Suppression of Drug Resistance in Dengue Virus

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ABSTRACT Dengue virus is a major human pathogen responsible for 400 million infections yearly. As with other RNA viruses, daunting challenges to antiviral design exist due to the high error rates of RNA-dependent RNA synthesis. Indeed, treatment of dengue virus infection with a nucleoside analog resulted in the expected genetic selection of resistant viruses in tissue culture and in mice. However, when the function of the oligomeric core protein was inhibited, no detectable selection of drug resistance in tissue culture or in mice was detected, despite the presence of drug-resistant variants in the population. Suppressed selection of drug-resistant virus correlated with coaggregation of the targeted drug-susceptible and drug-resistant core proteins. The concept of “dominant drug targets,” in which inhibition of oligomeric viral assemblages leads to the formation of drug-susceptible chimeras, can therefore be used to prevent the outgrowth of drug resistance during dengue virus infection.

IMPORTANCE Drug resistance is a major hurdle in the development of effective antivirals, especially those directed at RNA viruses. We have found that one can use the concept of the genetic dominance of defective subunits to “turn cousins into enemies,” i.e., to thwart the outgrowth of drug-resistant viral genomes as soon as they are generated. This requires deliberate targeting of larger assemblages, which would otherwise rarely be considered by antiviral researchers.

For the positive-strand RNA viruses that are considered priority pathogens (dengue virus, West Nile virus, yellow fever virus, Chikungunya virus, severe acute respiratory syndrome coronavirus, hepatitis A virus and the causative agents of several viral encephalopathies), no antiviral pharmaceuticals exist. Among negative-strand RNA viruses, the devastating Ebola outbreak in West Africa and the threat of new influenza virus pandemics further highlight the importance of developing effective vaccines and therapeutics that target any of these highly adaptable, genetically labile RNA viruses (1). Dengue virus, originally confined to tropical and subtropical areas, has begun to spread outside its former geographic limitations due to the changing ecology of mosquito populations in the face of urbanization and global warming. Previously dengue-free countries are now at risk: Japan, with no reported cases of dengue fever in 70 years, suffered a disease outbreak in 2014 (2). Due to the expanded range of each of the four dengue virus serotypes, it is now more likely for individuals who have been infected previously with one serotype to become infected with another. Antibodies from the first infection that do not neutralize the newly infecting serotype can enhance its symptoms, increasing the probability of dengue hemorrhagic fever, a severe and often fatal form of the disease (3). Given the demographics of infection by dengue virus and other emerging pathogens, it is highly desirable to reduce the complexity and expense of vaccines and treatments.

The only positive-strand RNA virus for which effective pharmacological treatment exists is hepatitis C virus (HCV), which serves as an illustration of the current paradigm for successful antiviral development. Anti-HCV drugs that target the NS3/4 protease, the NS5A nonstructural protein, and the NS5B polymerase are currently available. The high cost of these drugs, with individual treatments averaging $80,000 per patient, has contributed to a 13% increase in prescription drug spending in the United States alone since 2013 (4–6). This paradigm is not sustainable for the many infections that threaten human health and productivity worldwide.

The emergence of drug-resistant RNA viruses requires, first, that such viruses are generated and, subsequently, that the selective pressure of drug treatment promotes their selective amplification. All RNA viruses display error-prone replication strategies (7). With error rates of $10^{-4}$ to $10^{-5}$ per nucleotide copied, which result in accumulative mutation frequencies of greater than $10^{-4}$ per round of intracellular quasispecies generation (8), a large amount of preexisting diversity is ensured. In an inoculum of $10^5$ dengue viruses, genomes with mutations at each nucleotide position should be present. Understandably, most approaches to reducing the outgrowth of drug-resistant viral genomes aim to reduce the frequency of generation of those genomes. Viral escape from combination therapy, for example, should require multiple mutations to confer resistance to the drug combination. Similarly, if proteins or functions in the human host are targeted, there may be very few ways for the virus to escape its dependence on those human “host factors,” and therefore such viral genomes will be very infrequent (9).

