Molecular epidemiology of dengue virus serotypes 2 and 3 isolated in Brazil from 1991 to 2008

Epidemiologia molecular dos sorotipos 2 e 3 do vírus dengue, isolados no Brasil de 1991 a 2008

Epidemiología molecular de los serotipos 2 y 3 del virus dengue, aislados en Brasil de 1991 a 2008

ABSTRACT

The dengue virus (DENV1-4) causes dengue fever and dengue hemorrhagic fever/dengue shock syndrome (DHF/DSS) in tropical and subtropical areas. The aim of this study was to evaluate the circulating genotypes of DENV. This was accomplished by sequencing the PrM and E genes of Brazilian isolates of DENV2 and DENV3 that were obtained between 1991 and 2008 from various geographic regions. Phylogenetic analyses of DENV2 demonstrated that the genotype III (Southeast Asian/American), in spite of several nucleotide and amino acid changes, was the only one that circulated over the past 19 years. Since its introduction in 2000, the DENV3 isolates that have been analyzed have all grouped into genotype III (Indian subcontinent) and there has been no evidence of DENV3 belonging to other genotypes in this study.

Keywords: Dengue Virus; Dengue Hemorrhagic Fever; Flavivirus.

INTRODUCTION

Dengue fever (DF) and dengue hemorrhagic fever/dengue shock syndrome (DHF/DSS) are major health problems in the tropical and subtropical regions. The increasing number of patients with severe clinical manifestations and the expansion of epidemic areas have led to extensive research on its causative agent, dengue virus (DENV), which is the most widespread and important arthropod-borne virus in terms of morbidity and mortality

2,3. A global dengue pandemic, which started during World War II, has progressively spread worldwide and involves predominantly tropical countries. According to the World Health Organization

4, dengue fever results in approximately 5 million hospitalized children and at least 50 thousand deaths are caused by DHF/DSS annually.

The DENV serotypes 1 to 4 (DENV1 to DENV4) belong to the Flavivirus genus of the family Flaviviridae. The genome of flaviviruses is a positive-sense, single-stranded RNA approximately 11 Kb in length, from which a single polyprotein is produced. This polyprotein is...
proteolytically processed by cellular and viral proteases to generate ten viral proteins. The gene order of the dengue polyprotein is 5’-C-prM(M)-E-NS1-NS2A-NS2B-NS3-NS4A-NS4B-NS5-3’.

Dengue infections can be manifested from asymptomatic to subclinical pictures through DF and DHF/DSS, all of which may be observed during a dengue outbreak. Nonetheless, dengue infections usually result in two well-defined syndromes, DF and DHF. The mechanism by which the DENV-infected host exhibits benign or severe disease remains to be elucidated. The antibody-dependent enhancement (ADE) pathway postulates that circulating antibodies from a primary infection bind to a heterologous virus in a secondary infection facilitating penetration of virus particles into mononuclear cells. Further studies have shown that host immune system mediators, such as cytokines and complement, may have a direct role in the pathogenesis of plasma leakage, a defining feature of DHF.

In Brazil, DENV2 was first isolated in 1989 from an imported case from Uganda, Africa. Prior to that case, DENV1 and DENV4 caused an epidemic in Boa Vista, Roraima State, Brazil. In 1986 DENV1 caused epidemics in Rio de Janeiro and additional cities. In 1990, the first autochthonous DENV2 outbreak occurred in the State of Rio de Janeiro. The introduction of DENV2 increased the number of severe disease cases with the appearance of several cases of DHF/DSS. Furthermore, DENV3 was isolated in São Paulo State from an imported case that occurred in 2000. One year after its introduction into Brazil, DENV3 caused a large dengue outbreak in Rio de Janeiro State and has quickly spread to several other Brazilian States.

In this study, we have sequenced the C, prM/M and E genes of both DENV2 and DENV3 in 27 Brazilian strains from distinct geographic areas and at different periods of collection between 1991 and 2008 to investigate the molecular epidemiology of these serotypes that are circulating in the country.

MATERIALS AND METHODS

VIRUS ISOLATION

Twenty-seven DENV strains (14 DENV2 and 13 DENV3) were obtained from the virus collection at the Department of Arbovirology and Hemorrhagic Fever in the Instituto Evandro Chagas (IEC). These viruses were isolated from sera collected from patients that were clinically diagnosed as cases of DF, DHF/DSS and encephalitis during the outbreaks that occurred between the period of 1991 to 2008 and from mosquitoes captured in the endemic area (Figure 1). Viruses were grown in cultured Aedes albopictus cells (clone C6/36). Relevant information on each isolate is summarized in table 1.

