Detection of post-vaccination enhanced dengue virus infection in macaques: An improved model for early assessment of dengue vaccines

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Abstract

The need for improved dengue vaccines remains since the only licensed vaccine, Dengvaxia, shows variable efficacy depending on the infecting dengue virus (DENV) type, and increases the risk of hospitalization for severe dengue in children not exposed to DENV before vaccination. Here, we developed a tetravalent dengue purified and inactivated vaccine (DPIV) candidate and characterized, in rhesus macaques, its immunogenicity and efficacy to control DENV infection by analyzing, after challenge, both viral replication and changes in biological markers associated with dengue in humans. Although DPIV elicited cross-type and long-lasting DENV-neutralizing antibody responses, it failed to control DENV infection. Increased levels of viremia/RNAemia (correlating with serum capacity at enhancing DENV infection in vitro), AST, IL-10, IL-18 and IFN-γ, and decreased levels of IL-12 were detected in some vaccinated compared to non-vaccinated monkeys, indicating the vaccination may have triggered antibody-dependent enhancement of DENV infection. The dengue macaque model has been considered imperfect due to the lack of DENV-associated clinical signs. However, here we show that post-vaccination enhanced DENV infection can be detected in this model when integrating several parameters, including characterization of DENV-enhancing antibodies, viremia/RNAemia, and biomarkers relevant to dengue in humans. This improved dengue macaque model may be crucial for early assessment of efficacy and safety of future dengue vaccines.
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Competing interests: The authors declare the following conflicts: MAC, OT, SB, KS-O, YV, CL and LW are employees of the GSK group of companies. DWV used to be an employee of the GSK group of companies at the time when the study was performed. M-PM worked under contract with GSK at the time when the study was performed. KS-O, DWV, YV, CL and LW report owning shares and/or restricted shares in the GSK group of companies. MBB, RSM, RCP, YDSM, LGAM, LD-M, MF, AH and EC are employees of Fiocruz which was contracted by GSK in the context of this study.

risk factor for severe dengue. Thus, if vaccination does not elicit optimal DENV-specific immunity, a vaccine might, instead, increase the risk of severe dengue in vaccinated individuals, as seen with the only licensed vaccine (Dengvaxia) in children naïve to DENV at vaccination. It is thus crucial to assess dengue vaccine safety at the earliest development stages, ideally in the preclinical stage. The dengue macaque model has been used to assess preclinical efficacy of dengue vaccines, with post-challenge DENV replication as the sole efficacy endpoint. However, this model had not predicted the Dengvaxia-associated safety signals. Here we characterized, in macaques, a dengue purified and inactivated vaccine (DPIV) candidate for its immunogenicity and efficacy/safety. Using a multiparameter approach, including characterization of viral replication and biomarkers relevant to dengue/severe dengue in humans, we were able to detect vaccine-associated safety signals in this model. While these results enabled us to discontinue at an early stage the DPIV development, this improved dengue macaque model may also be instrumental for early assessment of efficacy/safety of future dengue vaccines.

Introduction

Dengue viruses 1–4 (DENV-1–4) are mosquito-borne flaviviruses annually responsible for 50–100 million dengue cases in humans, that have been classified as dengue fever (DF), dengue hemorrhagic fever (DHF) and dengue shock syndrome (DSS), and further re-classified as dengue (with or without warning signs) and severe dengue [1,2]. Pre-existing sub-optimal immunity to DENV is thought to be the strongest risk factor for DHF/DSS, for which antibody-dependent enhancement (ADE) of DENV infection is proposed to be the early underlying mechanism. ADE of DENV infection is posited to occur when pre-existing antibodies bind but do not efficiently neutralize DENV virions, resulting in DENV immune complexes that interact with FcγRs, thus facilitating not only entry but also viral replication (due to FcγR-associated suppression of intra-cellular innate immunity) in FcγR-bearing cells [3,4]. ADE of DENV infection has been observed in infants born to DENV-immune mothers, with a peak of incidence of DHF/DSS correlating with the waning of maternally-acquired antibodies [5,6]. The reported association between specific pre-existing DENV-antibody titers and the risk of DHF/DSS further confirmed the role of ADE in severe dengue [7].

The pre-existing immunity risk factor for DHF/DSS has complicated dengue vaccine development. It is believed that a dengue vaccine needs to elicit protective immune responses against all 4 DENV types while not inducing DENV-enhancing antibodies that could increase the risk of severe disease in vaccinated individuals [8,9]. However, the only licensed dengue vaccine, Dengvaxia, was shown to increase the risk of hospitalization for severe dengue in children naïve to DENV before vaccination, thus further emphasizing the need to assess dengue vaccine safety at the earliest development stages prior to human vaccination [10].

DENV vaccine development has also been impaired by the lack of an optimal animal model reproducing human dengue disease. Although several non-human primate species sustain DENV replication after experimental infection, they rarely develop clinical signs [11–13]. Despite this, the macaque is widely accepted as the most suitable model for preclinical characterization of dengue vaccines. Prior to clinical development, all vaccine candidates to date were tested for efficacy in this model using post-challenge viremia as the sole surrogate for disease [14–19]. Nevertheless, the only dengue vaccine for which both preclinical and clinical efficacy results were reported (Dengvaxia) showed almost 100% efficacy at preventing post-challenge viremia in macaques whereas its efficacy in humans was substantially lower. Although the
vaccine was further reported not to prevent post-challenge DENV-2 RNAemia in macaques, no signs of enhanced DENV infection were detected, thus not reflecting the vaccine-associated increased risk for severe dengue reported in children naïve to DENV at vaccination [10,17,20–22]. This suggests that, when determined using the sole post-challenge viral replication, dengue vaccine efficacy in macaques may not predict efficacy and safety in humans.

We hypothesized that the limited predictability of the dengue macaque model might be related to the use of DENV challenge strains isolated long ago and subjected to serial cell passages (thus likely to differ from circulating strains) and/or to the viremia levels that are substantially lower in macaques compared to humans (which could result in an over-estimation of vaccine efficacy in macaques). Another limitation may be the fact that the biomarkers modified in dengue patients are not evaluated, as surrogates of dengue clinical signs, when assessing vaccine efficacy in macaques. Therefore, we previously selected recent and minimally passaged Brazilian DENV clinical isolates (including DENV-1 0111/2011 and DENV-2 0126/2010) which induce robust viremia in macaques, and are associated with changes in cytokine/chemokine profiles sharing some similarities with those reported in DF patients [23]. Finally, the limited predictability of the dengue macaque model may also be explained by the short intervals usually allowed between vaccination and DENV challenge, ranging from 1 (in most studies) to 5–6 months post-vaccination [14,15,17–19]. Indeed, to more accurately predict long-term vaccine efficacy, the challenge should be performed at a sufficiently late time-point post-vaccination, when the dengue-associated short-lasting heterotypic immunity has waned to low/undetectable level, and the vaccine-elicited immunity has reached its low plateau level.

Here we evaluated, in rhesus macaques, a tetravalent dengue purified and inactivated vaccine (DPIV) candidate for its immunogenicity and efficacy to control infection following DENV challenge. We compared between vaccinated and non-vaccinated macaques, not only post-challenge DENV replication but also the changes in soluble immune mediators and hematological/biochemical parameters that are typically associated with dengue in humans.

**Results**

**DPIV elicited long-lasting DENV-neutralizing antibody responses against the four DENV types**

Three groups of monkeys (Gr.1-3) received two doses four weeks apart of DPIV (2 or 4 μg/DENV type) adjuvanted with aluminum hydroxide (Alum) or the adjuvant system AS03B (Table 1). The DENV-neutralizing antibody (DENV-nAb) titers, as measured in sera collected throughout the whole study, are shown in Fig 1. All tested vaccine formulations elicited DENV-nAb responses against the four DENV types that, at month 8 post-second immunization, were still detectable and did not significantly differ between groups (Fig 1 and S1–S3 Tables). As the measured DENV-nAb titers did mostly not differ between months 5 (day 168/173) and 8 (day 254) post-second immunization (S1 Table), we assumed the DENV-nAb response detected eight months following vaccination was representative of long-term immunity.

**DPIV failed at preventing DENV infection while signals of enhanced DENV replication were detected in a few vaccinated macaques**

We next aimed at assessing DPIV efficacy to prevent post-challenge DENV replication. Characterization of dengue vaccine efficacy in macaques was previously performed by measuring, after DENV challenge, viremia [14,15,17] or both viremia and RNAemia, with viremia being the primary measure to conclude on vaccine efficacy [16,19]. However, we and others previously reported low or no detectable post-challenge DENV viremia in vaccinated macaques.
whereas RNAemia was detected at levels similar or higher than those detected in non-vaccinated monkeys. Although this discrepancy was explained by possible detection of viral RNA derived from degraded or neutralized DENV particles, it was not elucidated [16,19]. In our previous study, viremia had been measured using frozen-thawed sera [16]. As freeze-thawing of sera may reduce infectious titer of enveloped viruses-containing samples, here we assessed whether DENV viremia quantification was impacted by freeze-thawing of sera, while hypothesizing that RNAemia quantification might not be impacted.

