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Genomic constellations of RVA detected in Brazil from 1986 to 2016: a temporal and geographical distribution and occurrence of reassortments

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ABSTRACT

Introduction: Species A rotavirus (RVA) infections are a major cause of severe gastroenteritis in children of <5 years worldwide. In Brazil, before vaccination, RVA was associated with 3.5 million episodes of acute diarrheal disease per year. Due to the segmented nature of their genomes, rotaviruses can exchange genes during co-infections, and generate new virus strains and new reinfections. **Objective:** To evaluate the genomic diversity of RVA isolated in Brazil in 30 years, between 1986 to 2016, to investigate possible changes in the frequency of genotype constellations before and after the implementation of the vaccine. Methods: In total, 4,474 nucleotide sequences were obtained from the Virus Variation Database. Genomic constellation was compared, and the proportion of rotavirus genotypes was analyzed by time and geographic region. **Results:** Our results showed that major known genotypes were circulating in the country during the period under analysis, with a prevalence of the G1P[8] Wa-like genotype, decreasing only in the period immediately after the introduction of the vaccine. Regarding the geographical distribution, most of our constellations, 62 (39.2%), and 50 (31.6%) were concentrated in the North and Northeast regions. Our analysis also showed the circulation of multiple strains during the periods when the DS-1-like and AU-1-like genotypes were co-circulating with the Wa-like genotype. **Conclusion:** Therefore, it is likely that inter-genogroup reassortments are still occurring in Brazil and so it is important to establish an efficient surveillance system to follow the emergence of novel reassorted strains that might not be targeted by the vaccine.

Keywords: rotavirus; gene rearrangement; genomic library; phylogeny; demography; vaccines.

INTRODUCTION

Rotavirus A (RVA) remains the main viral agent that causes acute gastroenteritis in children \leq 5 years old worldwide, affecting children from both developed and developing countries¹⁻⁵. In Brazil, before the vaccine introduction, RVA was responsible for approximately 650,000 outpatient visits, 92,000 admissions, and 850 deaths per year in children under five years of age⁶. Due to the importance of RVA, in 2006 two vaccines with proven efficacy were licensed. In the same year, one of these vaccines was implemented in the National Immunization Program in Brazil, the monovalent based on G1P[8] rotavirus vaccine Rotarix® (GlaxoSmithKline Vaccines, Rixensart, Belgium)⁷. The vaccine was available free of charge to all children of eligible age to reduce the number of deaths caused by gastroenteritis in children. Surveillance studies show that the goal was achieved by reducing considerably the number of hospitalizations and deaths related to gastroenteritis, especially in children up to 12 months⁷⁻⁹.

RVA possesses a double-stranded RNA genome with 11 gene segments. The segment of the genome enables the reassortment between and within human and animal strains, favoring greater genomic diversity of this virus¹⁰.

The traditional classification is based on the genes that encode the outer capsid proteins, VP4 (P-genotype) and VP7 (G-genotype)¹⁰. More recently, the genome classification of RVA strains has been enlarged to include all 11 genes: Gx-P[x]-Ix- Rx-Cx-Mx-Ax-Nx-Tx-Ex-Hx concerning the sequences of the genes VP7-VP4-VP6-VP1-VP2-VP3-NSP1-NSP2-NSP3-NSP4-NSP5/6, respectively¹¹. Based on this classification, the human RVAs possess one of the following genotype constellations along with the G and P genotypes: Wa-like (I1-R1-C1-M1-A1-N1-T1-E1-H1) of porcine origin, DS-1-like (I2-R2-C2-M2-A2-N2-T2-E2-H2) of the bovine origin or, more rarely, AU-1-like (I3-R3-C3-M3-A3-N3-T3-E3-H3) possibly with canine or feline origin¹⁰⁻¹⁴.

The Wa-like genotype is responsible for more than 50% of diarrhea cases in children worldwide and thus has a fundamental role in the emergence of new cases^{15,16}. Also in Brazil, previous studies have shown a prevalence of the G1P[8] Wa-Like genotype. However, some reassortment events between different constellations have been described previously, forming new strains that may be of epidemiological importance but are not identified with the analysis of only G and P genotypes^{10,17}. The classification of the entire viral genome makes it possible to visualize evolutionary events that are not under pressure from the vaccine, but that may be of great clinical or epidemiological importance described previously there are already some studies describing the genotype constellation of strains of RVA circulating in Brazil^{10,17,19}, there is still no study that makes a long temporal evaluation of all available strains to verify temporal changes in the genotypes. In addition, Brazil is a country that has a continental proportion and there are no studies that verify the regional differences in the RVA genomic constellation over time.

