11.7 kb and encodes four nonstructural proteins (NSP1-4), six structural proteins (capsid [C], envelope [E] proteins, and trans-frame), and two open reading frames (ORFs). Based on its whole-genome phylogeny, MAYV has three genotypes (D, L, and N) which are highly conserved with approximately 17% nucleotide divergence across all three genotypes, and 4% among D strains. It is thought that genotypes D and L diverged approximately 150 years ago, and genotype N...
diverged approximately 250 years ago 11. Genotype D has a diverse distribution in South America and the Caribbean, genotype L was detected in certain parts of Brazil whereas N genotype was found only in a localized region in Peru 12. Further, the ability of MAYV to recombine with other strains and related viruses may arise in new lineages 34.

Most of the cellular and molecular mechanisms related to MAYV is based on studies with other alphaviruses. To bridge this critical knowledge gap on the biology of this new world emerging, yet neglected, arthritis causing alphavirus we used mosquito and vertebrate cells to characterize its biology. This study provides a comprehensive investigation on the lifecycle of MAYV, its replication characteristics, and cellular tropism to provide insight into the interaction of MAYV with its mosquito and human host. Given MAYV is expanding its geographical existence it is a potential public health problem this study will lay down the foundation of biological understanding valuable for therapeutic and preventive interventions.

Results

Growth kinetics of MAYV in vertebrate and invertebrate cells. To determine the in vitro host range, growth kinetics, production of infectious viral particles, and cytopathology of MAYV, single- and multi-step growth curve analyses were performed in mammalian and insect cells (Fig.2A). The low MOI (0.1) growth curve demonstrates the release of infectious viruses in two-bursts. The first burst appears between 6-12 hours post infection (hpi) in mammalian cells and after 12hpi in mosquito cells. The second burst appears approximately at 24hpi in Vero and BHK-21 cell and after 30hpi for Huh
7.5 and mosquito cells. It is also important to note that, except for BHK-21, other vertebrate and invertebrate cells showed a distinctive latent, exponential, and logarithmic phase.

In all tested cell lines, MAYV replicated to high titers (>10⁷ FFU/mL) with the exception of the Aag2 cell line (from Aedes aegypti) where the maximum titer was 10⁶ FFU/ml. BHK-21 produced almost 1.5-log higher viral titer in comparison to other mammalian cells, and Sua5b produced more than 2-log higher virus titer at 24hpi and 48hpi. All mammalian cells caused cytopathic effects detectable after 12hpi which was characterized by cell rounding and detachment. Conversely, no obvious cytopathic effect was detected in mosquito cell lines.

Physical characterization of MAYV. To understand and compare the thermal stability of MAYV to other alphaviruses such as Sindbis virus (SINV) and ONNV, heat inactivation kinetics under different temperature conditions was investigated. Viral particles produced in Vero cells were treated at temperatures ranging from 25 to 55 °C for 3h, and virus infectivity was determined by focus forming assay. MAYV displayed higher thermal stability at 50°C compared to the ONNV, but lower than SINV (Fig. 2B). After 3h of incubation at 55°C all viruses were completely inactivated.

Plaque size is a measure of viral replication efficiency and genetic heterogeneity among the virus population. MAYV plaque morphology was examined in Vero cells at 72h post infection and compared to ONNV and SINV plaques. MAYV formed large (~0.8mm) and
small (~0.5 mm) plaques, while the other alphaviruses had only one type of plaque diameter (small for SINV and large for ONNV) (Fig. 2C). The large MAYV plaques were morphologically similar to ONNV, with a distinct and prominent outline. Conversely, SINV plaques were smaller, hazily outlined and less prominent.

**pH-dependent receptor-mediated endocytic entry of MAYV.** Electron dense cup-shaped structures (or pits) are prominent features during the entry phase of MAYV, as seen in electron micrographs (see below). In the RME process, these pits eventually mature and form the early endosome. At a later stage, entry features a pH-dependent fusion of viral envelope with the endocytic vesicle and release of the viral genome into the cytoplasm.

To assess the role of RME in the entry of MAYV, we used dynasore, a pharmacological dynamin GTPase inhibitor. Dynamin is a GTPase essential for pinching off of the endosomes from the cytoplasmic membrane. Vertebrate and invertebrate cells were pretreated (1h pre-infection) with dynasore and infected with MAYV. MAYV internalization was significantly inhibited by dynasore pretreatment and its presence in a dose-dependent manner indicating the role of RME in the MAYV entry process (Fig. 2D).

Further, to test whether the release of MAYV into the host cytoplasm upon entry is pH-dependent, we evaluated the effect of two known pharmacological inhibitors: Ammonium chloride and Bafilomycin. The former is a weak base and a lysosomotropic agent because of its propensity to accumulate in lysosomes, and the latter blocks the v type ATPase and alters the late endosomal vesicles' pH. Pretreatment of cells (1h pre-infection) and maintaining the chemicals in the culture medium with increasing
concentrations revealed a significant dose-dependent reduction in virus particle production (Fig. 2D).