A different approach to suppress drug resistance is to assume that drug-resistant genomes will inevitably be generated but that...
drug targets can be identified for which the selection for drug resistance will be blunted. Usually, drug resistance is dominant, i.e., a newly generated drug-resistant viral genome will be amplified and selected in the presence of a drug, even though it is generated in the same cell as its drug-susceptible parents and siblings. However, in some cases, defective genomes, such as drug-susceptible genomes, are dominant. This often occurs when the defect is in a protein that forms high-order oligomers. Then, in a cell that contains both defective and functional genomes, the final assemblage will contain both defective and functional subunits. Often, such chimeric structures are nonfunctional (10). Applying this principle to viral infections, we have shown that, although mutant poliovirus genomes that are resistant to capsid inhibitor V-073 could be readily generated, they were not selected in cultured cells or in mice (11). We termed viral targets such as the poliovirus capsid, for which drug-susceptible genomes are dominant over their drug-resistant siblings, “dominant drug targets.”

To identify dominant drug targets in the dengue virus genome, we tested the effects of two different antiviral compounds: MK-0608, a nucleoside analog that inhibits the viral NS5 polymerase, and ST-148, a planar compound that inhibits the function of core protein, an oligomeric structural protein. Both in tissue culture and in mice, the selection of MK-0608-resistant viruses was readily observed. However, viruses resistant to ST-148 were not selected in tissue culture in the presence of drug-susceptible virus, and resistant viruses did not emerge during treatment of infected mice with ST-148. Thus, dengue virus core protein is a dominant drug target, and pharmaceuticals targeted against it will be less likely to be plagued by the emergence of resistant virus.

RESULTS

Susceptibility to core inhibitor ST-148 is genetically dominant.

To identify potential dominant drug targets for dengue virus, we sought published antiviral compounds that target oligomeric viral structures. Byrd et al. in 2013 (12) reported the inhibition of all four dengue virus serotypes by a compound termed ST-148, which was identified in a high-throughput screen (Fig. 1A). Dengue virus core protein was determined to be the compound’s target in direct binding experiments and by the mapping of a drug-resistant mutation, S34L, to the coding sequences for core protein 11.

FIG 1 Coinfections with drug-susceptible and ST-148-resistant viruses. (A) Structure of the core inhibitor ST-148 (12) and space-filling model of the core dimer with ST-148-resistant mutation C-S34L. (B) Resistant mutation C-S34L was introduced into the dengue virus serotype 2 16681 infectious clone and tested for resistance to 10 μM ST-148. After 24 h, supernatants were collected and titers were determined. (C) Illustration of coinfection experiments, the results of which are shown in panels D and E. The black lines indicate drug-susceptible, wild-type RNA genomes; red lines denote drug-resistant RNA genomes, present at constant MOIs. (D) Drug-susceptible wild-type virus was mixed with drug-resistant C-S34L virus at various ratios and used to infect BHK-21 cells in the presence of 10 μM ST-148 for 24 h. Yields of total (blue) and drug-resistant (red) virus are shown. (E) Total viral RNA was extracted from supernatants and quantified by qRT-PCR. (F) Proportions of drug-resistant and drug-susceptible RNA genomes were determined by restriction digestion of amplicons of 476 bp that were generated by bulk RT-PCR from RNA extracted from supernatants of singly or doubly infected BHK-21 cells. Wild-type virus had been engineered to contain a silent AlIII restriction site in the core coding region. (G) Amplicons were cleaved with AlIII and run in 1% agarose–TBE gels. The proportions of resistant and susceptible genomes were determined from the percentage of uncleaved (476-bp) and cleaved (344-bp) PCR products, respectively. Results shown are a representative example of coinfection with wild-type and NS5-A60T virus. All data shown are averages ± standard deviations of results from two to three biological replicates. Statistical analysis of the yield of drug-resistant virus in the coinfections, compared to single infection, was via an unpaired Student’s t test.
distribution would predict that 1/n at an average of 1 FFU (focus-forming unit) per cell. A Poisson distribution would predict that 1/e cells would be uninfected and the rest infected with one or two C-S34L drug-resistant viruses. The number of wild-type viruses per cell ranged from 0 to an average of 3 FFU/cell, an MOI at which almost all cells will be infected by wild-type virus. After one cycle of infection, 24 h, supernatants were harvested and the yields of total and ST-148-resistant virus were determined. The addition of increasing amounts of drug-susceptible wild-type virus resulted in a dose-dependent decrease of total and resistant virus (Fig. 1D). Even when most cells were infected by only one drug-resistant and one drug-susceptible virus, the yield of drug-resistant virus was significantly reduced (Fig. 1D). Therefore, for the core protein inhibitor ST-148, drug susceptibility is genetically dominant.