Therefore, this study aimed to evaluate the genomic diversity of RVA isolated in Brazil over a period of 30 years to investigate possible temporal and/or regional changes in the genomic constellation prevalence.

METHODS

Data collection and development of a local database

To create a local database, 4,474 nucleotide sequences, with a length between 201 to 3302 bp of the 11 segments RVA infecting the human host and 158 fully sequenced RVA isolates between the years of 1986-2016, with sequence length between 528 to 3302 bp, were collected from the Rotavirus database at the Virus Variation Resource

(https://www.ncbi.nlm.nih.gov/genome/viruses/variation/). GenBank (https://www.ncbi.nlm.nih.gov/genbank/) was used to retrieve information from the country region to which the strains belonged. A local database was developed to store the nucleotide sequences in FASTA and XLSX format along with the DNA content information, GenBank accession numbers, collection date, segment, country region, and isolate/strain.

Analysis of the gene segments and constellations for time and geographic region

The distribution of gene segments and rotavirus genotype constellations in Brazil was analyzed before and after the vaccine introduction in two periods, 1986-2006 and 2006-2016, respectively. In addition, among all RVA strains, 158 fully sequenced, with ORF between 528 to 3302 bp, were recovered and classified into genomic constellation Wa-like, DS-1-like, AU-1-like, or mixed genotypes.

The proportion of gene segments and rotavirus genomic constellations were analyzed by time before the vaccine introduction (1986-1995, 1996-2000, and 2001-2006) and after (2007-2008, 2009-2011, and 2012-2016). Also were analyzed by geographic region of the country. Brazil has five geographic regions, South and Southeast (subtropical/tropical climate and high socioeconomic indicators); Northern and Northeast (equatorial/tropical climate and low socioeconomic indicators); and Central-West (tropical climate and intermediate socioeconomic indicators).

Sequence alignments and phylogenetic reconstruction of the RVA sequences

Nucleotide sequences for each rotavirus genome segment were aligned using the Muscle algorithm with the default parameters²⁰, which is incorporated into the MEGA5 software²¹. The phylogenetic relationships of RVA sequences were determined by

Bayesian inference using MrBayes v3.2.7^{22,23}, with 12,000,000 generations for the Markov chain Monte Carlo (MCMC) algorithm. A 25% discarded burnin was set to eliminate iterations at the beginning of the MCMC run. For Bayesian Inference tree reconstruction, the general time reversible (GTR) model using Gamma distribution (+G) and the proportion of invariable sites (+I) was used, which was indicated by jModelTest v2.1.10²⁴. This evolutionary model was set as the substitution rates of variation of the sequences for NSP1, VP1, VP3, VP4,0, and VP7 datasets. For NSP2 and NSP4 segments, the 3-parameter model (TPM)1uf+G and TPM2uf+I+G model was used, respectively. For NSP3 and VP2 segments, the transitional model (TIM)1+I+G and TIM3+G were used, respectively. The model used for phylogenetic reconstruction of the NSP5 and VP6 segments was Hasegawa-Kishino-Yano (HKY)+G and Tamura-Nei (TRN)+I+G, respectively. The tree for each viral segment was edited and visualized with Itol v 5.6.3 (https://itol.embl.de/).

RESULTS

A total of 4,474 nucleotide sequences were obtained from the Rotavirus database at the Virus Variation Resource. Regarding the length of the sequences, 4,214 had a length above 500 bp. The sequences smaller than 500 bp were distributed among the 10 RVA genotypes. The NSP1, NSP2, and NSP3 genotypes, with one sequence, respectively. Genotypes VP4, VP1, VP3, and VP7 had two, five, five, and seven sequences, respectively, with 201 to 425 bp. The VP2, NSP4 and VP6 genotypes had the most sequences with less than 500 bp. For VP2 38 sequences were found, with lengths>425 and <500 bp. The NSP4 and VP6 genotypes presented, respectively, 52 and 148 sequences, with lengths>209 and <500 bp.