**Spatial distribution of viral replication complex and viral particles.** To demonstrate the intracellular spatial distribution pattern of the viral replication complex, Huh-7.5 and C6/36 cells were infected with MAYV and assayed with dsRNA antibodies 6hpi. dsRNA is an intermediate in the replication complex of RNA viruses which was found to be distributed throughout the cytoplasmic compartment of both cell lines but absent in the filopodial extensions (detected using phallloidin). Z-stack 3D analysis revealed that replication units (dsRNA puncta) were distributed both in cytoplasm and cell membrane indicated by the distribution of dsRNA puncta both in and above the plane of the nucleus (Fig. 3A, 3B). This corroborates with the distribution of replication vesicles both in the cytoplasm and plasma membrane, as seen under TEM, and is described later.

To detect the induction of filopodial nanofiber-like extensions in MAYV infected cells and its ability to infect neighboring cells, a confocal immunofluorescence analysis was performed. Phallloidin and a CHIKV cross-reactive antibody were used to detect F-actin and MAYV E2 glycoprotein, respectively, in Huh-7.5, and C6/36 mosquito cell lines 24hpi MAYV infection. Both cell types exhibited 2-3 nanofiber like extensions per cell that connected to neighboring cells with a high density of viral particles at the surface of the membrane (Fig. 3C). MAYV E2 glycoproteins were primarily localized at the cell membrane and throughout the cytoplasm. Nevertheless, partial co-localization between E2 and actin was found in discrete areas. This indicated that MAYV might be able to transmit from cell-to-cell via filopodia.
**MAYV utilizes macropinocytosis for its entry.** Filopodial cup-like extensions engulfing virus particles were spotted during the ultrastructural study in both BHK-21 and C6/36 cells, suggesting a process of macropinocytosis (Fig. 4A). To assess the involvement of macropinocytosis in MAYV entry, a functional fluid uptake assay was performed. Both vertebrate (BHK-21 and Huh-7.5) and invertebrate (C6/36 and Aag2) cell lines were incubated with 1uM FITC-dextran, and intake of FITC labeled dextran evaluated by detection of fluorescence signal in the presence or absence of MAYV. An increased FITC-dextran uptake was noted in vertebrate cells infected with MAYV, but no changes were detected in mosquito cells (Fig. 4B).

For further validation, a quantitative MAYV infection assay was performed in the presence of EIPA, a pharmacological inhibitor of macropinocytosis that inhibits Na+/H+ exchange. A dose-dependent reduction in infection in both BHK-21 and Huh-7.5 cells was observed, but not was not observed in mosquito cells, reinforcing the role of macropinocytosis as an important entry pathway of MAYV in vertebrate cells (Fig. 4C).

**Ultrastructural analysis of BHK-21 and C6/36 cells infected with MAYV.** To provide ultrastructural details, thin-section transmission electron microscopy of BHK-21 and C6/36 cells infected with MAYV was performed (Fig. 5A). In BHK-21 cells, it was observed that MAYV replicated and matured in replication spherules, cytoplasmic vesicles (CPV) and egressed by both budding and exocytosis. The inner membrane of CPV-I, associated with bulb-shaped spherules, corresponds to the invaginations of the vacuole membranes. Inside the spherule, a central dense mass of possibly replicating viral RNA was often seen with a narrow neck connected it to the cytoplasm. Spherules
protruding toward extracellular space were also observed. Nucleocapsids (NCs) are associated with type II cytoplasmic vesicles (CPV-II) containing multivesicular inclusions, amorphous material, and occasional intact-looking virions. Different types of CPV-II were seen coupled with NCs on the cytoplasmic side of the vacuole and/or the interior of the double-membrane CPV-II. CPV-I and CPV-II were both present in the infected cells 6hpi, though CPV-II was more abundant 12hpi viral budding occurred predominantly from PM.

In C6/36 cells, viral entry through RME was seen (Fig. 5B). Furthermore, similar to the infection of BHK-21 cells, replication spherules, NC-containing vesicles, budding virus, and exocytic vesicles carrying mature viral particles were detected in the infected mosquito cells. In contrast to the series of membrane attached spherules facing inside the CPV-I of BHK-21 cells, in C6/36 cells most of the clustered spherules were unattached to the vesicular membrane. NCs were also observed inside and nearby the CVP-II membrane, demonstrating that MAYV maturation and replication occur in the same cellular space. Intermediary CPV-I and CPV-II replication spherules and internally budded viral particles were observed in both early and late phases of infection. These modified membrane structures were also seen near the rough endoplasmic reticulum and Golgi complexes. Furthermore, NCs close to the PM for budding and internally budded mature viral particles inside secretory intraluminal vesicles were seen.