To test whether a reduction in the accumulation of encapsidated RNA mimicked the reduction in infectivity, viral RNA was extracted from the virus preparations shown in Fig. 1D and quantified by quantitative reverse transcription-PCR (qRT-PCR) (Fig. 1E). The cDNA of the wild-type virus had been engineered to contain a new restriction site so that the proportions of drug-susceptible and drug-resistant genomes could be determined molecularly (Fig. 1F and G). AflII restriction digestion of amplified cDNA products resulted in no change in total or drug-resistant RNA in single infections versus coinfections (Fig. 1E). Therefore, the dominant inhibition of the production of infectious virus was not caused by a failure to synthesize viral RNA or to package it into secreted particles. Instead, particles are generated that are noninfectious. This is consistent with the mechanism proposed for ST-148 inhibition, i.e., targeting core protein assembly and virion morphogenesis.

Biochemical dominance of ST-148-susceptible core proteins. The most likely mechanism for the ability of drug-susceptible core protein to inhibit the function of drug-resistant core protein is the formation of chimeric oligomers. To test whether the drug-susceptible and drug-resistant proteins could coassemble, their cellular localizations (Fig. 2A) in single infections and coinfections were monitored. Scatturo et al. (13) found previously that wild-type core protein was predominately associated with cytoplasmic membranes in the absence of ST-148 (Fig. 2B, top row). In the presence of ST-148, the wild-type core protein was also found in nuclear fractions, which might contribute to the compound’s antiviral activity (13). In contrast, the

![FIG 2](https://example.com/fig2.png)

**FIG 2** Fractionation of dengue virus core protein upon single or double infection with ST-148-susceptible and -resistant viruses in the presence or absence of drug. BHK-21 cells were singly or doubly infected with wild-type or C-S34L virus at multiplicities of infection of 3 FFU/cell in the presence or absence of 10 μM ST-148. Infections were allowed to proceed for 24 h, and cells were fractionated into cytoplasmic (C), membrane (M), and nuclear (N) fractions. Lysed cell samples prior to fractionation were used as “input” (I) controls. (A) Proteins from input, C, M, and N fractions, displayed on 12.5% SDS-acrylamide gels, from cells infected with wild-type virus. Immunoblotting was used to verify fractionation, using antibodies against GAPDH (C fraction), mitofilin (M), and lamin (N). (B) Equal amounts of total protein from fractionated cells after single or double infections were loaded on 12.5% SDS-acrylamide gels and blotted for the presence of core protein. (C) Quantification of the results of the experiment shown in panel B, as the percentage of membrane-associated core protein in the presence of ST-148, drug susceptibility is genetically dominant.

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C-S34L mutant protein, which localized almost exclusively with cytoplasmic membranes in the absence of drug, did not relocalize in the presence of drug (Fig. 2B, middle row). To determine whether the drug-susceptible protein could cause the relocalization of drug-resistant protein, or vice versa, we monitored total core protein in 1:1 infections (Fig. 2B, bottom row). The localization of core in the mixed infection in the presence of drug was not the sum of the individual infections (Fig. 2B, lanes 4 to 6). Instead, it was indistinguishable from that of the drug-susceptible virus alone (Fig. 2B, top row right). The different localization patterns of wild-type and C-S34L mutant core proteins (Fig. 2B, top and middle rows) are quantified in Fig. 2C. Figure 2D shows the expected amounts of core membrane association if the patterns are additive (Fig. 2D, left). The observed pattern (Fig. 2D, right) instead mimicked that of drug-susceptible virus alone (Fig. 2D, right). Therefore, the wild-type, drug-susceptible phenotype is dominant over the drug-resistant phenotype in the presence of ST-148. That the drug-resistant protein fractionates differently when in the presence of the drug-susceptible protein supports the hypothesis that the two kinds of proteins coassemble.