RNA EXTRACTION AND RT-PCR AMPLIFICATION

Genomic RNA samples were extracted using Trizol reagent (Invitrogen, USA) according to the manufacturer’s instructions. cDNAs were synthesized and amplified with a standard two step RT-PCR assay with specific primers for DENV2 and DENV3 that were designed to generate overlapping products corresponding to the C/prM/M/E structural genomic region as previously reported. Initially, 0.2-1 μg of viral RNA and 20 μM of antisense primer were heated at 90 °C for 90 s and then placed on ice. RT mix containing 1X RT buffer (200 mM Tris-HCl (pH 8.4), 500 mM KCl), 0.2 mM dNTP (deoxynucleotides) mixture, 100 mM DTT (dithiothreitol), 40 U RNAse inhibitor (Invitrogen, USA) and 200 U Superscript II Reverse transcriptase (Invitrogen, USA) was added to the pre-heated RNA and the volume adjusted to 20 μL. The RT reaction was carried out at 45 °C for 1 h, followed by heating at 94 °C for 10 min. The RT products were used as templates for PCR amplification. Reactions were composed of 20 μM each of the forward and reverse primers, 1X PCR buffer, 1 mM of MgCl2, 0.2 mM of dNTP mixture, 5 U/μL of Platinum Taq Polymerase (Invitrogen) and the final volume adjusted to 50 μL with RNase-free water. The samples were placed in a thermal cycler (Perkin Elmer 9600, USA) at 94 °C for 90 s (for initial denaturation) followed by 35 cycles of: 94 °C for 30 s, 55 °C for 30 s, and 72 °C for 120 s and a final extension step at 72 °C for 5 min. RT-PCR products were visualized in a 1.5% agarose gel stained with ethidium bromide (5 μg/mL).
NUCLEOTIDE SEQUENCING AND SEQUENCE ANALYSES

The RT-PCR products were purified using the QIAquick gel extraction kit (Qiagen, Inc., Valencia, CA). Purified cDNA was used as the template for sequencing using the Big Dye Terminator 3.0 kit (Applied Biosystems, Inc., USA) according to the manufacturer’s instructions. Sequencing was performed using the ABI Prism 377 (Applied Biosystems) equipment. Nucleotide sequences were analyzed and edited with SeqMan software (DNASTAR, Lasergene software package). Phylogenetic analysis were generated by progressive pairwise multiple sequence alignments using CLUSTAL W (Megalign software; Lasergene software package DNASTAR).

A 2271-base sequence spanning the C-prM-E structural protein genes was used for comparing the 14 isolates described here and 22 previously published sequences of DENV2 retrieved from GenBank (http://www.ncbi.nlm.nih.gov). For DENV3, 1,977 bases from the M and E genes were used to compare our 13 isolates with the 27 previously published sequences of DENV3. The DENV2 sequence (strain Jamaica access number M20558) was used as an outgroup.

Phylogenetic analyses were performed with bootstrap values of 1 thousand pseudoreplicas using the neighbor-joining method (NJ) Kimura-2-parameter (MEGA 2.1 software).

RESULTS

ANALYSIS OF DENV2

We have determined the full nucleotide sequence of the structural genome of 14 DENV2 isolates from patients exhibiting different patterns of disease severity (Table 1). Comparative differences in nucleotide numbers between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaica (Jam83) strain revealed high nucleotide sequence similarity (89.0 to 99.9%) among our isolates. The average similarity between all viruses studied and the DENV2 Jamaic...
The prM/M/E amino acid sequences of all 14 DENV2 isolates from Brazil showed that these proteins were highly conserved. Only five positions in the prM/M/E genes varied in the five isolates, the most variable being prM amino acid 14, which was a glycine (V→G) in four strains. Isolate BeH527822 accumulated two additional changes at prM/M 58 (Q→P) and 60 (E→D) as shown in Table 2. In the E gene, 12 amino acid positions were variable with E53 (P→L) and E269 (E→K) being observed in four and two isolates, respectively. The most variable isolates were BeH527822, BeH666426 and BeH547176 with changes at four positions where the only one that was common to them all was E53 (P→L) (Table 2).
Table 2 – Differences of amino acids among the prM/E genes sequences of Brazilian DENV2

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* Amino acid number from protein amino terminus; † Sample code.