**MAYV activates both mitochondrial dependent and independent apoptosis.** MAYV replication in Vero, BHK-21, and Huh-7.5 cells resulted in cytopathological changes characterized by cellular fusion and multinucleated giant syncytia, detachment of
infected cells, and eventually cell lysis and death. To assess the involvement of MAYV in the apoptosis pathway (Huh7.5 cells), a cell viability test, and a PARP and Caspase detection study were performed.

Cell viability of MAYV infected Huh-7.5 cells was quantitatively measured through neutral red uptake assay. Neutral red is a eurhodin dye, actively transported into live cells, which stains lysosomes and is subsequently measured to determine cell viability. Cell viability was reduced up to ~40% at 24hpi and more than 90% at 48hpi. A series of morphological changes, including cytoplasmic blebbing, mitochondrial swelling, chromatin condensation, and nuclear fragmentation, were observed in the ultrastructural analysis of MAYV infected cells (Fig. 6A, 6B).

Poly (ADP-ribose) polymerase (PARP) is an enzyme that suppresses nuclear fragmentation and apoptotic body formation. Under cellular stress such as viral infection, PARP is cleaved and compromises the cellular viability. Cleaved PARP, a marker of cells undergoing apoptosis, is detected at 12hpi of MAYV and foremost at 24hpi in the immunoblot of cells infected with MAYV. Additionally, stained cells with Hoechst 33342 revealed that MAYV induced chromatin condensation, nuclear fragmentation at 12hpi and is florid around 36hpi, correlating with the PARP immunoblot results (Fig. 6C).

Caspase-9, an intrinsic pathway marker, and caspase-8, which initiates the extrinsic pathway, were detected in MAYV infected cells through immunostaining at 12hpi and 24hpi, respectively. Furthermore, active caspase-3, which cleaves the PARP enzyme and culminates in nuclear fragmentation and cell death, was detected at 12hpi in the
immunoblot. This suggests that apoptosis which is led by the extrinsic pathway due to MAYV infection is subsequent to the early intrinsic pathway (Fig. 6D).

Finally, to test the hypothesis that MAYV utilizes the apoptosis pathway to maximize its spread, a co-culture experiment was designed. MAYV Huh7.5 infected cells pretreated with caspase inhibitor (ZVAD-FMK) were co-cultured with Vero cells that served as the infection center. The focus size formed by the caspase inhibitor treated cells was reduced up to 50% compared to the control group indicating that MAYV utilizes the apoptosis process to maximize the infection process (Fig. 6E).

Discussion

Like other members of alphaviruses, the life cycle of MAYV is rapid and newly infectious viruses can be detected in the culture supernatant as early as 3-5hpi in vertebrate cells and 7-9hpi in mosquito cells 15. The viral titers are similar in mosquito and vertebrate cells at 24hpi and the viral titers in supernatants reach 10^7–10^8 FFU/ml within 48h, depending on the cell types. Further in agreement with previous reports of transmission of MAYV by Anopheles mosquitoes 8, Sua5b, a cell line derived from Anopheles gambiae, supported efficient replication and produced high viral titer similar to Aedes albopictus C6/36 cells. The low viral titer in the Aag2 cells compared to other insect cells and early plateau (36hpi) is not surprising, as Aag2 cells are persistently infected with insect-specific bunyavirus Phasi Charoen-like virus and possibly interfered with the replication of MAYV 16. MAYV had also been reported to replicate to high titer in avian cell lines which migratory birds has been hypothesized to be its reservoir 17.
Serological diagnosis is critical not only for disease surveillance but also for bedside diagnostics and is preferred over molecular assays especially for resource limited setup. The gold standard for serological diagnosis is the plaque reduction neutralization test, which demonstrates the virus neutralizing capacity of serum samples. Before any tests, the serum samples are heat inactivated for 30 minutes at 56°C to inactivate complement proteins and any infectious viral particles, if there are any. Sometimes the inactivation time needs to be prolonged (e.g. 60 minutes for Western equine encephalitis virus and CHIKV) ensuring the sensitivity of the assays and the safety of laboratory personnel. We tested and compared the thermal stability of MAYV along with SINV and ONNV. After 3h of incubation at 45°C some of the virus particles were still infective but when the temperature increased to 55°C all of them lost their infectivity. The complete inactivation of MAYV could be achieved after 60 minutes of incubation at 55°C.