Mutations in the NS5 methyltransferase domain confer resistance to the nucleoside inhibitor MK-0608. Nucleoside analog 2′-C-methyl-7-deaza-adenosine (also termed MK-0608, or 7-DMA) (Fig. 3A) was originally developed as an inhibitor of hepatitis C virus polymerase (14) but also displays antiviral activity against dengue virus serotype 2 in cultured cells and in mice (15). Inhibition of dengue virus serotype 2 in BHK-21 cells is shown in Fig. 3. To select viruses resistant to MK-0608, dengue virus pools grown in C6/36 mosquito cells were serially passaged in BHK-21 cells in the presence or absence of increasing MK-0608 concentrations. For each passage, the multiplicity of infection was below 0.1 FFU per cell, which allowed any drug-resistant variants to propagate in the absence of coinfecting drug-susceptible virus. The amount of viral amplification at each passage is displayed as the ratio of total output virus to the initial inoculum (Fig. 3B). Sequence analysis of passaged viruses revealed mutations in all populations, some of which were identical in the drug-treated and untreated populations (Table 1). Such mutations likely conferred a growth advantage in BHK-21 cells; one of these, Q399H, has proven useful in adaptation to other mammalian cell lines (R. M. Deans et al., unpublished data). Notably, only the two viral pools passaged in the presence of MK-0608 acquired mutations in the NS5 coding region: A60T in pool 1 and Y201H in pool 2 (Table 1). When tested individually, the NS5-A60T and NS5-Y201H mutations...
TABLE 1 Coding mutations observed in viral populations passed in the absence or presence of MK-0608

| Coding region | Mutation(s) found in poola | Drug P1 | Drug P2 | Control P1 | Control P2 |
|---------------|-----------------------------|---------|---------|------------|------------|
| prM           | E                           | Glu194Lys | Ala195Val | Val232Ala | Ala232Thr/Asp/Ala |
|               | E                           | Asp196Thr | Ala196Val | Se596Pro  | Se596Pro   |
| NS1           |                             | Gln396His | Gly396Glu | Gly396Glu | Gly396Glu  |
| NS4A          |                             | Ile224Phe | Ala224Val | Ala224Val | Ala224Val  |
| NS4B          |                             | Ile238Phe | Ala238Val | Ala238Val | Ala238Val  |
| NSS           |                             | Ile224Phe | Leu224Phe | Tyr303His | Tyr303His  |

a Mutations found after sequence analysis of total RNA extracted from four viral pools: two pools were passaged 22 times in the presence of increasing concentrations of MK-0608 (drug P1 and drug P2) and two pools were passaged 17 times in the presence of control DMSO-containing medium (control P1 and control P2). Mutations present at greater than 10% of the population could be observed and are indicated.

The selection of drug-resistant viruses in the first intracellular infectious cycle. To quantify the frequency of drug-resistant viruses in the multiple rounds of amplification that occur during murine infection, we needed a mouse model that gave rise to high viral yields. A frequently used mouse model for dengue virus infection is intravenous inoculation into strain 129 IFNAR⁻/⁻ INFGR⁻/⁻ (alpha and gamma interferon [IFN]-deficient) mice (15, 17, 18). Virus could be readily detected in spleens of infected mice 4 days postinfection (Fig. 5E). C57BL/6 IFNAR⁻/⁻ INFGR⁻/⁻ mice were also susceptible to dengue virus infection, and the yield of virus obtained from their spleens following the same inoculum was on average 10-fold higher (Fig. 5E). As this higher yield facilitated the detection of subpopulations such as drug-resistant viruses, we employed C57BL/6 IFNAR⁻/⁻ INFGR⁻/⁻ mice to measure drug-resistant viruses during treatment of dengue virus-infected mice with ST-148 or MK-0608.

To assess whether ST-148-resistant viruses could be selected during murine infection, C57BL/6 IFNAR⁻/⁻ INFGR⁻/⁻ mice were inoculated with dengue virus and treated orally with ST-148, or vehicle solution as a control, twice daily for 4 days as described previously (12). Treatment with ST-148 led to a significant decrease in dengue virus yield (Fig. 5B). To detect the emergence of drug resistance, spleen homogenates from control and ST-148-treated mice were diluted and used to infect fresh monolayers of BHK-21 cells in the presence of ST-148 (Fig. 5A). After 2 days, supernatant titers were determined and RNA was quantified. Strikingly, no amplification of ST-148-resistant virus (Fig. 5C) or of RNA populations (Fig. 5D) was observed.