About geographical distribution, 1559 (34.8%) sequences were from the North region, 1098 (24.5%) were from the Northeast, 1179 (26.4%) were from the Southeast, 322 (7.2%) were from the South, 276 (6.2%) from the Central-West and 40 (0.9%) had no information about the region of origin (Supplementary file 1). In the North and Northeast regions, most of the gene segments obtained were from the post-vaccine period, with a predominance of the VP7 (217) and VP1 (203) genotypes for the North, and VP7 (186) and VP4 (177) for Northeast. The years 2009 and 2011 presented the largest number of representatives (Figure 1a, 1b). In Central-West, Southeast, and South regions, most of the gene the pre-vaccination period, especially between the years 2001 to 2006 (Figure 1c and 1e).

A total of 769 sequences reported G genotypes (*VP7* gene). G1 (241; 31.3%), G2 (232; 30.2%), and G12 (106; 13.8%) were the most frequent, followed by G9 (89; 11.6%), G3 (58; 7.5%), and G5 (29; 3.8%). Other strains included G4, G8, G10 and G26 (total of 14; 1.8%). Regarding the P genotype, 695 sequences were obtained: P[8] (426; 61.3%), P[4] (185; 26.6%), P[6] (61; 8.8%), P[9] (21; 3.0%), and the genotypes P[3] and P[19] with one sequence each (0.1%) (Figure 2). Figure 3 presents the cumulative distribution of the genotype combinations for the past 30 years. It was possible to identify 54 combinations of G and P genotypes. The most frequent combinations were G1P[8] (6.1%), G2P[8] (6%), G12P[8] (5%), G9P[8] (4.8%), G3P[8] (4.4%), G5P[8] (4.2), G1P[4] (4%), G4P[8] (4%), G8P[8] (4%), G10P[8] (4%), and G2P[4] (3.8%). There were 43 other G and P combinations representing 49.7% of the strains.

Based on the genome constellation classification, 158 human RVA strains detected in Brazil were distributed following the reference genome constellations: Wa-like, DS-1-like, or AU-1-like or mixed. In total, 120 (75.9%) Brazilian strains belonged to the Wa-like genome constellation, 25 (15.8%) belonged to the DS-1-like, 4 (2.5%)

belonged to the AU-1-like, and 9 (5.7%) were Wa-like mixed with DS-1-like and/or AU-1-like (Supplementary file 2).

Regarding mixed genomes (n=9), all had the prevalence of the Wa-like genome with reassortment events in specific genes forming the mixed genomes. Of these, 6 (66.7%) genomes were Wa-like + DS-1-like, four with T2 genotype (*NSP3* gene), one with R2 genotype (*VP1* gene), and one with M2, N2 and T2 genotypes (*VP3*, *NSP2* and *NSP3* genes, respectively). In addition, 2 (22.2%) genomes were Wa-like + Au-1-like, one with R3 genotype (*VP1* gene) and the other with T3 genotype (*NSP3* gene). One genome (11.1%), with reassortment among the three genomic genotypes, was also identified, Wa-like + DS-1-like + AU-1-like, containing the M3 and T2 genotypes (*VP3* and *NSP3*, respectively) (Supplementary file 2).

Out of the Wa-like genotypes, 108 (90%) were classified as G1P[8] genotype, and the other 12 (10.0%) were classified as G12P[6] genotype. The DS-1-like genomes were G12P[6] (22; 88.0%) and G2P[4] (3; 12.0%), whereas all AU-1-like genomes (4; 100.0%) were G3P[9] (Supplementary file 2).

The analysis of the genomic constellation by time and region showed that most of the isolates, 62 (39.2%), and 50 (31.6%), were concentrated in the North and Northeast regions and the period 2009-2011, with 59 (37.3%) and 28 (17.7%) constellations, respectively. The Southeast and South regions presented 28 (17.7%) and 12 (7.6%) constellations, respectively. Most of the isolates in the Southeast and South regions were identified between 1996-2000 and 2001-2006. In the period 1996-2000, only constellations identified in the Southeast region were registered in the analyzed database.

They represented 13 (8.2%) of the total constellations analyzed in this study. The number of constellations identified in the Southeast and South regions in the period 2001-2006 was 9 (5%) and 10 (6.3%), respectively. The periods 1986-1995, 2007-2008, and

2012-2016 had the lowest number of constellations identified, 9 (5%), 6 (3.8%), and 2 (1.3%), respectively. The Central-West region with 6 (3.8%) constellations presented the lowest frequency identified.