*In vitro* characterization of MAYV showed overt cytopathic effects in vertebrate cells, which facilitated the development of plaque assay. The viral strain used in this study forms a mixture of large and small plaques with clear and sharp boundaries demonstrating the presence of a swarm of viral population with different viral entry and replication capacities. In contrast, mosquito cells did not show any cytopathic effect, were chronically infected with MAYV, and continuously shed infectious viral particles. This demonstrates that MAYV has developed a delicate balance between the mammalian host and mosquito host that allows its persistent survival in nature.

The role of RME has been established as one on the major route of entry for alphaviruses. As the endosome migrates closer to the nucleus, the inner
compartment becomes acidic. This acidification causes the glycoproteins on the virus surface to undergo conformational changes and fuse with the endosomal membrane and releases the viral genome into the cytoplasm \(^{21}\).

Mxra8, an adhesion molecule primarily expressed on epithelial myeloid, and mesenchymal cells has been recently established as a receptor for CHIKV, ONNV, RRV and MAYV \(^{22}\). Electron dense clathrin-coated pits in the MAYV infected cells and significant reduction of viral titers both in the vertebrate and invertebrate cells upon chemical inhibition of dynamin-2, a key protein required for the formation of clathrin-coated pits and vesicles, suggested the role of RME in MAYV entry \(^{23}\). Further the use of NH\(_4\)Cl and Bafilomycin that alters the pH of the early and late endosomal vesicles, inhibited the release of viral genome into the cytoplasm and infection process in turn and production of infectious virus \(^{24}\). This indicates that the release of MAYV genome is mediated through pH-dependent RME.

Upon release of viral RNA into the host cytoplasm which has same polarity as that of cellular mRNA, it undergoes several rounds of translational events using host cellular machinery generating viral polyproteins. These are further processed by cellular and newly synthesized viral proteases creating the viral replication complex. The nonstructural proteins of alphaviruses induce intracellular membrane remodeling that results in the appearance of cytopathic vacuoles (CPVs) and has been greatly studied using the SFV model. The CPVs are of two types, namely, CPV type I (CPV-I) and CPV type II (CPV-II) \(^{25}\). Double labeling of organelle and viral nonstructural proteins showed that CPV-I are derivatives of late endosomes and lysosomes, while monensin treatment results in accumulation of E1/E2 glycoproteins of SFV in the trans-Golgi network (TGN),
indicating TGN origin of CPV-II 26, 27. The replication complexes are strategically concentrated in CPV-I and replication spherule for efficient viral genome replication and to escape the cellular antiviral response 28. These replication spherules that has access to the host cellular raw materials through an opening toward the cytoplasm were seen either inside CPV-I or transported to the plasma membrane 29. Both vertebrate and invertebrate cells had similar spherules with a central electron dense material possibly the MAYV replicating RNA. dsRNAs are unique to viral infected cells and are the markers of replication intermediates/replication complex 30. In the confocal image detection of dsRNA in plane with the nucleus indicated the replication of MAYV in the cytoplasmic CPV-I, which appears during the early stage of infection and above the plane of the nucleus indicating the presence of replication complex on the cell membrane that is further validated in the electron micrographs. Like other alphaviruses, CPV-II was the predominant vacuolar structure in the later stage of MAYV infection both in BHK-21 and C6/36 cells. Our electron micrograph analysis showed mainly three different populations of CPV-II in BHK-21 cells. The first population consists of numerous nucleocapsids (NCs) attached to the cytoplasmic face of membranes, second has NCs enclosed inside the vesicles, and the third were transporting vesicles containing E/E2 viral glycoproteins from the TGN to the viral budding sites on the plasma membrane. In contrast to previous reports NCs were seen in the cytoplasm close to the plasma membrane of infected BHK-21 and C6/36 cells. The assembly of NCs as matured viruses was observed to take place either by a budding process from the plasma membrane into the extracellular space or mature inside the CPVs transported through exocytosis pathway. The C6/36 cells had additional hybrid vesicles
containing both replication spherules and NCs, also having vesicles with mature virus particles.

Arthritogenic alphaviruses, including MAYV, have been associated with inflammation of the joints and often with other tissues. Inflammation and apoptosis go in parallel with disease severity. Although apoptosis eliminates infected cells, viruses have evolved to manipulate this cellular antiviral mechanism for maximizing the production of progeny virus and their spread to maintain a prolonged and high viremia in vertebrate host, a property that is relevant especially for arboviruses for their maintenance in the natural transmission cycle. CHIKV and SINV camouflage in apoptotic blebs to facilitate infection of neighboring cells. Apoptotic blebs bodies are quite evident in the MAYV infected cells. Most commonly blebs are seen during apoptosis and contains part of the cytoplasm (~2%) with or without organellar fragments for recycling by the phagocytic cells35,36. We designed a coculture-infection center assay that produced smaller diameter focus size in the presence of apoptosis inhibitor ZVAD-FMK that inhibits apoptosis at an early stage indicating the role of apoptosis in the spread of MAYV infection. Up-regulation of proapoptotic proteins or down-regulation of antiapoptotic proteins can alter mitochondrial membrane permeability that can promote release of cytochrome c31. Cytochrome c interacts with Apaf-1 and caspase-9 and forms a multiprotein complex apoptosome, leading to activation of caspase-3 and then apoptosis32,33. Increased activation of caspase 9 during MAYV infection indicates the involvement of mitochondrial intrinsic pathway. Further detection of active caspase 8, possibly activated by secretion of cellular or virus induced death signal by cell membrane death receptors,
demonstrate that the MAYV induced apoptosis is triggered through both intrinsic and extrinsic pathways.