Selection for MK-0608 resistance in murine infection. To document any selection for resistance to MK-0608 during murine infection, C57BL/6 IFNAR⁻/⁻ INFGR⁻/⁻ mice were infected and treated orally twice daily with MK-0608 at 40 mg/kg of body weight. After 4 days, MK-0608 treatments significantly decreased total viral yield (Fig. 5B). To determine whether this inhibitory pressure led to the selection of drug-resistant viruses, drug-resistant virus and RNA were quantified as for the ST-148 infection. Virus from control mice showed no increase in drug resistance when passaged in the presence of the drug. However, virus from MK-0608-treated mice showed significantly more amplification of drug-resistant viruses (Fig. 5C) and RNA (Fig. 5D). The amount of drug resistance observed correlated somewhat with the amount of selection pressure exerted (Fig. 5B to D, open symbols). However, this alone does not explain the large increase in resistance observed in MK-0608-treated mice. This highly significant selection for MK-0608 resistance contrasts with the lack of selection observed when the oligomeric core protein was targeted.

DISCUSSION
Dengue virus grows primarily in monocytes, macrophages, and dendritic cells of infected humans and mice (19–21). With intravenous inoculations such as those performed here, virus travels directly to the spleen, where it primarily infects macrophages of the marginal zone (19, 20). Upon infection by the natural intradermal route, on the other hand, dendritic cells present in the periphery become infected before migrating to the spleen (Fig. 6). In either case, in lymph nodes or the spleen, the virus replicates and spreads locally to nearby cells, including red pulp macrophages in the spleen, before dissemination to the bloodstream.

During these steps, there are several opportunities for drug-resistant viruses to arise and to undergo selective pressure. During the

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initial infection, the inoculum of 10^5 FFU/mouse used in this study should contain several viruses with any particular point mutation. Then, a robust amplification step occurs after the initial inoculum has seeded thousands of individual macrophages and dendritic cells (22), and viral genomes begin to amplify and diversify (23). Any new drug-resistant variants that arise during this period must do so in the presence of the drug-susceptible products made from parental and sibling genomes. It is at this step (boxed in Fig. 6) at which the suppression of...
The growth of drug-resistant viruses can be accomplished by the choice of a dominant drug target.

The building blocks of the dengue virus nucleocapsid assembly are thought to be the preassembled dimers of core protein observed in solution (24, 25). Interestingly, the C-S34L mutation, which confers resistance to the core-targeting antiviral compound ST-148, is not located at the dimer interface (Fig. 1), suggesting that both mutant and wild-type proteins are capable of forming higher-order oligomers in the presence or absence of the drug (12). Wild-type and S34L core proteins in mixed infections exhibited similar fractionation patterns, even when the patterns changed in the presence of ST-148 (Fig. 2). Thus, it is likely that the oligomers formed in either the presence or absence of the drug contain both drug-resistant and drug-susceptible core proteins.

For ST-148, drug-susceptible viruses were found to be dominant inhibitors of drug-resistant viruses in deliberate coinfections performed for single infectious cycles in cultured cells (Fig. 1D). Even at ratios of 1:1, significant inhibition of drug-resistant viral growth by coinfection with drug-susceptible virus was observed. Assuming random assembly, this argues that as much as 50% of an oligomeric assembly of core protein can be composed of drug-resistant subunits and remain phenotypically drug susceptible. Such a ratio certainly underestimates the effect that would be seen during an actual infection, during which it would be much more likely for a drug-resistant variant to be created in a cell that already contained dozens, or hundreds, of drug-susceptible genomes.

The finding of phenotypic dominance of drug-susceptible genomes is in marked contrast to the situation observed with MK-0608, a nucleoside inhibitor of dengue virus RNA replication. In this case, the presence of drug-susceptible viruses was found to have no effect on the growth of drug-resistant viruses under many coinfection conditions in culture (Fig. 4).