At month 8 post-second immunization, Gr.1, 2 and 4 (non-vaccinated control group) were each divided into two subgroups (n = 5) and challenged with either DENV-1 0111/2011 or DENV-2 0126/2010 (Table 1). These groups are further referred to as Gr.1-2 versus Gr.4. RNAemia was detected in all animals and, although the area under the curves (AUC) tended to be reduced in most vaccinated subgroups, the mean RNAemia peaks were 2.86- and 3.19-fold higher in Gr.2 compared to non-vaccinated Gr.4 after challenge with DENV-1 0111/2011 and DENV-2 0126/2010, respectively (Fig 2A). However, viremia had also been measured, for two selected days, using fresh sera, and the titers determined on fresh versus frozen-thawed sera were compared (Fig 2B). Unexpectedly, freeze-thawing of sera did reduce viremia titration only in sera derived from vaccinated, but not non-vaccinated macaques, suggesting that comparing viremia titers between vaccinated and non-vaccinated animals could be biased when using frozen-thawed sera. We then focused on the RNAemia to further compare Gr.1-2 versus Gr.4. RNAemia was detected in all animals and, although the area under the curves (AUC) tended to be reduced in most vaccinated subgroups, the mean RNAemia peaks were 2.86- and 3.19-fold higher in Gr.2 compared to non-vaccinated Gr.4 after challenge with DENV-1 0111/2011 and DENV-2 0126/2010, respectively. Furthermore, 7 out of 20 vaccinated macaques showed higher RNAemia peaks (1.02- to 22-fold) compared to the highest peaks detected in the corresponding non-vaccinated subgroups (Fig 2A and 2C and S4 Table).

To investigate whether the high-level RNAemia detected after challenge of Gr.1-2 was restricted to the newly selected isolates DENV-1 0111/2011 and DENV-2 0126/2010, Gr.3 and 5 were next divided into two subgroups and challenged, at month 8.5 post-second immunization,

### Table 1. Vaccine formulations and treatment groups.

| Group | Number of animals | First vaccine dose (Day 0) | Second vaccine dose (Day 28) | DENV challenge |
|-------|------------------|---------------------------|-----------------------------|----------------|
| 1     | 10               | DPIV-2 μg+ Alum           | DPIV-2 μg+ Alum             | M8 pII DENV-1 0111/2011 and DENV-2 0126/2010 |
| 2     | 10               | DPIV-2 μg+ AS03n          | DPIV-2 μg+ AS03n            | M8 pII DENV-1 0111/2011 and DENV-2 0126/2010 |
| 3a    | 9a               | DPIV-4 μg+ Alum           | DPIV-4 μg+ Alum             | M8.5 pII DENV-2 0126/2010 and DENV-2 S16803 |
| 4b    | 10               | NA                        | NA                          | M8 pII DENV-1 0111/2011 and DENV-2 0126/2010 |
| 5b    | 10               | NA                        | NA                          | M8.5 pII DENV-2 0126/2010 and DENV-2 S16803 |

*Group 3 (Gr.3) was included after Gr.1-2;  
Gr.4 and 5 were included, as non-vaccinated control groups, at the time of challenge of Gr.1-2 and Gr.3, respectively;  
One animal from Gr.3 died during the course of the experiment;  
Vaccine was administrated intra-muscularly;  
Gr.1, 2 and 4 and Gr.3 and 5 were subcutaneously challenged with the indicated DENV challenge strains at month 8 post-second immunization (M8 pII) and month 8.5 post-second immunization (M8.5 pII), respectively (n = 5/challenge subgroup but Gr.3 challenged with DENV-2 S16803, n = 4), NA, non applicable.

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with either DENV-2 0126/2010 or the WHO reference DENV-2 strain S16803, which has frequently been used in rhesus macaques to assess efficacy of dengue vaccine candidates [15–17] (Table 1). These subgroups are further referred to as Gr.3 or 5/DENV-2 0126/2010 (n = 5) and Gr.3 or 5/DENV-2 S16803 (n = 4 and 5, respectively). To ensure accurate characterization of post-challenge DENV replication, viremia and RNAemia were measured using both fresh and frozen-thawed sera. Freeze-thawing of sera did not impact RNAemia quantification but, as previously observed (Fig 2B), did reduce viremia titration only in sera derived from vaccinated macaques (Fig 3A and 3B). For further analysis and after having confirmed the positive correlation between viremia and RNAemia values (S1 Fig), we focused on viremia and RNAemia titers in fresh and frozen-thawed sera, respectively. As shown in Fig 3A and 3C and in S5 and S6 Tables, after DENV-2 0126/2010 challenge, reduced viremia/RNAemia were detected in

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**Fig 1. DPIV elicited broad and long-lasting DENV-nAb responses.** Three groups of rhesus macaques received two intra-muscular administrations, 28 days apart, of the indicated formulations. Sera collected before and after immunization were tested, in duplicate, using a plaque reduction neutralization test (PRNT) for their neutralizing activity against each of the four DENV types. The individual reciprocal serum dilutions associated with 50% reduction in plaque counts (PRNT50 titers) were determined. Shown are the geometric mean titers (GMT) and 95% confidence intervals (CI) (n = 10/group except for Gr.3 at day 254, n = 9). Dotted lines indicate the limit of detection. §Sera were collected in Gr.3 at day 168 instead of 173 in Gr.1-2.

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Fig 2. Viremia and RNAemia detected after challenge of Gr.1, 2 and 4 with either DENV-1 0111/2011 or DENV-2 0126/2010. At month 8 post-second immunization, Gr.1, 2 and 4 were divided into two subgroups each (n = 5) which were subcutaneously inoculated with approximately $10^5$ plaque-forming units (PFU) of either DENV-1 0111/2011 or DENV-2 0126/2010. (A) Shown are the individual viremia (expressed as plaque-forming units (PFU)/mL) and RNAemia (expressed as genome equivalent (ge)/mL) determined, using frozen-thawed sera, after inoculation with DENV-1 0111/2011 or DENV-2 0126/2010. Horizontal black and grey dotted lines indicate the threshold of detection for viremia and RNAemia, respectively. Horizontal green and red dashed lines indicate the lowest and highest RNAemia peaks detected in the corresponding non-vaccinated subgroup. Animals with RNAemia peaks higher than the highest peaks detected in non-vaccinated groups are indicated in red font. (B) Fresh sera collected at days 2 and 7 post-challenge were also tested for their viremia content, in parallel. Shown are the individual viremia titers determined using either fresh or frozen-thawed sera from days 2 and 7 post-challenge. Circle, square and triangle symbols correspond to values obtained with Gr.1, 2 and 4, respectively. Open and black symbols correspond to values obtained after challenge with DENV-1 0111/2011 or DENV-2 0126/2010, respectively. The horizontal dashed line indicates the...
vaccinated compared to non-vaccinated macaques, with one animal (AH85) protected from viremia/RNAemia. In contrast, after DENV-2 S16803 challenge, viremia/RNAemia were not reduced, and the mean viremia and RNAemia peaks were 4.06- and 6.99-fold higher in Gr.3 compared to Gr.5, respectively. Importantly, 3 out of 4 vaccinated macaques showed higher RNAemia peaks (1.3- to 83.6-fold) compared to the highest peak detected in the non-vaccinated subgroup.

Altogether, DPIV vaccination failed to prevent post-challenge DENV replication, and was associated with increased RNAemia/viremia peaks in 10 out of 29 vaccinated macaques. While

![Graphs showing viremia and RNAemia detection after challenge of Gr.3 and 5 with either DENV-2 0126/2010 or DENV-2 S16803.](https://doi.org/10.1371/journal.ppat.1007721.g003)
the vaccine failure to prevent DENV replication was further confirmed by the anamnestic responses detected approximately one month after challenge (S2 and S3 Figs), it could not be explained by a possible mismatch between the vaccine and challenge DENV strains as DPIV-elicited immunity neutralized all DENV challenge strains in vitro (S3 and S4 Figs).

**DPIV may have triggered ADE of DENV infection in some vaccinated macaques**

NAb responses are a major component of the DENV-specific immunity, with low and high pre-existing DENV-Ab/nAb titers correlating with severe dengue and, in contrast, with protection from dengue in humans, respectively [7,24,25]. Interestingly, negative correlations were observed between the pre-challenge DENV-nAb titers and the post-challenge RNAemia peaks (Fig 4A). This indicated an impact of the DENV-nAb response on the post-challenge RNAemia levels in DPIV-vaccinated macaques. Furthermore, the positive correlations detected, for the DENV-2 type, between the pre-challenge sera capacities at enhancing, in vitro, DENV infection and RNAemia peaks (Fig 4B) indicated that ADE of DENV infection may have occurred in some vaccinated macaques.

**Higher IFN-γ, IL-10 and IL-18, and lower IL-12 responses were detected after DENV challenge in vaccinated compared to non-vaccinated macaques**

While ADE of DENV infection is proposed to be the early mechanism underlying DHF/DSS, one of the late mechanisms triggering the short-lived plasma leakage and coagulopathy, the hallmarks of DHF/DSS in humans, has been proposed to be the DENV infection-associated cytokine storm, including the pro-inflammatory cytokines interleukin (IL)-2, IL-6, IL-8, IL-12, IL-18, interferon (IFN)-γ and tumor necrosis factor (TNF)-α, the anti-inflammatory cytokine IL-10, the chemokines macrophage inflammatory protein (MIP)-1α, MIP-1β, and monocyte chemoattractant protein (MCP)-1, and the vascular endothelial growth factor (VEGF)-A [1,26–29]. In addition, we recently showed that infection of rhesus macaques with DENV-1 0111/2011 or DENV-2 0126/2010 was associated with modifications of the serum cytokine profile sharing similarities with those associated with dengue in humans, including post-infection increases in MCP-1 and IFN-γ levels [23]. Therefore, DPIV efficacy was further assessed here by characterizing the immune mediator profiles after DENV challenge. Sera collected before and after DENV challenge were tested for their concentrations in IFN-γ, IL-1β, IL-2, IL-6, IL-8, IL-10, IL-12/23, IL-17A, IL-18, MIP-1α, MIP-1β, TNF-α, TGF-α, VEGF, MCP-1, GM-CSF and G-CSF. The results are shown in Figs 5, S5 and S6. In all groups, levels of IL-1β, IL-2, IL-17A, GM-CSF, MIP-1β, TGF-α and VEGF-A were not or only slightly modified after challenge (S5 Fig). Increased G-CSF levels were detected only in Gr.3 and 5, without statistically significant between-group difference (S5 Fig). Increased MIP-1α levels were detected in all groups, with higher increases detected in non-vaccinated than in vaccinated macaques except for Gr.2/DENV-1 0111/2011 (Fig 5A). Consistent with dengue-associated cytokine profiles reported in humans, increased IL-6, IL-8, TNF-α and MCP-1 levels were detected in most groups after challenge. However, statistically significant were only the lower increase in IL-6 levels detected at day 8 in Gr.3/DENV-2 0126/2010, and the higher increases in MCP-1 levels detected at day 5 in Gr.1 and 2/DENV-2 0126/2010, compared to the control groups (Fig 5A).