In conclusion, we present a comprehensive in vitro characterization of MAYV in both vertebrate host and invertebrate mosquito vector cells. Besides pH-dependent receptor mediated endocytosis (RME) MAYV uses macropinocytosis for entry into vertebrate cells. During acute infection in vertebrate cells MAYV achieves very high titer and utilizes the apoptotic pathway for efficient transmission to neighboring cells. In mosquito cells a burst in high viral titer that follows continuous virus production at reduced levels, indicating the evolutionary adaptation of arboviruses with their mosquito vector. Further it will be interesting to understand and identify the molecular signature(s) that makes them clinically different from their close geographical ancestors.

Methods

**Cell culture.** C6/36 (*Aedes albopictus*) cells, Aag2 (*Aedes aegypti*), and Sua5b (*Anopheles gambiae*) cells were maintained in Schneider’s insect cell culture medium. Vero (African green monkey kidney), BHK-21 (Baby Hamster Kidney) and Huh7.5 (Human liver) cells were maintained in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 50 units ml⁻¹ of penicillin and 50 µg ml⁻¹ of streptomycin. Vertebrate cells were cultured in a 37 °C incubator with 5% CO₂, while invertebrate (mosquito) cells were cultured at 28 °C incubator without CO₂.

**Viral growth kinetics by focus forming assay.** Approximately $3 \times 10^4$ Vero cells per well were seeded in a 96-well plate and incubated overnight. Virus samples were diluted in tenfold serial dilutions in DMEM without FBS supplemented with antibiotics; 30 µl
from each dilution was added to each well containing the cells and incubated at 37°C. One hour post infection, 100 µl of overlay media (1X DMEM medium, 10% FBS, 50 units ml⁻¹ penicillin, 50 µg ml⁻¹ streptomycin, 1% carboxymethyl cellulose) was added. Twenty-four hours post-infection, the overlay was removed; to fix the cell, 100 µl of 4% paraformaldehyde in phosphate buffered saline (PBS) was added and incubated at RT for 15 minutes. For antibody probing, the plate was first blocked for 30 minutes at RT with 50 µl of blocking solution (3% BSA, 0.25% triton x in PBS), followed by the addition of 30 µl of primary antibody (CHIK-48, anti-E1 protein) for 2 h at RT or overnight at 4°C. Plates were then washed three times with PBS. Secondary antibody (Alexa 488 Goat anti-Mouse, ThermoScientific) was added (30 µl per well) and incubated for 1 h at RT. Plates were washed three times with distilled water, air-dried and screened manually under 4x objective of Olympus microscope. Viral titers were expressed as FFU ml⁻¹.

**Heat treatment of virus particles.** Approximately 10⁶ FFU of MAYV, SINV and ONNV were subjected to a thermal gradient treatment from 30 to 60 °C for 3 h with a thermocycler (Bio-Rad T100 Thermal Cycler), after which samples were immediately titrated on Vero cells. A non-heat-treated virus control kept at 4°C for 3 h was also included. The ratio of the number of FFU in heat-treated versus the non-heat-treated viruses was calculated to determine the relative infectivity.

**Plaque assay:** Vero cells (5×10⁵ cells/well) were grown overnight to a confluent monolayer in 6 well plates and infected with serial dilutions of MAYV or ONNV or SINV-infected culture supernatant. Virus were allowed to adsorb on the cell surface for 1 hour at 37°C with 5% CO₂; subsequently monolayers were rinsed with DMEM without FBS
and overlay medium (1% methylcellulose in DMEM with 5% FBS) was added. The plates were incubated at 37 °C in a 5% CO₂ incubator for 72 h. At the end of the incubation period, overlay media was removed and fixed with 1 ml of 4% PFA solution at RT for 15 minutes. After one washing with PBS, methylene blue prepared in methanol was added onto the fixed cells. After 30 minutes of incubation, plates were cleaned in tap water, and plaques size were measured.