ANALYSIS OF DENV3

Nucleotide sequence homology among the DENV3 strains ranged from 89.7 to 100%. The deduced prM/M and E amino acid sequences for all ten DENV3 isolates showed that their proteins were highly conserved, with a similarity ranging from 93.2 to 100%. The average sequence homology was 94.8% and 96.6% for the nucleotide and amino acid sequences respectively.

The phylogenetic tree constructed for the DENV3 isolates was based on the alignment of sequences at the E/NS1 region (992-2550 nt), using the criteria for division of the genotypes described by Lanciotti. All DENV3 isolates were identified as members of genotype III (Indian subcontinent) and clustered separately in two subclades (A and B). Subclade A contains the Latin American strains and is genetically more closely related to subclade B strains isolated in Sri Lanka (L11437, L11438) (Figure 3).

The variability of amino acid sequences for the DENV3 isolates was restricted to one position in the M gene (M86 H → R), whereas six positions were found to vary in the E gene (Table 3). The most variable were E6 (I → V), E89 (Q → P) and E246 (Q → P). Isolates BeH657637 and BeH665993 had accumulated changes in three of the six variable amino acid positions (Table 3).

Table 3 – Differences of amino acids among the M/E genes sequences of Brazilian DENV3

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* Amino acid number from protein amino terminus; † Sample code.
DISCUSSION

The nucleotide sequence divergence (10.5%) obtained in this study for DENV2 suggests an accumulation of mutations during the period of high circulation of this DENV serotype\(^{26}\). Indeed, DENV2 has circulated in Brazil for almost two decades; however, there has been a surprisingly low incidence of DHF cases when compared to other countries. After introduction of DENV3 in 2002, the number of DHF has increased sharply, suggesting a high virulence of this serotype\(^{26}\). Previous studies on DENV2 from the Americas had shown that the appearance of DHF was related to the introduction of an Asian genotype into the Caribbean Region\(^{27}\). Our data confirms that this DENV2 genotype has been predominant in Brazil throughout the 19 years of its circulation.

According to the classification of Rodriguez-Roche et al\(^{22}\) and Vasilekos and Weaver\(^{23}\), the DENV2 is grouped into three genotypes, with the circulation of genotypes III and IV in the Americas. Genotype III circulated in all countries of the American Region having, as its prototype, the Jamaican strain, which has been associated with DHF cases. The genotype IV (American) has been associated with strains of lower virulence and only associated with DF. It is of interest that this genotype has circulated only in the American subcontinent\(^{28}\). In our study, none of the Brazilian sequences studied were included in the DENV2 genotype IV (American). In fact, all of sequences were included in the genotype III (Figure 2).

The first studies comparing nucleotide sequence of the gene of 12 DENV2 showed no correlation between disease
severity and specific nucleotide or amino acid sequence. Further analysis of the complete genome of the Southeast Asian DENV2 also failed to identify specific sites that could determine virulence. Leitmyer et al. conducted a more consistent analysis of the genetic diversity of DENV2. This study identified an amino acid substitution at position E390 as a primary determinant of severe dengue. In the American genotype, the cases of DF were found to be associated with the presence of an aspartic acid (Asp-D), which may alter interactions with cellular receptors. However, the results obtained with 14 sequences of DENV2 in our study identified an Asn at position E390 in all the sequences studied when the sequences were compared within the 14 strains and with strains of Asian and American genotypes. The results indicate that the Brazilian DENV2 analyzed have the potential to cause DHF, which is consistent with the data reported with isolates from Venezuela.

The change of amino acid E390 is located in the carboxyl-terminal domain III (aa 303-395) on the lateral surface, which is believed to contain residues that are involved in tropism and virulence in different flaviviruses. In the E gene of 14 Brazilian viruses analyzed, 12 changes were present, and only five amino acids substitutions were significantly associated with a change of character and polarity. Those positions were: E208 (Val/V → Glu/E), 262 (Thr/T → Lys/K), 273 (Ser/S → Leu/L), 274 (Ser/S → Ala/A) and 405 (Thr/T → Pro/P). All amino acid changes are shown in table 2. The changes were present in the ectodomain III or outside of the ectodomain of the protein such as in the residue 405 of sample H666426/Goi4191. Some changes shown in table 2, such as E262 and 264 that are present in H547176/ROR1811 and the viral strain from Venezuela (AF100466), suggest that the virus may have originated from this Country, thus confirming the existence of two modes of entrance of the DENV2 into Brazil.