The most notable changes detected after challenge were related to IFN-γ, IL-10, IL-12 and IL-18. IL-10 and IL-18 levels were increased in all vaccinated groups and INF-γ levels were increased in all vaccinated groups but Gr.2/DENV-1 0111/2011, whereas these cytokines were not or minimally increased in non-vaccinated control groups (Fig 5A). In contrast, increased IL-12 levels were detected in all non-vaccinated groups but not, or only slightly, in vaccinated

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Furthermore, when comparing the maximum cytokine levels, irrespective of the vaccine formulation and the challenge strain, the peak levels of IFN-γ, IL-18 and IL-10 detected after challenge were significantly higher in vaccinated compared to non-vaccinated monkeys, whereas the peak levels of IL-12 were significantly lower in vaccinated compared to non-vaccinated groups (Fig 5B and 5C).

When further assessing the relationship between the maximum levels of IFN-γ, IL-12, IL-18 and IL-10 and the RNAemia peak levels, as shown in S6 Fig, both IL-10 and IL-18 levels were found to positively correlate with RNAemia peak levels (statistically significant for IL-10 and borderline significant for IL-18). Furthermore, the two vaccinated macaques that had shown the highest RNAemia peak increases compared to controls (AF125, 22-fold and AG179, 86.3-fold) also showed the highest increases in IFN-γ, IL-18, IL-10 levels, and the strongest down-regulation in IL-12 levels among all macaques. In contrast, the only vaccinated macaque
Fig 5. Post-challenge immune mediator profiles among vaccinated versus non-vaccinated macaques. Sera collected before and after DENV challenge were tested for their concentration in the indicated immune mediators. (A) Shown are the mean changes from baseline and SEM (n = 5 but Gr.3/DENV-2 S16803, n = 4). The log_{10} transformed changes from baseline were analyzed using an ANCOVA model. The p-values compare vaccinated to their corresponding non-vaccinated control subgroups with color codes referring to the vaccinated groups (*, p<0.05; **, p<0.01; ***, p<0.001). (B) Heat map representation of normalized scores of individual maximum changes from baseline in immune mediator levels. Monkeys were grouped by DENV challenge strain/wave.
that was fully protected from DENV replication (AH85) did not show any increase in IFN-γ and IL-10 levels but showed slightly increased IL-12 and IL-18 levels (Fig 5B). Altogether, these results showed that a lack of protection from DENV replication was associated with increased IFN-γ, IL-10 and IL-18 levels together with the absence/down-regulation of IL-12, all exacerbated in the case of increased RNAemia/viremia. In contrast, the combination of increased IL-12 and IL-18 levels, without increased IL-10 levels, might be associated with protection from DENV infection.

Higher AST levels were detected after DENV challenge in vaccinated compared to non-vaccinated macaques

The main hematological and biochemical parameters monitored in clinical practice to predict dengue disease evolution are platelet and white blood cell (WBC) counts, hematocrit (HCT) and serum liver aminotransferases, both alanine and aspartate aminotransferases (ALT and AST), with thrombocytopenia, leukopenia, hematocrit increase >20% and elevated serum levels of AST/ALT being typical biological features of severe dengue in humans [30]. Therefore, hematological and biochemical parameters were characterized using blood samples collected before and at day 7 post-DENV challenge. The results are shown in S7 Fig. When comparing the changes from baseline, irrespective of the vaccine formulation and the DENV challenge strain, serum levels of AST were significantly higher in vaccinated compared to non-vaccinated macaques (S7B Fig). While no other statistically significant difference was detected, we cannot exclude that some differences might have been present at other time-points post-challenge.

Discussion

To inform further clinical development, the DPIV vaccine candidate was assessed in rhesus macaques for its immunogenicity and efficacy to control DENV infection after a late challenge performed approximately eight months post-second immunization. While all tested vaccine formulations elicited long-lasting and cross-type DENV-Ab responses, they failed to prevent, not only post-challenge DENV replication, but also the biological changes that have been consistently associated with dengue severity in humans. In addition, increased DENV RNAemia/viremia, correlating with serum capacity at enhancing DENV infection in vitro, were observed in some vaccinated compared to non-vaccinated macaques, indicating the vaccine may have triggered ADE of DENV infection.

Serum/plasma DENV loads have been reported as positively correlating with dengue disease severity in humans [31–33], and so were DENV RNA levels [34,35]. Therefore, the non-reduced viremia/RNAemia observed after DPIV vaccination (except in Gr.3/DENV-2 0126/2010), and increased RNAemia peaks in some animals, might translate into lack of protection from disease and possibly enhanced disease in humans.
The post-challenge DENV replication was determined by measuring both viremia and RNAemia using either fresh or frozen-thawed sera. RNAemia quantification was not impacted by freeze-thawing of sera. In contrast and unexpectedly, the measured viremia titers were drastically lower when using frozen-thawed compared to fresh sera only for vaccinated but not non-vaccinated macaques. Although the limited remaining sera samples, together with the low viremia titers, prevented us from investigating the mechanism(s) underlying the lower detection of viremia in DPIV-immune frozen-thawed sera, we hypothesize that the viral particles may be captured within immune complexes that might be more prone to freeze-thawing-driven degradation, when compared to naked viral particles in sera from non-immunized animals. Importantly, if in the present study viremia had only been determined using frozen-thawed sera, and RNAemia had not been assessed, the wrong conclusion that DPIV immunization confers protection would have been made, and the enhanced DENV replication would have remained undetected. If, in previous dengue vaccine preclinical studies, viremia was assessed using frozen-thawed sera, post-challenge DENV replication was likely underestimated in vaccinated animals, and vaccine efficacy may have been overestimated. In addition, possible signals of vaccine-associated enhanced DENV replication would have been less likely detected. This might contribute to explain the discrepancy between the preclinical and clinical efficacy results reported for Dengvaxia [10,17,21,22]. Furthermore, beyond dengue vaccine studies in macaques and to avoid possible major biases in viral load quantification in preclinical/clinical/epidemiological studies, investigations should urgently be performed to determine whether such a phenomenon also occurs in humans (and in other animal models), and may be extended to other viruses/pathogens. In the meantime, our results strongly suggest that if measurement of viremia using fresh sera is not possible, measurement of viral genome should be favored.

IFN-γ, IL-10 and IL-18 have been frequently reported to be higher in the serum of DHF/DSS patients compared to DF patients and may thus be considered as immune markers of severe dengue in humans [27,29,32,36–45]. In contrast, IL-12 has been reported to be lower or not detectable in serum of DHF/DSS patients compared to DF patients [46,47], and its absence/reduced level may therefore be also considered as a marker of severe dengue.

IL-12 is a pro-inflammatory cytokine, mainly produced by phagocytic and antigen-presenting cells following exposure to intra-cellular pathogens including viruses. IL-12 plays a major role in early antiviral immunity through promoting IFN-γ production and thus Th1-type responses. In contrast, the absence/reduced levels of IL-12, driven by IL-10 that acts as the main inhibitor of IL-12, greatly contribute to skewing of the cytokine balance towards a Th2-biased response [48].

IL-18 is a pro-inflammatory cytokine, an important function of which is the induction, in concert with IL-12, of IFN-γ production [49,50]. However, in the absence of IL-12, IL-18 induces Th2-type cytokines and may therefore, depending upon the context, either promote Th1 or Th2 responses [49–51]. Interestingly, increased IL-18 levels together with decreased IL-12 levels were previously reported in human immunodeficiency virus (HIV)-infected patients to result in decreased IFN-γ production and Th2-biased immune responses, both being associated with increased HIV replication and development of acquired immunodeficiency syndrome [52]. Hence, it is likely that the combination of increased IL-18 levels and decreased IL-12 levels may also play an important role in dengue immunopathogenesis.

IFN-γ is mainly produced by activated T lymphocytes and NK cells and plays a crucial role in antiviral immunity, both through induction of effector molecules including nitric oxide and enhancement of antigen presentation and apoptosis [53]. The role of IFN-γ in DENV infection is controversial with several studies showing an association of elevated IFN-γ levels with DHF/DSS [36,39,42,54] whereas others showed that sustained IFN-γ production was associated with
protection from dengue-associated clinical manifestations [55]. Although the peak levels of IFN-γ were significantly increased in vaccinated compared to non-vaccinated groups, only one group, i.e. Gr.3/DENV-2 0126/2010, showed sustained IFN-γ production after challenge (Fig 5A). As this group was also the one with better controlled DENV-replication (Fig 3A and 3C), our results corroborate the report by Gunther et al [55]. Given the non-reduced/enhanced DENV replication in the other vaccinated groups, we speculate that unlike sustained IFN-γ production, short-lasting IFN-γ production, albeit high, might fail to control DENV replication.