Based on the genomes deposited in the database, before the introduction of the vaccine, only the Wa-like and AU-1-like constellations were present in Brazilian strains, with the period 2001-2006 housing 41 of the 64 constellations present in the 1986-2006 time interval (Figure 4). After the introduction of the vaccine, the Wa-like constellations showed a small increase and the strains of the DS-1-like constellation appeared for the first time in the country. The period 2009-2011 presented the largest number of representatives, with 59 constellations of the Wa-like type and 22 DS-1-like. The number of mixed constellations doubled after the introduction of the vaccine, from three to six (Figure 4).

The phylogenetic analysis of RVA based on each of the 11 gene segments has shown that all RVA genes presented a specific pattern of clustering of the isolates from Brazil according to the genotypes and their respective lineages. It was also possible to observe the existence of a temporal pattern of clustering for the lineages in all reconstructed phylogenies. In addition, the internal groups of most lineages presented a regional cluster. However, the temporal clustering was the most evident (Supplementary file 3-13). When analyzing the phylogeny of NSP1, it was possible to identify a cluster of sequences belonging to the South (KM026548), Southeast (KM026553), and Midwest (KM026547) regions, that clustered together in black. The [P8] genotype of VP4 also showed this clustering pattern. In this segment, an orange sequence (KM027056) from the Southeast region is highlighted, clustering together with sequences from the Midwest region (KM027057, KM027059, and JX437037), in yellow. These clusters highlighted here had in common the period, 1988-1998. Other segments analyzed also presented this

clustering pattern, highlighted sequences, or marked with different colors within the same genotype. A sequence in A3 of NSP1 (JQ715659) from the State of Pará did not present a clustering pattern based on year or geographical region. The posterior probabilities for the external nodes were above 95% and, for most internal nodes present in the 11 gene segments, the posterior probability values were greater than 75%. The VP4 segment had the lowest branch support for several internal nodes of the phylogenetic tree (Supplementary files 3-13).

DISCUSSION

Despite the existence of two licensed and effective vaccines, RVA remains an important agent of acute gastroenteritis and diarrhea-related deaths in children under 5 years of age, especially in undeveloped countries¹⁻⁴. In Brazil, the vaccine has been available since 2006 and has maintained high levels of coverage, which has caused a decrease in the number of hospitalizations and deaths from diarrhea in the country⁹. However, despite the vaccine's effectiveness, new cases of RVA are reported annually in several studies^{10,25-28}. Based on the analyzed data, during 30 years of surveillance for Brazilian RVA, G1P[8] genotype was the most prevalent when considering the entire study period, but there is a sharp decrease in the circulation of this genotype shortly after the introduction of the vaccine in the country (2007-2008 period). In the same period, it is also possible to see an increase in G2P[x] genotypes, especially G2P[4], which is a heterotypic strain compared to the vaccine Rotarix® (G1P[8] strain) (Figure 3). This decrease in the G1P[8] genotype and increase in the G2P[4] between 2007-2008 was also seen in countries where the Pentavalent Rotateq® vaccine was introduced, such as Australia²⁹, as well as countries that had not introduced any vaccine against RVA, such

as Argentina³⁰, which demonstrates that these fluctuations of genotypes over time can be a natural viral mechanism of maintenance in the human population³¹.

Before the vaccine, G1P[8] represented 6.7%, 14.4%, and 33.9% of the strains in the 1986-1995, 1996-2000, and 2001-2006 periods, respectively. After the vaccine introduction, it represented 1.3%, 32.9%, and 10.6% of the strains in 2007-2008, 2009-2011, and 2012-2016, respectively. These variations in the prevalence of the G1P[8] genotype, with a decrease mainly in the periods following the start of vaccination, have already been described by Santos et al.²⁵ in a systematic review of publications between 1986-2015. Here, we have analyzed the nucleotide sequences for all 11 RVA genes, and the genomic constellations, in databases and evaluated their distribution by time (1986-2016) and by geographic region of the country. Therefore, we conclude that there is no linear reduction of the frequency of this genotype in the total circulating RVA after the implementation of the vaccine, although there is a reduction in the total number of deaths and hospitalizations due to infection with this virus.