Over the past two decades, DENV3 has caused epidemics of DHF in Southeast Asia, East Africa and Latin America. The first phylogenetic analysis of the E protein of this dengue serotype resulted in the recognition of four distinct genotypes. In general, genotypes can be grouped by distinct geographic origin: American, Indian subcontinent, Thailand and Southeast Asian/South Pacific. The DENV3 genotype that circulated in the Americas until 1989 had low epidemic potential and was isolated only from DF patients. Between 1980 and 1990, two new genotypes were introduced into the South Pacific and Americas: the Southeast Asian genotype (genotype I) is associated with large epidemics of DHF in Tahiti and Fiji, and the Indian subcontinent genotype (genotype III) was introduced into Central America in the 1990s in Nicaragua.

In 1994 DENV3 resurfaced in the Americas, causing a small outbreak associated with classic DF in Panama. The virus spread toward Northern Central America and reaching Nicaragua and Mexico. Seven years after its introduction into the Americas, DENV3 spread throughout South America reaching Venezuela, Paraguay and Brazil and causing large epidemics.

The 13 Brazilian DENV3 isolates from our study that were collected from different geographical regions, together with those obtained from neighboring countries, have all been grouped into genotype III (Indian subcontinent). They are genetically different from the DENV3 strain that previously circulated in the Americas in the 1960s belonging to the genotype V. It is possible that this accumulated variability is the result of the rapid dispersion and increase in viral replication. This hypothesis is supported by the bootstrap values of around 84% shown in the phylogenetic tree based on the neighbor-joining method (Figure 3). The variability of the DENV3 (genotype III) strain may be due to multiple introductions into Brazil, which give rise to subgroups according to origin and year of isolation of the viral strain (Figure 3).

Local transmission of DENV3 was initially detected in Rio de Janeiro State in December 2000. In the summer of 2001, the first autochthonous epidemic occurred in Rio de Janeiro City, with a large number of DHF cases, followed by the spread of this genotype throughout Brazil. All viruses isolated belonged to genotype III, similar to that circulating in Sri Lanka, which is in agreement with previously described studies. Our results show, based on the chronology of the virus isolation, that the dispersion of the DENV3 genotype III in Brazil occurred from the Southeast to the Northern region. These results differ from those of Figueiredo et al., which recorded the circulation of the DENV3 genotype I in the State of Minas Gerais, Southeast to Brazil and Araújo et al. suggesting the formation of a new genotype (V) that grouped the DENV3 strains from Brazil, China and Japan. The differences in results show that the circulation of new genotypes appeared to be restricted to a specific region in Brazil or to a certain period of time. This raises some important questions: Why have other strains of these genotypes failed to be identified? Does this strain have a potential transmission vector? Therefore, the emergence of other genotypes or novel genotypes should be further investigated. This should include analyzing the complete genome sequences of DENV3 and its epidemiological profile.

CONCLUSION

In summary, our study suggests that the low genetic variability for DENV2 circulating in Brazil may explain the low incidence of severe cases of the disease that have been reported to date. However, the explosive spread of DENV2 genotype III, which has been found to be more variable than DENV3, may have accumulated genetic changes associated with an increase of virulence, immune status and other important factors previously described. This would explain the higher number of severe dengue cases since the introduction of DENV2 and DENV3 into Brazil.
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Epidemiologia molecular dos sorotipos 2 e 3 do vírus dengue, isolados no Brasil de 1991 a 2008

RESUMO
O vírus dengue (DENV1-4) causa a dengue clássica e a febre hemorrágica da dengue / síndrome de choque da dengue (FHD/SCD) em regiões tropicais e subtropicais. O objetivo deste estudo foi avaliar os genótipos circulantes de DENV2 e DENV3 obtidos em distintas regiões geográficas no período de 1991 a 2008. As análises filogenéticas de DENV2 demonstraram que o genótipo III (Sudeste da Ásia/América), apesar das diversas alterações nucleotídicas e de aminoácidos, foi o único a circular durante os últimos 19 anos. Desde a sua introdução no estudo, em 2000, todas as amostras isoladas de DENV3 analisadas foram agrupadas no genótipo III (subcontinente indiano). Não foram encontradas evidências de que o DENV3 pertença a outros genótipos investigados.

Palavras-chave: Virus da Dengue; Febre Hemorrágica da Dengue; Flavivirus.

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