IL-10 is a Th2-type anti-inflammatory cytokine, mainly produced by monocytes/macrophages, type 2 T-helper and regulatory T cells, which acts as a major suppressor of antiviral immunity. Serum IL-10 level has been consistently reported as being drastically elevated in DHF/DSS patients, compared to DF patients, and is accepted as the strongest predictive immune marker of DHF/DSS [27,29,32,37,42,56,57]. In summary, an exacerbated IL-10 production may down-regulate IL-12, resulting, in the presence of increased IL-18 levels, in suppression of IFN-γ and thus in impaired antiviral immunity.

The positive correlations detected between the capacities of pre-challenge sera at enhancing DENV infection in vitro and the post-challenge RNAemia peaks indicated that ADE of DENV infection may have occurred in some vaccinated macaques. ADE of DENV production has been postulated to be mediated through promotion of virus entry into FcγR-bearing cells, resulting in an increased infected cell mass [3,4,58]. In addition, DENV infection via the FcγR-mediated pathway was shown, in vitro, to be associated with up-regulation of IL-10 expression and, in contrast, down-regulation of IL-12 expression, resulting in suppression of innate intracellular antiviral responses, and enhanced DENV replication. [3,59–63]. The combination of increased IL-10 and decreased IL-12 levels observed in some vaccinated macaques thus further supports that the vaccination may have triggered ADE of DENV infection.

Varying degrees of liver involvement have been reported in dengue patients and are believed to result from DENV-triggered hepatocyte apoptosis, hypoxic damage due to impaired liver perfusion associated with fluid leakage, oxidative stress and/or immune-mediated injury [64,65]. Elevation of liver aminotransferases is commonly reported during dengue in humans and serum ALT and AST levels are established biomarkers used to monitor dengue-related liver injury, with more frequent and higher AST level raise, when compared to ALT, making AST a more robust biomarker to monitor liver injury in dengue patients [30,66–68]. Furthermore, AST and ALT levels were consistently reported as being significantly higher in DHF/DSS patients compared to DF patients and are thus accepted as markers of disease severity [67,69,70].

Altogether, the higher AST, IL-10, IFN-γ, IL-18 levels and, in contrast, lower IL-12 levels detected after DENV challenge, in vaccinated compared to non-vaccinated macaques (Figs 5 and S7), reflect the overall biochemical/hematological and immune mediator profiles observed in humans experiencing severe dengue while further indicating that FcγR-mediated DENV infection may have occurred in vaccinated macaques.

In summary, although DPIV was immunogenic given as two doses four weeks apart, it failed to confer protection from DENV infection and may have triggered ADE of DENV infection. These results do not support further development of DPIV in humans. Despite the limitations of the dengue macaque model, we demonstrate here that, when assessing vaccine efficacy by performing a late DENV challenge, by accurately measuring post-challenge viral replication (both viremia and RNAemia, and viremia on fresh sera) and using a multiparameter approach, safety signals associated with a dengue vaccine candidate can be detected in this model. While there is room for further improvement of the dengue macaque model (for example by measuring the DENV infection-associated soluble nonstructural protein 1 which has been shown to
directly trigger endothelial hyperpermeability and vascular leakage [71–73]), we believe the improved model described here may already be crucial to assess preclinical efficacy and safety of next-generation dengue vaccine candidates, and therefore de-risk large-scale human vaccination.

**Materials and methods**

**Cell culture and viruses**

Vero cells (ATCC No CCL-81) were grown at 37˚C in a humidified 5%-CO₂ incubator in medium 199 with Earle’s salts supplemented with 5%-FBS (Gibco). U937 cells (ATCC No CRL-1593.2) were grown at 37˚C in a humidified 5%-CO₂ incubator in RPMI 1640 medium supplemented with 10%-FBS (Gibco). The DENV-1 0111/2011 and DENV-2 0126/2010 strains [23] were kindly provided by Dr Ana Bispo de Filippis (Instituto Oswaldo Cruz, Rio de Janeiro, Brazil), and used both for *in vivo* viral challenge and plaque-reduction neutralization test (PRNT). The DENV-1 WestPac-74 and DENV-2 S16803 strains were kindly provided by Dr Kenneth Eckels (Walter Reed Army Institute of Research, Silver Spring, USA), and used for *in vivo* viral challenge, PRNT, and ADE assay (DENV-2 S16803), and for ADE assay (DENV-1 WestPac-74). The DENV-1 60305 [74], DENV-2 44/2 [75], DENV-3 16562 [74] and DENV-4 TVP360 strains were used in PRNT.

**Vaccine formulations**

DPIV antigen is based on a combination of attenuated viruses from all four DENV types amplified in Vero cells, purified by size-exclusion chromatography, and inactivated using both UV-treatment and formaldehyde. The four attenuated DENV strains used to produce DPIV have been previously described [76–79]. The same amount of antigen, either 2 or 4 μg, was used for each DENV type. The tetravalent DPIV antigen was then formulated with either Al (OH)₃ or the GSK proprietary Adjuvant System AS03B containing α-tocopherol and squalene as an oil in water emulsion.

**Ethics statement**

The study protocol (protocol P-58/14-2 with license LW-49/14) was approved by the Institutional Ethical Committee for Use of Animals (CEUA-Fiocruz) and conducted in strict accordance with the recommendations from the Guide for Care and Use of Laboratory Animals of the Brazilian Society of Science in Laboratory Animals and the National Council for the Control of Animal Experimentation. The experiment was performed in a biohazard level 2 animal facility (temperature 20–22˚C; humidity 50–60%; light/dark cycle 12 h/12 h). Monkeys were acclimated for 14 days before study start. Monkeys were housed individually but retained in a social environment through visual contact with other monkeys. Polished stainless-steel mirrors, PVC and wooden teethers, as well as foraging tray containing food such as pieces of cereal bars, raisins, rice grains or sunflower seeds were given as environmental enrichment. Monkeys had free access to water and received a commercial diet (Nuvilab Primates 6030 Nuvital) supplemented with fresh fruits and vegetables. Throughout the study, monkeys were observed twice a day by animal care and veterinary staff for health and well-being assessment. Monkeys were anesthetized with ketamine (8–10 mg/kg) prior to immunization, virus inoculation and blood drawing. At the end of the study, monkeys were anesthetized by intra-muscular injection of ketamine (20 mg/kg) prior to being euthanized by intra-peritoneal injection of thiopental sodium (50 mg/kg).
Animals and experimental procedures

Fifty male or female adult rhesus macaques (Macaca mulatta) of Indian origin, 3-10-year-old and weighing 4.9–9.5 kg at the study start, flavivirus-naïve and colony-born in captivity in the Non-human Primates Breeding Service from the Institute of Science and Technology in Biomodels of the Oswaldo Cruz Foundation (Rio de Janeiro, Brazil), were used in this study. The 50 rhesus macaques were randomly allocated into five groups (n = 10/group) using the Sās Proc Optex software and taking into account the sex, the weight and the age. The male (M)/female (F) sex ratio was as follows: 7 M/3 F in Gr.1, 2 and 4, and 6 M/4 F in Gr.3 and 5. The sample size was determined based on 80% power analysis to detect a geometric mean ratio (GMR) of 3 for between-group geometric mean titer (GMT) comparisons, as well as to detect a GMR of 30 for between-group RNAemia area under the curves (AUC) comparisons while estimating, based on our previous similar experiment [16], at 0.38 and 0.75 the standard deviations for PRNT and qRT-PCR assays, respectively. Monkeys were anesthetized with ketamine (8–10 mg/kg) prior to immunization, virus inoculation and blood drawing. For vaccination, Gr.1, 2 and 3 received two intra-muscular injections of 0.5 mL of adjuvanted DPIV 28 days apart, as described in Table 1. Gr.4 and 5 were included as non-vaccinated control groups at the time of the DENV challenge of Gr.1-2 and 3, respectively. Blood samples were collected before each immunization, at days 56, 112, 168 or 173, 224 and 254 post-first immunization, as well as before DENV challenge, on a daily basis until at least day 12 post-challenge, and at day 39 post-challenge. Prior to DENV challenge, each group was subdivided into two groups into which macaques were randomly allocated using the same methodology than that described above. At month 8 post-second immunization, monkeys from Gr.1, 2 and 4 were subcutaneously challenged with 0.5 mL of sterile culture medium containing approximately 10^5 plaque-forming units (PFU) of either DENV-1 0111/2011 (further referred to as Gr.1, 2 or 4/DENV-1 0111/2011 subgroups) or DENV-2 0126/2010 (further referred to as Gr.1, 2 or 4/DENV-2 0126/2010 subgroups). At month 8.5 post-second immunization, Gr.3 and 5 were subcutaneously challenged with 0.5 mL of sterile culture medium containing either approximately 10^5 PFU of DENV-2 0126/2010 (further referred to as Gr.3 and 5/DENV-2 0126/2010 subgroups) or 4x10^4 PFU of DENV-2 S16803 (further referred to as Gr.3 and 5/DENV-2 S16803 subgroups). The M/F sex ratio and sample size/subgroup were as follows: 3 M/2 F and n = 5 in Gr.1, 2 or 4/DENV-1 0111/2011, in Gr.3 or 5/DENV-2 0126/2010 and in Gr.5/DENV-2 S16803, 4 M/1 F and n = 5 in Gr.1, 2 or 4/DENV-2 0126/2010, and 3 M/1 F and n = 4 in Gr.3/ DENV-2 S16803. After inoculation, titer of the residual viral inoculum was confirmed by plaque assay. One monkey from Gr.3 became ill and died before the end of the study (at month 5 post-second immunization). When performing post-mortem histological analysis, this macaque was diagnosed with hydrocephalic brain. No additional microscopic lesion was observed in both the central nervous system and the extraneural organs. Although the hydrocephalus etiology was not determined, we hypothesized it to be congenital and most likely not related to the DPIV vaccination.