It is intriguing that the two mutations conferring MK-0608 resistance identified in this work mapped to the methyltransferase domain of NS5 (Fig. 3D). The methyltransferase and polymerase domains are known to interact functionally and structurally (16, 26). Thus, it is possible either that the A60T and Y201H mutations exert allosteric effects on the polymerase active site or that MK-0608 inhibits methyltransferase activity directly. Whatever the particular step of RNA synthesis inhibited by MK-0608, it does not appear to be a dominant drug target. It is possible that other aspects of NS5 function, such as RNA binding, elongation, or translocation, will be targets (27).

After the first cycle of infection, any disseminated drug-resistant viruses could in principle be selected during viral spread within an animal (Fig. 5). One mechanism by which this might be thwarted is if viral spread is predominately local, and any drug-resistant virus that escapes the first infected cell infects neighboring cells in the company of its drug-susceptible relatives.

During 4 days of viral growth in dengue virus-infected mice, selection for ST-148-resistant viruses was not observed, even though, under the same circumstances, resistance to the nucleoside inhibitor MK-0608 increased significantly (Fig. 5C). Although it is possible that mutations leading to ST-148 resistance are not frequently generated or that ST-148-resistant viruses show reduced fitness in the mouse, these hypotheses seem unlikely, because generation of the C-S34L mutant virus requires only a single nucleotide change from the wild-type virus and it was readily selected at a low MOI in cultured cells (12). Instead, we argue that the mechanism of suppression of ST-148-resistant virus in the
mouse is the same as that observed in tissue culture. The continued suppression of drug resistance over 4 days of infection could indicate that viral spread is local, such as might occur in the lymph nodes or spleen.

The present report describes the application of the concept of “dominant drug targets,” initially described for poliovirus (11, 27), to dengue virus, a Category A pathogen for which no current treatment exists. The intracellular replication cycle of dengue virus differs from poliovirus in several ways relevant to the efficacy of dominant drug targeting. Dengue virus is an enveloped virus that continuously buds from infected cells; therefore, the intracellular copy number of viral genomes and their products is likely to be lower than that of poliovirus. The RNA replication complexes of dengue virus are sequestered within endoplasmic reticulum invaginations, which might preclude access of the products of sibling genomes to each other. Nonetheless, the dominance of ST-148 susceptibility argues that core proteins encoded by different genomes interact directly. The creation of dominant inhibitors of drug-resistant viruses by their own families, while macabre, is a novel application of the “quasispecies” concept (7, 28). If implemented, the use of dominant drug targets should reduce the number of drugs necessary for efficacious inhibition of viral infection.

MATERIALS AND METHODS

Cells and viruses. Baby hamster kidney cells (BHK-21, clone 15) were cultured as monolayers in Dulbecco’s modified Eagle’s medium supplemented with 10% (vol/vol) calf serum, 100 units of penicillin/ml, 100 μg streptomycin/ml, and 10 mM HEPES (pH 7.2) at 37°C and 5% CO₂.

Aedes albopictus C6/36 cells were cultured as monolayers in Leibovitz’s L-15 medium supplemented with 10 mM HEPES (pH 7.2), 100 units of penicillin/ml, 100 μg streptomycin/ml, and 10% (vol/vol) fetal bovine serum at 30°C.

Dengue virus 2 serotype 16681 was propagated from infectious cDNA clone pD2IC/30P-A, a gift from Eva Harris, University of California, Berkeley, and originally developed by Kinney et al. 1997 (29). Dengue virus serotype PL046 (17) was also generated from an infectious cDNA (a gift from Sujan Shresta, La Jolla Institute for Allergy and Immunology). All viruses were grown in C6/36 cells, and titers were determined in BHK-21 cells. For mouse experiments, virus was concentrated by ultracentrifugation at 53,000 g for 2 h at 4°C and resuspended in cold endotoxin-free phosphate-buffered saline supplemented with 10% fetal bovine serum.