**Effect of lysosomotropic drug on MAYV entry:** BHK-21 (3X10⁴ cells/well) and Huh7.5 and C6/36 (5X10⁴ cells/well) cells were seeded in 96 well plates, the day before treatment. Cells were pretreated for 3h in serum-free media containing mentioned concentrations of ammonium chloride or bafilomycin A1. Following incubation, cells were infected with each virus at an MOI of 1 in the presence of each compound for 1 h at 37°C. Cells were washed, complete media containing each compound was added, and were incubated at 37°C for 16h. Additionally, no drug and drug with no virus control were included. After incubation, cells were fixed with 4% PFA, permeabilized, processed for immunofluorescence using virus-specific antibodies, and virus-positive cells were quantified.

**Immunofluorescence analysis of infected cells.** Huh7.5 or C6/36 cells were seeded in two well chamber slides at a density of 2 × 10⁵ cells per well. Cells were then infected with MAYV at a MOI of 1 or mock-infected and incubated at 37 °C (Huh7.5) or 28 °C (C6/36). After 1 h, cells were washed with DMEM or Schneider’s insect cell culture medium without FBS and replaced with fresh growth medium and then incubated at 37 °C (Huh7.5) or 28 °C (C6/36). Cells were fixed with 4% PFA for 15 minutes at RT at 12hpi for detection of dsRNA or at 24hpi for E1 protein detection. Cells were then
blocked with 500µl of blocking solution (3% BSA with 0.25% tritonx in PBS) for 30 min.
at room temperature before incubation with CHIK-48 (anti-E2) (1:500) or dsRNA
antibody (1:100) diluted in blocking buffer overnight. Next day cells were washed three
times with PBS/T and incubated with Alexa Fluor 488 or 595 secondary antibody (Life
Technologies) diluted in PBS/T for 1 h at RT. After incubation, cells were washed three
times with PBS/T. Finally, for nuclear staining, cells were incubated with 500 ul of PBS
with Hoechst stain for 2 minutes, followed by a final washing with distilled water. Cells
were mounted using 1.5mm cover glass with ProLong Diamond Antifade Mountant (Life
Technologies). Images were taken using a Zeiss LSM 800 confocal microscope.

**TEM of infected cells.** Cell pellets were fixed in 3% glutaraldehyde in PBS for 1 h at
room temperature, washed with 0.1 M cacodylate buffer, and incubated in 1% OsO4 (in
0.1 M cacodylate buffer) for 40 min at room temperature. Samples were then washed
once with 0.1 M cacodylate buffer and once in 80% acetone for a further incubation
overnight at 4 °C in 2% uranyl acetate/80% acetone. The following day, serial
dehydration and resin infiltration steps were performed as follows: 2 × 10 min with 80%
acetone, 2 × 10 min with 90% acetone, 3 × 20 min with 100% acetone, 1 × 90 min with
50% Epon/50% acetone, 1 × 90 min with 75% Epon/25% acetone and 1 × 90 min with
100% Epon. Epon was replaced by fresh 100% Epon with polymerization accelerator
BDMA and embedded at 65 °C for 72 h. Resin blocks were then trimmed using a
DiATOME Ultra Diamond Knife on a Leica EM UC7 ultramicrotome, from which 50nm
sections were obtained and mounted on EM copper grids with carbon coating. Sections
were post-stained in 2% uranyl acetate in water and Reynolds’ lead citrate for 1 min
each and then processed for TEM imaging using either a FEI Tecnai F20 S/TEM electron microscope.

**Cell cytotoxicity and viability assay.** Neutral red uptake assay was used to evaluate the MAYV induced cell cytotoxicity/death and cytotoxicity of chemicals in Huh7.5 cells. In brief, Huh7.5 cells were seeded into 96-well plates at a density of $2.5 \times 10^4$ cells per well and allowed to attach for 12 h. The outer perimeter wells of the plate were left blank as these wells have been reported with a decreased cell growth. Cells were then infected with 0.1 or 1 moi of MAYV at different time points in order to harvest the plates at once for 3,6,12,24 and 48 h of post infection. Cell plates were washed once and replenished with 100ul of neutral red medium (40 ug ml$^{-1}$) which was prepared a day before and incubated at 37°C. After 2 h of incubation at 37 degree, plates were washed (with PBS) and 150 ul of neutral red destain solution (50% ethanol (96%), 49% deionized water, 1% glacial acetic acid) was added to each well. To extract the neutral red from the cells, plates were agitated for 10 minutes on a microtiter plate shaker. Optical density of the plates was measured at 540 nm in a microtiter plate reader spectrophotometer. Each plate had blanks which contain no cell reference and cells without virus as no treatment reference. Cell cytotoxicity was measured using the following formula: $\%$ viable cells = $(\text{Abs}_{\text{samp}} - \text{Abs}_{\text{blank}}) / (\text{Abs}_{\text{control}} - \text{Abs}_{\text{blank}}) \times 100$.