Plaque-reduction neutralization test (PRNT)

The PRNTs were performed using different DENV strains when compared to those DPIV is derived from. The following strains were used: DENV-1 60305, DENV-2 44/2, DENV-3 16562 and DENV-4 TVP360 or the DENV challenge strains (DENV-1 0111/2011, DENV-2 0126/2010 and DENV-2 S16803). One hundred DENV PFU were mixed with equal volumes of serially diluted sera and incubated for 1 h at 37˚C. The mixtures were added for 1 h at 37˚C onto Vero cells previously seeded into 6-well plates, and subsequently replaced by maintenance medium supplemented with 2%-carboxyl-methyl-cellulose (CMC). Seven (DENV-2) or eight
(DENV-1, DENV-3 and DENV-4) days later, cells were fixed overnight with 5%-formaldehyde solution prior to crystal violet staining. PFU were counted by the naked eye and the percent neutralization was determined relative to the number of PFU counted with the virus control (corresponding to 0% neutralization). PRNT50 titers, corresponding to the reciprocal serum dilution associated with 50% reduction in plaque counts, were determined using a linear model. All serum samples were tested in duplicate. In the absence of detection of neutralizing activity, the corresponding sample was assigned an arbitrary titer corresponding to half the limit of detection, namely 5.

Antibody-dependent enhancement (ADE) assay
Heat-inactivated macaque sera, collected on day 254 (before challenge), were diluted 4-fold starting from 1:2 prior to being incubated for 2 h at 37˚C with either DENV-1 WestPac-74 (MOI = 1) or DENV-2 S16803 (MOI = 10) in maintenance medium without FBS. As negative controls for DENV infection enhancement, viruses were mixed with medium only with sera derived from DENV/DPIV-naïve macaques (from Gr.4 and 5). A total of 5x10⁴ U937 cells were then added to each well containing the sera/virus mixtures, and incubated for 1 h at 37˚C. The medium was next removed and replaced by maintenance medium with 10%-FBS, and the cells were incubated for 48 h at 37˚C. The cells were then harvested, washed and fixed/permeabilized (BD Cytofix/Cytoperm, BD Biosciences) prior to intracellular staining for the DENV envelop using the pan-flavivirus 4G2 mouse monoclonal antibody (8 μg/mL) followed by 0.7 μg/mL phycoerythrin-conjugated rat anti-mouse IgG2a (Biolegend). The proportion of infected cells was then determined by flow-cytometry (the data were acquired using either FACs LSRII or Fortessa, and analyzed using the FlowJo 9.9.6 software, all from BD Biosciences). DENV infection enhancement was determined relative to the infection percent detected with virus alone (without serum). The cut-off for DENV infection enhancement was set at 2.5-fold infection increase compared to virus alone.

Virus titration by plaque assay
Serial dilutions of macaque sera were added onto Vero cells previously seeded into 6-well plates. After 1 h at 37˚C, the diluted samples were replaced by maintenance medium supplemented with 2%-CMC. Seven days later cells were fixed overnight with a 5%-formaldehyde solution prior to crystal violet staining. PFU were counted by the naked eye and infectious virus titers were expressed as PFU/mL. All serum samples were tested in duplicate. In the absence of PFU detection, the corresponding sample was assigned an arbitrary titer corresponding to half the limit of detection, namely 2.5 PFU/mL.

DENV genome equivalents quantification by real-time RT-PCR
Viral RNA was extracted from 200 μL of monkey sera using the LSI MagVet Universal Isolation Kit (YSI-Thermo Fisher Scientific). DENV genome equivalents were quantitated by real-time RT-PCR using the AgPath-ID One-Step RT-PCR kit (Ambion). Each RT-PCR reaction mixture contained 2.5 μL of RNA, 1.67 μL of Detection Enhancer, 2X RT-PCR Buffer, 25X RT-PCR Enzyme Mix (all from the kit), 20 U of RNAsin (Ambion), as well as 10 and 5 pmol of DENV type-specific forward/reverse primers and probes, respectively. Primers and probes used were: DENV-1, forward 5’-GCA-TTY-CTA-AGA-TTT-CTA-GCC-ATA-CC-3’, reverse 5’-TCG-CT C-CAT-TCT-TCT-TGA-ATG-AG-3’, probe 5’-AAC-AGC-AGG-AAT-3’; and DENV-2, forward 5’-CTG-CAR-GGA-CGA-GGA-CCA-TT-3’, reverse 5’-GGG-ATT-GTT-AGG-AA A-CGA-AGG-A-3’, probe 5’-AAA-CTG-TTC-ATG-GCC-CTG-GTG-3’. RT was performed using an ABI 7500 Real-Time PCR system (Applied Biosystems) at 45˚C for 10 min, followed
by an incubation step at 95˚C for 10 min and 40 cycles of 15 s at 95˚C and 1 min at 60˚C. All serum samples were tested in duplicate. In the absence of RNAemia detection, the corresponding sample was assigned an arbitrary titer corresponding to half the limit of detection, namely 80 ge/mL.

**Cytokine quantification**

The seventeen following cytokines were assessed: IFN-γ, IL-1β, IL-2, IL-6, IL-8, IL-12/23, IL-17A, IL-18, TNF-α, IL-10, G-CSF, GM-CSF, MCP-1, MIP-1α, MIP-1β, TGF-α, and VEGF-A. Undiluted sera were tested in duplicate using the MILLIPLEX MAP Non-Human Primate Cytokines Magnetic Bead Panel Kit (Merck). Data were acquired and analyzed using the Luminex 200 reader and the BioPlex Manager software (BioRad). Results were expressed as pg/mL and log10-transformed for statistical analysis. In the case of a cytokine level \( \leq 1 \), the level was arbitrary set at 1 prior to log10-transformation for statistical analysis. When no signal was detected, the corresponding sample was assigned the arbitrary value of half the limit of detection for the corresponding cytokine. The changes from baseline in cytokine concentrations were analyzed in an ANCOVA model with group, time and group-by-time interaction as factors and baseline value as covariate, the structure of correlation between time-points being of the Toeplitz type. For each of the four DENV challenge waves (Gr.1, 2 and 4 challenged with DENV-1 0111/2011, Gr. 1, 2 and 4 challenged with DENV-2 0126/2010, Gr.3 and 5 challenged with DENV-2 0126/2010 and Gr.3 and 5 challenged with DENV-2 S16803), a non-parametric test was used to compare the maximum change from baseline in cytokine levels between vaccinated versus non-vaccinated macaques. For heat map representation, normalized scores for maximum change from baseline in cytokine levels were determined by challenge wave.

**Laboratory testing for hematological and biochemical parameters**

Hematological analyses were performed using the automatic counter Sysmex pocH Diff (Roche Diagnóstica Brasil Ltda, São Paulo, Brazil). Biochemical analyses were performed using the auto-analyses system Vitros 250 XRC (Johnson–Johnson Clinical Diagnostics, Rochester, NY). The following parameters were evaluated: red and white blood cell counts (RBC and WBC), alanine aminotransferase (ALT), aspartate aminotransferase (AST) and creatinine (CREA). Baseline levels were determined from blood samples collected before DENV challenge.

**Statistical analysis**

All statistical analysis methods are described in the corresponding method sections and/or figure and table legends.

**Supporting information**

*S1 Fig. Correlations between viremia and RNAemia.* Black square and circle symbols correspond to values obtained from Gr.3 after challenge with DENV-2 S16803 and DENV-2 0126/2010, respectively. Grey square and circle symbols correspond to values obtained from Gr.5 after challenge with DENV-2 S16803 and DENV-2 0126/2010, respectively. Pearson and Spearman correlations were performed to compare AUC and peak levels, respectively (***, \( p < 0.001 \)).

(TIF)
S2 Fig. DENV-neutralizing antibody responses as determined before and after DENV challenge. Sera collected before challenge (on day 254 post-first immunization) and 39 days after challenge (day 309 and 321 for Gr.1, 2 and 4 and Gr.3 and 5, respectively) were tested, in duplicate, in plaque reduction neutralization test (PRNT) for their neutralizing activity against each of the four DENV serotypes. The DENV challenge strains are indicated above graphs. The individual reciprocal serum dilutions associated with 50% reduction in plaque counts (PRNT50 titers) were determined. The geometric mean titers (GMT) and 95% confidence intervals (CI) are shown for each of the three vaccinated groups (n = 5/group except for Gr.3/DENV-2 S16803 for which n = 4).

(TIF)

S3 Fig. Neutralizing antibody responses against DENV challenge strains as determined before and after challenge. Sera collected before challenge (on day 254 post-first immunization) and 39 days after challenge (day 309 and 321 for Gr.1, 2 and 4 and Gr.3 and 5, respectively) were tested, in duplicate, in plaque reduction neutralization test (PRNT) for their neutralizing activity against each of the DENV challenge strains. The DENV strains used to challenge the different subgroups are indicated above graphs. The individual reciprocal serum dilutions associated with 50% reduction in plaque counts (PRNT50 titers) were determined. The geometric mean titers (GMT) and 95% confidence intervals (CI) are shown (n = 5/subgroup except for Gr.3/DENV-2 S16803 for which n = 4).

(TIF)

S4 Fig. DPIV-elicited antibody responses potently neutralized in vitro the three DENV challenge strains. Sera collected before immunization and at days 56 and 254 post-first immunization were tested, in duplicate, in plaque reduction neutralization test (PRNT) for their neutralizing activity against the DENV challenge strains, which are indicated above graphs. The individual reciprocal serum dilutions associated with 50% reduction in plaque counts (PRNT50 titers) were determined. The geometric mean titers (GMT) and 95% confidence intervals (CI) are shown for each of the three vaccinated groups (n = 5/group except for Gr.3/DENV-2 S16803 at day 254 for which n = 4).