Compounds and antibodies. MK-0608 (2’-C-methyl-7-deazaadenosine) was purchased from Carbosynth (San Diego, CA). ST-148 was purchased from ChemBridge Corporation (San Diego, CA). Anti-dengue virion antibody that recognizes all four serotypes of dengue virus was purchased from Abcam (Cambridge, MA). Anti-core antibody was a kind gift from Andrea Gamarnik (Instituto Leloir, Argentina). Anti-glyceraldehyde-3-phosphate dehydrogenase (GAPDH) and anti-lamin A/C antibodies were purchased from Santa Cruz Biotechnology (Santa Cruz, CA). Anti-mitofilin antibody was purchased from Proteintech (Chicago, IL).

Plasmids and RNA transcription. Viral cDNA manipulation and generation of infectious RNA for tissue culture experiments have been described elsewhere (30). A silent mutation in the core protein region of the genome that introduced an AflII restriction site in the wild-type genome and all drug resistance mutations were introduced using the QuikChange site-directed mutagenesis kit (Agilent Technologies, Santa Clara, CA). For mouse experiments, serotype 2 strain PL046 was used. Envelope muta-
tions N124D and K128E, which confer increased pathogenicity in dengue virus-susceptible 129 IFNAR−/− IFNGR−/− mice (31), were introduced as described above. Each amplified DNA segment was sequenced in its entirety to ensure that no adventitious mutations were introduced and was then reintroduced into the infectious cDNA backbone to generate infectious RNA, as described by Mateo et al. in 2013 (30).

**Cell fractionation, protein precipitation, and quantitation.** Fractionation of cultured cells was performed using the Qproteome cell fractionation kit (Qiagen, Valencia, CA) following the manufacturer’s instructions. Proteins from each fraction were methanol precipitated, resuspended in 8 M urea, 100 mM Tris-HCl buffer (pH 8.8), and quantified using the DC protein assay kit (Bio-Rad, Hercules, CA).

**Immunoblotting.** Cell lysates were separated by gel electrophoresis on 12.5% or 15% SDS-polyacrylamide gels and transferred to Immobilon polyvinylidene difluoride (PVDF) membranes (Millipore, Billerica, MA) for 90 min at 100 V in a MiniProtein III transfer tank (Bio-Rad, Hercules, CA). Immunoblots were incubated with anti-core antibody at a dilution of 1/1,000, followed by incubation with alkaline phosphatase-conjugated goat anti-rabbit immunoglobulin (Jackson ImmunoResearch, West Grove, PA) at a dilution of 1/20,000. Immunoblots were imaged on a PhosphorImager (Bio-Rad), and band quantitation was conducted with ImageQuant software (Bio-Rad).

**RNA transfection, RNA extraction, and qRT-PCR.** Methods used for RNA transfections in BHK-21 cells and qRT-PCR assays have been described elsewhere (30).

**Determination of proportion of Affili-marked and unmarked genomes.** Supernatants from infected cells were used to extract viral RNA as described by Mateo et al. (30). Total RNA was used as a template for bulk RT-PCR with a SuperScript III one-step RT-PCR system with platinum Taq (Invitrogen, Grand Island, NY) and the following primers: forward, 5′ CGTTGACCCAGAAACAGATTCTT 3′; reverse, 5′ CTTGTCGTC TGACGATCAGTGGT 3′. The resulting amplicon was purified using the NucleoSpin extract II and PCR cleanup kit (Macherey-Nagel, Deer Park, NY) and digested with Affili. The resulting product were run in 1% agarose–Tris-borate–EDTA (TBE) gels and quantified using Alphalmage EP software (Alphalnnotech, San Jose, CA).

**Infectivity assays.** Virus titrations were conducted on BHK-21 cells as described by Mateo et al. (30). To determine titers for only resistant viruses in the populations analyzed, the polymerase inhibitor MK-0608 was added to the overlay at a concentration of 10−5 M ST-148, also resuspended in DMSO, was added at a final concentration of 10−5 M. Infections were indicated (or with 0.5% DMSO as a negative control) for 30 min at 37°C and allowed to proceed for 48 h, after which supernatants were harvested and titers were determined. Viral RNA was extracted and quantified as described above.

**Statistical analysis.** Data were analyzed with Prism software (GraphPad Software, Inc.). Statistical significance was determined by using a two-tailed paired t test for experiments in cultured cells and the Mann-Whitney test for the data illustrated in Fig. 5.

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