**Western blot:** MAYV infected and mock infected cells were harvested at the time points stated above by a cell scraper and pelleted by centrifugation. Cell pellets were washed twice with PBS and lysed in RIPA buffer (20 minutes on ice) with protease inhibitor cocktail. The lysate was cleared by centrifugation at 14,000 rpm for 20 min at 4°C and resolved on 10% or 12% sodium dodecyl sulfate polyacrylamide gel electrophoresis.
Proteins were transferred to nitrocellulose membranes (0.45, Bio-Rad). Membranes were blocked in 5% milk in TBS-tween 20 (50 mM Tris-HCl pH 7.4, 250 mM NaCl, 0.1% Tween-20) for 1 hour. Membranes were probed with primary antibody overnight at 4°C, then with corresponding HRP-conjugated secondary antibodies in 5% milk/TBS-tween. Signals were detected with the enhanced chemiluminescence method (GE healthcare) or AP-based colorimetric kit (Bio-Rad).

Infection Center assay
At 6hpi Huh7.5 cells were washed, suspended with the aid of trypsin, centrifuged, and resuspended in DMEM supplemented with ZVAD-FMK or DMSO. The infected cells were counted and diluted. Then 0.03 ml of the infected cell dilution was added onto the Vero cell monolayer prepared on the 96 well culture plate. To permit the infected cells to settle down on the Vero cells, the medium was removed after 2 hours of incubation, and 100ul of methylcellulose overlay supplemented with ZVAD-FMK or DMSO was added. At 24hpi, cells were fixed and processed as described for focus forming assay and pictured using an epifluorescence microscope. The diameters of the foci were measured using ImageJ with arbitrary units.

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References


Figure Legends:

Fig. 1: Distribution of D (yellow) and L (green) strains of Mayaro virus in South America and geographical areas at risk of future MAYV outbreaks highlighted in blue. The map was generated based on the literature (Sinka et al 2012, Powers et al 2006 and Brustolin et al 2018).

Fig. 2: Growth kinetics, thermostability, plaque morphology and cellular entry of Mayaro virus. A) Vertebrate (Huh7.5; Human, Vero; Monkey and BHK-21; Hamster) and invertebrate (Ae. albopictus; C6/36, Ae. aegypti; Aag2 and An. gambiae; Sua5b) cells were infected with MAYV at an MOI of 0.1 and 1. Culture supernatants were harvested every 2-hour intervals up to 12-hour post infection, then at every 6-hour up to 36-hour post infection and a final sample was harvested at 48-hour post infection. Viral titers were quantified in the harvested samples using focus forming assay using Vero cells and growth curves were plotted. B) Mayaro, Sindbis and O'nyong'nyong virus thermostability. Stocks of indicated viruses were incubated at 4, 25, 30, 35, 40, 45, 50 and 55°C for 180 minutes and the number of infectious particles determined by focus forming assay C) Known titer stocks of MAYV, ONNV and SINV (15-20 PFU/100ul) were used to produce plaques in 6 well plates on Vero cell monolayer as detailed in the method section. The plaque diameters were measured. D) Effect of Dynasore, NH4Cl and Bafilomycin on MAYV entry process. Huh7.5 and C6/36 cells were treated before and after infection with the indicated doses of the chemicals. MAYV was adsorbed to cells at 1 MOI for one hour and non-adsorbed virus was removed, virus in the culture
supernatant were collected at 24hpi for Huh7.5 and 36hpi for C6/36 and titrated. Data from three independent experiments were plotted, error bars indicate standard deviations of the means.

Fig. 3: Spatial distribution of MAYV and its replication intermediate (dsRNA): A) Huh7.5 (upper panel) and C6/36 (lower panel) cells were infected with 1 MOI of MAYV, 6hpi cells were fixed and stained for dsRNA (green), filamentous actin (F-actin) filament with phalloidin (red) and nucleus (blue) with Hoechst stain. B) Z-stack analysis shows viral dsRNA (replication complex) are localized above and on same plane with reference to nucleus. C) MAYV on the intercellular extensions: Huh7.5 (upper panel) and C6/36 (lower panel) cells were infected with MAYV, incubated at 37°C for 12 h for Huh7.5 and 24h for C6/36 cells, and fixed. Cells were permeabilized and stained for viral E2 envelope protein (green) and phalloidin to detect F-actin (red) and nucleus (blue) with Hoechst stain. Images from one optical section are shown and are representative of three independent experiments.