(TIF)

S5 Fig. Serum levels in IL-1β, IL-2, IL-17, MIP-1β, G-CSF, GM-CSF, TGF-α and VEGF-A were not or slightly modified after challenge with DENV-1 0111/2011, DENV-2 0126/2010 or DENV-2 S16803. Sera collected before (baseline) and at days 1, 4, 6, 8, 10 and 14 after challenge were tested, in duplicate, for their concentration in the indicated soluble mediators. Results were expressed as pg/mL. When no signal was detected, the corresponding sample was assigned the arbitrary value of half the limit of detection for the corresponding mediator. Shown are the mean changes from baseline and SEM from 5 (Gr.1, 2, 4/DENV-1 0111/2011, Gr.1-5/DENV-2 0126/2010, Gr.5/DENV-2 S16803) and 4 (Gr.3/DENV-2 S16803) animals. For statistical analysis, the log_{10}-transformed changes from baseline were analyzed using an ANCOVA model with group, time and group-by-time interaction as factors and baseline values as covariates. The calculated p-values compare, by DENV challenge strain, vaccinated groups to their corresponding non-vaccinated control groups with color codes referring to the vaccinated groups (*, p<0.05; **, p<0.01; ***, p<0.001).

(TIF)

S6 Fig. Relationship between RNAemia peaks and maximum changes from baseline in IFN-γ, IL-10, IL-12 and IL-18 levels. The relationship between the RNAemia peaks and the maximum changes from baseline in IFN-γ, IL-10, IL-12 and IL-18 levels was assessed using a linear regression model performed on log_{10}-transformed values. Shown are all individual
values together with, for the vaccinated groups, the linear regression lines. The statistical significance of the linear regression slopes to be different from 0 was assessed across the different vaccinated groups/challenge waves. The measured $p$ values are indicated. No $p$ value could be calculated for IFN-$\gamma$ due to inter-group interference.

(TIF)

S7 Fig. Post-challenge changes from baseline in hematological and biochemical parameters among vaccinated versus non-vaccinated macaques. Whole anticoagulated venous blood samples, collected before (baseline) and at day 7 post-DENV challenge, were tested for the indicated hematological and biochemical parameters (ALT, alanine aminotransferase; AST, aspartate aminotransferase; GGT, gamma glutamyl transferase; HCT, hematocrit; WBC, white blood cells; MCH, mean corpuscular hemoglobin; MCHC, mean corpuscular hemoglobin concentration; MCV, mean corpuscular volume). (A) Heat map representation of normalized scores of individual changes from baseline. Monkeys were grouped by DENV challenge strain/wave, further divided based on their vaccination status, and ranked, within each subgroup, based on their maximum RNAemia level, monkeys with the lowest and the highest RNAemia peaks being on the left and the right sides, respectively. Score normalization was performed by DENV challenge strain/wave so that normalized scores can only be compared between vaccinated and non-vaccinated macaques within each DENV challenge strain/wave. The only parameter for which the change from baseline was further shown to significantly differ between vaccinated and non-vaccinated macaques is shown in red font. (B) An ANOVA model was used to compare, across the DENV challenge strains/waves, the changes from baseline in hematological/biochemical parameters between vaccinated and non-vaccinated macaques. Shown are the individual values for AST (*, $p<0.05$).

(TIF)

S1 Table. Between-time-point PRNT50 comparisons.

(DOCX)

S2 Table. Between-DENV type PRNT50 comparisons.

(DOCX)

S3 Table. Between-group PRNT50 comparisons.

(DOCX)

S4 Table. RNAemia area under the curves, peaks and durations after challenge of Gr.1-2 and Gr.4 with either DENV-1 0111/2011 or DENV-2 0126/2010 (frozen-thawed sera).

(DOCX)

S5 Table. Viremia area under the curves, peaks and durations after challenge of Gr.3 and Gr.5 with either DENV-2 0126/2010 or DENV-2 S16803 (fresh sera).

(DOCX)

S6 Table. RNAemia area under the curves, peaks and durations after challenge of Gr.3 and Gr.5 with either DENV-2 0126/2010 or DENV-2 S16803 (frozen-thawed sera).

(DOCX)

S1 Data. Hematology.

(DOCX)

S2 Data. Immune mediators.

(XLSX)
S3 Data. Individual PRNT50.
(XLSX)

S4 Data. Individual PRNT50 against challenge DENV strains.
(XLSX)

S5 Data. Viremia_Gr1_Gr2_Gr4.
(XLSX)

S6 Data. Viremia_Gr3_Gr5.
(XLSX)

S7 Data. RNAemia_Gr1_Gr2_Gr4.
(XLSX)

S8 Data. RNAemia_Gr3_Gr5.
(XLSX)

S9 Data. ADE.
(XLSX)

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References
1. Guzman MG, Harris E. Dengue. Lancet. 2015; 385: 453–465. https://doi.org/10.1016/S0140-6736(14)60572-9 PMID: 25230594
2. Stanaway JD, Shepard DS, Undurraga EA, Halasa YA, Coffeng LE, Brady OJ, et al. The global burden of dengue: an analysis from the Global Burden of Disease Study 2013. Lancet Infect Dis. 2016; 16: 712–723. https://doi.org/10.1016/S1473-3099(16)00026-8 PMID: 26874619
3. Halstead SB, Mahalingam S, Marovich MA, Ubol S, Mosser DM. Intrinsic antibody-dependent enhancement of microbial infection in macrophages: disease regulation by immune complexes. Lancet Infect Dis. 2010; 10: 712–722. https://doi.org/10.1016/S1473-3099(10)70166-3 PMID: 20883967
4. Halstead SB. Dengue Antibody-Dependent Enhancement: Knowns and Unknowns. Microbiol Spectr. 2014; 2.
5. Chau TN, Hieu NT, Anders KL, Wolbers M, Lien IB, Hieu LT, et al. Dengue virus infections and maternal antibody decay in a prospective birth cohort study of Vietnamese infants. J Infect Dis. 2009; 200: 1893–1900. https://doi.org/10.1086/648407 PMID: 19911991
6. Kikis SC, Nimmanitya S, Nisalak A, Burke DS. Evidence that maternal dengue antibodies are important in the development of dengue hemorrhagic fever in infants. Am J Trop Med Hyg. 1988; 38: 411–419. PMID: 3354774
7. Katzelnick LC, Gresh L, Halloran ME, Mercado JC, Kuan G, Gordon A, et al. Antibody-dependent enhancement of severe dengue disease in humans. Science. 2017; 358: 929–932. https://doi.org/10.1126/science.aan6836 PMID: 29097492
8. Murphy BR, Whitehead SS. Immune response to dengue virus and prospects for a vaccine. Annu Rev Immunol. 2011; 29: 587–619. https://doi.org/10.1146/annurev-immunol-031210-101315 PMID: 21219187
9. Vannice KS, Durbin A, Hombach J. Status of vaccine research and development of vaccines for dengue. Vaccine. 2016; 34: 2934–2938. https://doi.org/10.1016/j.vaccine.2015.12.073 PMID: 26973072
10. Sridhar S, Luedtke A, Langevin E, Zhu M, Bonaparte M, Machabert T, et al. Effect of Dengue Serostatus on Dengue Vaccine Safety and Efficacy. N Engl J Med. 2018.
11. Clark KB, Onlamoon N, Hsiao HM, Perng GC, Villinger F. Can non-human primates serve as models for investigating dengue disease pathogenesis? Front Microbiol. 2013; 4: 305. https://doi.org/10.3389/fmicb.2013.00305 PMID: 24130557
12. Sariol CA, White LJ. Utility, limitations, and future of non-human primates for dengue research and vaccine development. Front Immunol. 2014; 5: 452. https://doi.org/10.3389/fimmu.2014.00452 PMID: 25309540

13. St John AL, Abraham SN, Gubler DJ. Barriers to preclinical investigations of anti-dengue immunity and dengue pathogenesis. Nat Rev Microbiol. 2013; 11: 420–426. https://doi.org/10.1038/nrmicro3030 PMID: 23652323

14. Blaney JE Jr., Matro JM, Murphy BR, Whitehead SS. Recombinant, live-atenuated tetravalent dengue virus vaccine formulations induce a balanced, broad, and protective neutralizing antibody response against each of the four serotypes in rhesus monkeys. J Virol. 2005; 79: 5516–5528. https://doi.org/10.1128/JVI.79.9.5516-5528.2005 PMID: 15827166

15. Clements DE, Collier BA, Lieberman MM, Ogata S, Wang G, Harada KE, et al. Development of a recombinant dengue virus vaccine: immunogenicity and efficacy studies in mice and monkeys. Vaccine. 2010; 28: 2705–2715. https://doi.org/10.1016/j.vaccine.2010.01.022 PMID: 20997152

16. Fernandez S, Thomas SJ, De La Barrera R, Im-Erbsin R, Jarman RG, Baras B, et al. An adjuvanted, tetravalent dengue virus purified inactivated vaccine candidate induces long-lasting and protective antibody responses against dengue challenge in rhesus macaques. Am J Trop Med Hyg. 2015; 92: 698–706. https://doi.org/10.4269/ajtmh.14-0268 PMID: 25646261

17. Guirakhoo F, Pugachev K, Zhang Z, Myers G, Levenbook I, Draper K, et al. Safety and efficacy of chimeric yellow Fever-dengue virus tetravalent vaccine formulations in nonhuman primates. J Virol. 2004; 78: 4761–4775. https://doi.org/10.1128/JVI.78.9.4761-4775.2004 PMID: 15078958

18. Koraka P, Benton S, van AG, Stittelaar KJ, Osterhaus AD. Efficacy of a live attenuated tetravalent canarypox dengue virus vaccine in cynomolgus macaques. Vaccine. 2007; 25: 5409–5416. https://doi.org/10.1016/j.vaccine.2007.04.079 PMID: 17560694

19. Osorio JE, Brewoo JN, Silengo SJ, Arguello J, Moldovan IR, Tary-Lehmann M, et al. Efficacy of a tetravalent chimeric dengue vaccine (DENVax) in Cynomolgus macaques. J Am Trop Med Hyg. 2011; 84: 978–987. https://doi.org/10.4269/ajtmh.2011.09-0592 PMID: 21633037

20. Barban V, Mantel N, De MA, Pagnon A, Pradezynski F, Lang J, et al. Improvement of the Dengue Virus (DENV) Nonhuman Primate Model via a Reverse Translational Approach Based on Dengue Vaccine Clinical Efficacy Data against DENV-2 and -4. J Virol. 2018; 92.