The Wa-like genomic constellation was the most prevalent during the entire study period, a fact that was already expected considering that this is the genotype that is most frequently reported in human infections around the world³²⁻³⁶. However, in specific periods, the other genotypes were also identified in Brazil. AU-like-1, which is a rarer genotype, was only found in the period between 1996-2000, and the DS-1-like genotype was identified in two consecutive periods analyzed (2007-2008 and 2009-2011). Despite low frequencies, mixed genotypes, which had reassortment events, were also identified in the same periods in which the genotypes DS-1-like and AU-like-1 were identified (Figure 4).

Some DS-1-like and AU-like-1 genotypes, along with mixed genotypes, have already been described in other studies in Brazil as well as in other regions of the

world^{16,18,22,24,35,37}. Recently, the spread of a mixed equine-like G3P[8] strain in Brazil has been identified, showing that even recently these mixed strains are still emerging and need to be identified³⁷.

Some studies have reported the existence of inter-genogroup reassortments among different animal and human RVA genogroups^{14,15,38}. Reassortment events have been reported in the NSP3 gene in strains from Maranhão and Rio de Janeiro^{14,39}. Our results indicate reassortment events in the NSP2, NSP3, NSP4, VP2, and VP3 genes in strains collected from Brazil that belong to the N2, T2, E2, C2, and M3 genotypes, associated with the co-circulation of DS-1 like and AU-1-like genomic constellations. Despite the evolutionary importance of the reassortment events in RVA genomes, the effects of these events on RVA vaccines still need to be addressed in further studies.

Our analysis of the genomic constellations by time and geographic region showed that before the vaccine was introduced, there was a greater number of genotype constellations in the Southeast and Northeast regions. The northern region had no constellations available in the pre-vaccine periods, which may indicate a lack of interest in monitoring RVA in this region before the vaccine introduction. In the post-vaccine periods, the Northeast continued with the second highest number of available constellations, but the Southeast has decreased the available sequences and the North has emerged as the region with the most sequences available, which demonstrates an inconsistency in RVA surveillance in the country that can make it difficult monitoring the epidemiology of the virus as well as the effectiveness of the vaccine. The Central-West region had the least number of constellations available throughout the analyzed period, including no constellation obtained in the post-vaccination period, which shows how little we know about the vaccine impacts on the complete RVA genome constellations circulating in this region. It is important to remember that the human Wa-

like strain of RVA has the same phylogenetic origin as porcine strains of RVA and the appearance of new strains is likely to happen during co-infection in pig cells¹³. Then, in environments where there is the human manipulation of these animals, as well as with cattle and canines, there is a potential for the emergence of new mixed strains between human and animal hosts. These new strains may not be able to infect human cells, which is more likely. However, it is necessary to maintain control measures, with good sanitary conditions, maintenance of vaccination to control the spread of the virus, and the maintenance of surveillance for viral circulation and analysis of the RVA genome to identify these changes and implement measures effective control systems¹³.

In this study, based on the analyzed genomic data, it was possible to conclude that strains of mixed genotype or animal origin have not been successful in maintaining themselves in the human population in Brazil over the years, with the Wa-like genotype being the most prevalent throughout the analyzed period. However, the evolutionary history between pathogen and host is dynamic and it is necessary to remain vigilant in this process to verify any event that could change the course of what is observed today.

Phylogenetic analysis and evolutionary history were inferred for all 11 genome segments of the RVA, which exhibited a pattern similar to temporal and regional. Sequences from different regions were identified in some internal groups. Thus, it was possible to verify that even in a continental country, such as Brazil, the temporal grouping overlaps the regional grouping in the analyzed phylogenies. Perhaps this is explained by the constant temporary migrations from one region to another in the country⁴⁰.

Although our findings are relevant, our study presents a limitation. Our data were obtained from a genomic database, in which some periods may be more represented than others since the genomic sequencing of rotavirus has become more accessible in recent years. Therefore, we have many more available sequences in recent years than in some

years in the past. Also, some regions may be over- or under-represented and the results here presented may not represent the real situation of each region. Even with these limitations, these results are relevant because it presents the distribution of circulating genotypes in Brazil over a period of 30 years, as well as the possibility that intergenogroup reassortment is occurring in Brazil and so it is important to establish an efficient surveillance system to follow the emergence of novel genotype constellations that might not be targeted by the vaccine.

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Figure 2: Sequence's disposition of the G and P genotypes in the period from 1986 to 2016.



Figure 3: Distribution of the combined G and P genotypes of Rotavirus A in Brazil from 1986

to 2016.