Fig. 4: Macropinocytosis as an additional entry mechanism for MAYV in vertebrate hosts. A) Macropinocytic membrane projections chasing virus particles. BHK-21 and C6/36 cells infected with MAYV at 5 MOI; 3hpi cells were harvested, processed, sectioned and scanned under TEM. B) MAYV enhance FITC-dextran uptake in vertebrate cells. BHK-21, Huh7.5, C6/36 cells were pre-treated with MAYV (lower panel) at MOI of 1 or mock infected for one hour. Cells were then
washed and incubated with medium containing FITC-labeled dextran 10,000 MW (1 mg/ml). After 20 minutes, cells were washed, fixed and imaged. C) EIPA, macropinocytosis inhibitor inhibits infection of MAYV in vertebrate cells. Indicated vertebrate and invertebrate cells were pre-treated with the different concentrations of EIPA, followed by incubation with MAYV at MOI of 0.1 in the continued presence of the drug. Control cells received DMSO instead of the drug. Cells were washed and fixed and probed with appropriate primary and secondary antibodies after 12hpi for vertebrate cells and 24hpi for invertebrate cells. Percent virus infection was calculated and plotted as bar diagram.

Fig. 5: Electron microscopy analysis of MAYV lifecycle in vertebrate and invertebrate cells. A) BHK-21 cells infected with MAYV at an MOI of 5, fixed and sectioned for TEM analysis. i) and ii) Virus replication inside the replication spherules in the cytopathic vesicle and on the cell membrane. iii) and iv) formation and maturation of nuclear core (NCs) in cytoplasm. v) and vi) release of viral particles by membrane budding and exocytosis. RS=Replication Spherules; CPV: Cytopathic Vacuoles. B) Electron microscopy analysis of MAYV lifecycle in C6/36 cells. C6/36 cells were infected with Mayaro virus at an MOI of 5 and fixed for TEM analysis. i) Virus entry via clathrin coated pits like vesicle. ii) and iii) Virus replication inside the replication spherules in the cytopathic vesicle and on the cell membrane. ii) and vi) Virus replication and maturation in cytopathic vesicle I and II hybrid vesicles. v) formation and maturation of nuclear core (NCs) in
cytoplasm. Release of viral particles by vii) membrane budding and viii) exocytosis. RS=Replication Spherules; CPV: Cytopathic Vacuoles

Fig.6: Both intrinsic and extrinsic apoptotic pathways are activated during MAYV infection. A) At 3,6,9,12,24,36 and 48hpi Huh7.5 cell viability was measured by neutral red uptake assay. Data from three independent experiments were plotted, error bars indicate standard deviations of the means. C) TEM analysis showing apoptotic bleb and nuclear fragmentation of MAYV infected Huh7.5 cell. C) The cleavage of PARP at 3,6,12, 24 and 48hpi monitors by Western blot analysis in the cell lysate of mock and MAYV Huh7.5 cells. Hoechst 33342 staining of MAYV infected Huh7.5 cells (36hpi) observed under UV light. Arrows (bottom righthand side panel) indicate nuclei that contain condensed chromatin. D) Immunofluorescence detection of Caspase 9 (intrinsic apoptotic marker) and Caspase 8 (extrinsic apoptotic marker) at 36hpi MAYV infected Huh7.5 cells. E) Focus size in Vero cells by MAYV infected Huh7.5 cells treated with apoptosis inhibitor (zVAD-FMK) or vehicle control (DMSO).
Fig 1.

- Current distribution of Mayaro virus
- Distribution of Anopheles mosquitoes (competent vector for Mayaro virus)
Fig 2.

A

Huh 7.5
Human liver
1 moi
0.1 moi

Vero
African green monkey kidney
1 moi
0.1 moi

SHK-21
Baby hamster Kidney
1 moi
0.1 moi

Sua5B
Anopheles gambiae
1 moi
0.1 moi

C6/36
Aedes albopictus
0.1 MOI
1 MOI

Aag2
Aedes aegypti

B

MAYV
ONNV
SIN

Virus titer in log_{10} FFU/mL

Temperature (°C)

C

Control
MAYV

Plaque diameter in mm

MAYV
MAYV
SINV
ONNV

Early stages virus entry inhibitors

Dynasore

NH₄Cl

Bafilomycin

Viral genome
Fig 4.

**A**

BHK-21

C6/36

**B**

Functional uptake assay

Dextran

(FITC)

**C**

Pharmacological inhibition assay

(EIPA)

Macropinocytosis Blocker
BHK-21 cell

Fig 5A.
C6/36 cell

Fig 5B.
**Fig 6.**

(A) Time course and percent cell survival. Cell survival was measured at 0.1 and 1.0 moi.

(B) MAYV infected cells show nuclear fragmentation.

(C) Western blot analysis of Caspases 9, 8, and 3 in mock and MAYV-infected cells.

(D) Immunofluorescent staining of Caspases 9, 8, and 3 in mock and MAYV-infected cells.

(E) Infection of Huh cell line with MAYV, followed by co-culture with Vero cells and treatment with zVAD-FMK. The focus diameter was measured as a marker of infection. The graph shows a significant difference between control and zVAD-FMK treatment (p<0.0001).