21. Capeding MR, Tran NH, Hadinegoro SR, Ismail HI, Chotpitayasunondh T, Chua MN, et al. Effectiveness and safety of a novel tetravalent dengue vaccine in healthy children in Asia: a phase 3, randomized, observer-masked, placebo-controlled trial. Lancet. 2014; 384: 1358–1365. https://doi.org/10.1016/S0140-6736(14)61096-6 PMID: 25018116

22. Villar L, Dayan GH, Arredondo-Garcia JL, Rivera DM, Cunha R, Deseda C, et al. Efficacy of a tetravalent dengue vaccine in children in Latin America. N Engl J Med. 2015; 372: 113–123. https://doi.org/10.1056/NEJMoa1411037 PMID: 25365735

23. Borges MB, Marchevsky RS, Mendes YS, Mendes LG, Duarte AC, Cruz M, et al. Characterization of recent and minimally passaged Brazilian dengue viruses inducing robust infection in rhesus macaques. PLoS One. 2018; 13: e0196311. https://doi.org/10.1371/journal.pone.0196311 PMID: 29694440

24. Katzelnick LC, Montoya M, Gresh L, Balmaseda A, Harris E. Neutralizing antibody titers against dengue virus correlate with protection from symptomatic infection in a longitudinal cohort. Proc Natl Acad Sci U S A. 2016; 113: 726–733. https://doi.org/10.1073/pnas.1522136113 PMID: 26728879

25. Salje H, Cummings DAT, Rodriguez-Barruquer I, Katzelnick LC, Lessler J, Klungthong C, et al. Reconstruction of antibody dynamics and infection histories to evaluate dengue risk. Nature. 2018; 557: 719–723. https://doi.org/10.1038/s41586-018-0157-4 PMID: 29795354

26. Green S, Rothman A. Immunopathological mechanisms in dengue and dengue hemorrhagic fever. Curr Opin Infect Dis. 2006; 19: 429–436. https://doi.org/10.1097/01.qco.0000244047.31135.fa PMID: 16940865

27. Lee YH, Leong WY, Wilder-Smith A. Markers of dengue severity: a systematic review of cytokines and chemokines. J Gen Virol. 2016; 97: 3103–3119. https://doi.org/10.1099/jgv.0.00637 PMID: 27902384

28. Rothman AL. Immunity to dengue virus: a tale of original antigenic sin and tropical cytokine storms. Nat Rev Immunol. 2011; 11: 532–543. https://doi.org/10.1038/nri3014 PMID: 21760609

29. Srikiatkhachorn A, Mathew A, Rothman AL. Immune-mediated cytokine storm and its role in severe dengue. Semin Immunopathol. 2017; 39: 563–574. https://doi.org/10.1007/s00281-017-0625-1 PMID: 28401256

30. Wang XJ, Wei HX, Jiang SC, He C, Xu XJ, Peng HJ. Evaluation of aminotransferase abnormality in dengue patients: A meta analysis. Acta Trop. 2016; 156: 130–136. https://doi.org/10.1016/j.actatropica.2015.12.013 PMID: 26739659
31. Kumar Y, Liang C, Bo Z, Rajapakse JC, Ooi EE, Tannenbaum SR. Serum proteome and cytokine analysis in a longitudinal cohort of adults with primary dengue infection reveals predictive markers of DHF. PLoS Negl Trop Dis. 2012; 6: e1887. https://doi.org/10.1371/journal.pntd.0001887 PMID: 23209847
32. Libraty DH, Endy TP, Houng HS, Green S, Kalayanarooj S, Suntayakorn S, et al. Differing influences of virus burden and immune activation on disease severity in secondary dengue-3 virus infections. J Infect Dis. 2002; 185: 1213–1221. https://doi.org/10.1086/340365 PMID: 12001037
33. Vaughan DW, Green S, Kalayanarooj S, Innis BL, Nimmannitya S, Suntayakorn S, et al. Dengue viremia titer, antibody response pattern, and virus serotype correlate with disease severity. J Infect Dis. 2000; 181: 2–9. https://doi.org/10.1086/315215 PMID: 10608744
34. Paik T, Dutta SK, Mandal S, Saha B, Tripathi A. Differential clinical symptoms among acute phase Indian patients revealed significant association with dengue viral load and serum IFN-gamma level. J Clin Virol. 2014; 61: 365–370. https://doi.org/10.1016/j.jcv.2014.09.003 PMID: 25288310
35. Srikiatkhachorn A, Wichit S, Gibbons RV, Green S, Libraty DH, Endy TP, et al. Dengue viral RNA levels in peripheral blood mononuclear cells are associated with disease severity and preexisting dengue immune status. PLoS One. 2012; 7: e51335. https://doi.org/10.1371/journal.pone.0051335 PMID: 23284680
36. Bozza FA, Cruz OG, Zagne SM, Azeredo EL, Nogueira RM, Assis EF, et al. Multiplex cytokine profile from dengue patients: MIP-1beta and IFN-gamma as predictive factors for severity. BMC Infect Dis. 2008; 8: 86. https://doi.org/10.1186/1471-2334-8-86 PMID: 18578883
37. Brasier AR, Ju H, Garcia J, Spratt HM, Victor SS, Forshey BM, et al. A three-component biomarker panel for prediction of dengue hemorrhagic fever. Am J Trop Med Hyg. 2012; 86: 341–348. https://doi.org/10.4269/ajtmh.2012.12-0469 PMID: 23202872
38. Her Z, Kam YW, Gan VC, Lee B, Thein TL, Tan JJ, et al. Severity of Plasma Leakage Is Associated With High Levels of Interferon-gamma-Inducible Protein 10, Hepatocyte Growth Factor, Matrix Metalloproteinase 2 (MMP-2), and MMP-9 During Dengue Virus Infection. J Infect Dis. 2017; 215: 42–51. https://doi.org/10.1093/infdis/jiw94 PMID: 28077592
39. Kurane I, Innis BL, Nimmannitya S, Nisalak A, Meager A, Janus J, et al. Activation of T lymphocytes in dengue virus infections. High levels of soluble interleukin 2 receptor, soluble CD4, soluble CD8, interleukin 2, and interferon-gamma in sera of children with dengue. J Clin Invest. 1991; 88: 1473–1480. https://doi.org/10.1172/JCI115457 PMID: 1939640
40. Malavige GN, Huang LC, Salimi M, Gomes L, Jayaratne SD, Ogg GS. Cellular and cytokine correlates of severe dengue infection. PLoS One. 2012; 7: e50387. https://doi.org/10.1371/journal.pone.0050387 PMID: 23097331
41. Mustafa AS, Elbishbishi EA, Agarwal R, Chaturvedi UC. Elevated levels of interleukin-13 and IL-18 in patients with dengue hemorrhagic fever. FEMS Immunol Med Microbiol. 2001; 30: 229–233. https://doi.org/10.1111/j.1574-695X.2001.tb01573.x PMID: 1135143
42. Pandey N, Jain A, Garg RK, Kumar R, Agrawal OP, Lakshmana Rao PV. Serum levels of IL-8, IFN-gamma, IL-10, and TGF beta and their gene expression levels in severe and non-severe cases of dengue virus infection. Arch Virol. 2015; 160: 1463–1473. https://doi.org/10.1007/s00705-015-2410-6 PMID: 25860648
43. Sierra B, Perez AB, Alvarez M, Garcia G, Vogt K, Aguirre E, et al. Variation in inflammatory/regulatory cytokines in secondary, tertiary, and quaternary challenges with dengue virus. Am J Trop Med Hyg. 2012; 87: 538–547. https://doi.org/10.4269/ajtmh.2012.12-0531 PMID: 22802438
44. van Wilgenburg B, Scherwitzl I, Hutchinson EC, Leng T, Kurioka A, Kulicke C, et al. MAIT cells are activated during human viral infections. Nat Commun. 2016; 7: 11653. https://doi.org/10.1038/ncomms11653 PMID: 27337592
45. Yong YK, Tan HY, Jen SH, Shankar EM, Natkunam SK, Sathar J, et al. Aberrant monocyte responses predict and characterize dengue virus infection in individuals with severe disease. J Transl Med. 2017; 15: 121. https://doi.org/10.1186/s12967-017-1226-4 PMID: 28569153
46. Chaturvedi UC, Agarwal R, Elbishbishi EA, Mustafa AS. Cytokine cascade in dengue hemorrhagic fever: implications for pathogenesis. FEMS Immunol Med Microbiol. 2000; 28: 183–188. https://doi.org/10.1111/j.1574-695X.2000.tb01474.x PMID: 10865168
47. Pacsa AS, Agarwal R, Elbishbishi EA, Chaturvedi UC, Nagar R, Mustafa AS. Role of interleukin-12 in patients with dengue hemorrhagic fever. FEMS Immunol Med Microbiol. 2000; 28: 151–155. https://doi.org/10.1111/j.1574-695X.2000.tb01470.x PMID: 10799806
48. Trinchieri G. Cytokines acting on or secreted by macrophages during intracellular infection (IL-10, IL-12, IFN-gamma). Curr Opin Immunol. 1997; 9: 17–23. PMID: 9039773
49. Nakashishi K, Yoshimoto T, Tsutsui H, Okamura H. Interleukin-18 is a unique cytokine that stimulates both Th1 and Th2 responses depending on its cytokine milieu. Cytokine Growth Factor Rev. 2001; 12: 53–72. PMID: 11312119

Enhanced dengue virus infection in vaccinated macaques...
50. Walker W, Aste-Amezaga M, Kastelein RA, Trinchieri G, Hunter CA. IL-18 and CD28 use distinct molecular mechanisms to enhance NK cell production of IL-12-induced IFN-gamma. J Immunol. 1999; 162: 5894–5901. PMID: 10229825

51. Hoshino T, Wiltrout RH, Young HA. IL-18 is a potent coinducer of IL-13 in NK and T cells: a new potential role for IL-18 in modulating the immune response. J Immunol. 1999; 162: 5070–5077. PMID: 10227975

52. Iannello A, Samariani S, Debbache O, Tremblay C, Toma E, Boulasse M, et al. Role of interleukin-18 in the development and pathogenesis of AIDS. AIDS Rev. 2009; 11: 115–125. PMID: 19654853

53. Zhang SY, Boisson-Dupuis S, Chappier A, Yang K, Bustamante J, Puel A, et al. Inborn errors of interferon (IFN)-mediated immunity in humans: insights into the respective roles of IFN-alpha/beta, IFN-gamma, and IFN-lambda in host defense. Immunol Rev. 2008; 226: 29–40. https://doi.org/10.1111/j.1600-065X.2008.00698.x PMID: 19161414

54. Sierra B, Perez AB, Vogt K, Garcia G, Schmolke K, Aguirre E, et al. Secondary heterologous dengue infection risk: Disequilibrium between immune regulation and inflammation? Cell Immunol. 2010; 262: 134–140. https://doi.org/10.1016/j.cellimm.2010.02.005 PMID: 20219186

55. Gunther VJ, Putnak R, Eckels KH, Mammen MP, Scherer JM, Lyons A, et al. A human challenge model for dengue infection reveals a possible protective role for sustained interferon gamma levels during the acute phase of illness. Vaccine. 2011; 29: 3895–3904. https://doi.org/10.1016/j.vaccine.2011.03.038 PMID: 21443963

56. Malavige GN, Jeewananda C, Alles KM, Salimi M, Gomes L, Kamaladasa A, et al. Suppression of virus specific immune responses by IL-10 in acute dengue infection. PLoS Negl Trop Dis. 2013; 7: e2409. https://doi.org/10.1371/journal.pntd.0002409 PMID: 24040431

57. Tsai TT, Chuang YJ, Lin YS, Wan SW, Chen CL, Lin CF. An emerging role for the anti-inflammatory cytokine interleukin-10 in dengue virus infection. J Biomed Sci. 2013; 20: 40. https://doi.org/10.1186/1423-0127-20-40 PMID: 23800014

58. Guzman MG, Alvarez M, Halstead SB. Secondary infection as a risk factor for dengue hemorrhagic fever/dengue shock syndrome: an historical perspective and role of antibody-dependent enhancement of infection. Arch Virol. 2013; 158: 1445–1459. https://doi.org/10.1007/s00705-013-1645-3 PMID: 23471635

59. Chareonsirisuthigul T, Kalayanarooj S, Ubol S. Dengue virus (DENV) antibody-dependent enhancement of infection upregulates the production of anti-inflammatory cytokines, but suppresses anti-DENV free radical and pro-inflammatory cytokine production, in THP-1 cells. J Gen Virol. 2007; 88: 365–375. https://doi.org/10.1099/vir.0.82537-0 PMID: 17291552

60. Chen RF, Wang L, Cheng JT, Yang KD. Induction of IFNalpha or IL-12 depends on differentiation of THP-1 cells in dengue infections without and with antibody enhancement. BMC Infect Dis. 2012; 12: 340. https://doi.org/10.1186/1471-2334-12-340 PMID: 22216989

61. Tsai TT, Chuang YJ, Lin YS, Chang CP, Wan SW, Lin SH, et al. Antibody-dependent enhancement infection facilitates dengue virus-regulated signaling of IL-10 production in monocytes. PLoS Negl Trop Dis. 2014; 8: e3320. https://doi.org/10.1371/journal.pntd.0003320 PMID: 25412261

62. Ubol S, Halstead SB. How innate immune mechanisms contribute to antibody-enhanced viral infections. Clin Vaccine Immunol. 2010; 17: 1829–1835. https://doi.org/10.1128/CVI.00316-10 PMID: 20876821

63. Ubol S, Phuklia W, Kalayanarooj S, Modhiran N. Mechanisms of immune evasion induced by a complex of dengue virus and preexisting enhancing antibodies. J Infect Dis. 2010; 201: 923–935. https://doi.org/10.1086/651018 PMID: 20158392

64. Gil L, Martinez G, Tapanes R, Castro O, Gonzalez D, Bernardo L, et al. Oxidative stress in adult dengue patients. Am J Trop Med Hyg. 2004; 71: 652–657. PMID: 15569800

65. Martina BE, Koraka P, Osterhaus AD. Dengue virus pathogenesis: an integrated view. Clin Microbiol Rev. 2009; 22: 564–581. https://doi.org/10.1128/CMR.00035-09 PMID: 19622889

66. Krishnamurthi C, Kalayanarooj S, Cutting MA, Peat RA, Rothwell SW, Reid TJ, et al. Mechanisms of hemorrhage in dengue without circulatory collapse. Am J Trop Med Hyg. 2001; 65: 840–847. PMID: 11791984

67. Lee LK, Gan VC, Lee VJ, Tan AS, Leo YS, Lye DC. Clinical relevance and discriminatory value of elevated liver aminotransferase levels for dengue severity. PLoS Negl Trop Dis. 2012; 6: e1676. https://doi.org/10.1371/journal.pntd.0001676 PMID: 22679523

68. Mohan B, Patwari AK, Anand VK. Hepatic dysfunction in childhood dengue infection. J Trop Pediatr. 2000; 46: 40–43. https://doi.org/10.1093/troped/46.1.40 PMID: 10730040

69. Cui L, Pang J, Lee YH, Ooi EE, Ong CN, Lee YS, et al. Serum metabolome changes in adult patients with severe dengue in the critical and recovery phases of dengue infection. PLoS Negl Trop Dis. 2018; 12: e0006217. https://doi.org/10.1371/journal.pntd.0006217 PMID: 29364889
70. Senaratne T, Carr J, Noordeen F. Elevation in liver enzymes is associated with increased IL-2 and predicts severe outcomes in clinically apparent dengue virus infection. Cytokine. 2016; 83: 182–188. https://doi.org/10.1016/j.cyto.2016.04.010 PMID: 27155816

71. Beatty PR, Puerta-Guardo H, Killingbeck SS, Glasner DR, Hopkins K, Harris E. Dengue virus NS1 triggers endothelial permeability and vascular leak that is prevented by NS1 vaccination. Sci Transl Med. 2015; 7: 304ra141.

72. Glasner DR, Ratnasiri K, Puerta-Guardo H, Espinosa DA, Beatty PR, Harris E. Dengue virus NS1 cytokine-independent vascular leak is dependent on endothelial glycocalyx components. PLoS Pathog. 2017; 13: e1006673. https://doi.org/10.1371/journal.ppat.1006673 PMID: 29121099

73. Puerta-Guardo H, Glasner DR, Harris E. Dengue Virus NS1 Disrupts the Endothelial Glycocalyx, Leading to Hyperpermeability. PLoS Pathog. 2016; 12: e1005738. https://doi.org/10.1371/journal.ppat.1005738 PMID: 27416066

74. Rico-Hesse R. Microevolution and virulence of dengue viruses. Adv Virus Res. 2003; 59: 315–341. PMID: 14696333

75. Miagostovich MP, Sequeira PC, Dos Santos FB, Maia A, Nogueira RM, Schatzmayr HG, et al. Molecular typing of dengue virus type 2 in Brazil. Rev Inst Med Trop Sao Paulo. 2003; 45: 17–21.

76. Blaney JE Jr., Hanson CT, Firestone CY, Hanley KA, Murphy BR, Whitehead SS. Genetically modified, live attenuated dengue virus type 3 vaccine candidates. Am J Trop Med Hyg. 2004; 71: 811–821. PMID: 15642976

77. Durbin AP, Karron RA, Sun W, Vaughn DW, Reynolds MJ, Perreault JR, et al. Attenuation and immunogenicity in humans of a live dengue virus type-4 vaccine candidate with a 30 nucleotide deletion in its 3'-untranslated region. Am J Trop Med Hyg. 2001; 65: 405–413. PMID: 11716091

78. Whitehead SS, Hanley KA, Blaney JE Jr., Gilmore LE, Elkins WR, Murphy BR. Substitution of the structural genes of dengue virus type 4 with those of type 2 results in chimeric vaccine candidates which are attenuated for mosquitoes, mice, and rhesus monkeys. Vaccine. 2003; 21: 4307–4316. PMID: 14505913

79. Whitehead SS, Falgout B, Hanley KA, Blaney JE Jr., Markoff L, Murphy BR. A live, attenuated dengue virus type 1 vaccine candidate with a 30-nucleotide deletion in the 3' untranslated region is highly attenuated and immunogenic in monkeys. J Virol. 2003; 77: 1653–1657. https://doi.org/10.1128/JVI.77.2.1653-1657.2003 PMID: 12502885