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Received 5 October 2000/Returned for modification 2 January 2001/Accepted 5 March 2001


Group A rotaviruses (RV) are the leading cause of severe gastroenteritis in infants and young children worldwide and are associated with 600,000 to 800,000 deaths each year, mostly in developing countries (9).

Belonging to the Reoviridae family, RV are characterized by the presence of 11 segments of double-stranded RNA (dsRNA) enclosed in a triple-shelled protein capsid. Serotypes of RV are defined by the two outer proteins: the protease-sensitive protein, VP4, and the major glycoprotein, VP7. The two outer proteins induce neutralizing antibodies and designate the G (VP7) and P (VP4) viral serotypes (8). Using genotyping methods to determine P and G genotypes, the most common combinations found in humans worldwide are P[8]G1, P[4]G2, P[8]G3, and P[8]G4 (9). Each serotype of RV induces serotype-specific protective immunity, which has been used as the basis for the development of an RV vaccine including the four most important human strains (13). However, uncommon genotypes and/or serotypes infecting humans in developed and developing countries throughout the world have been detected (9, 19). In Brazil, uncommon RV serotypes and/or genotypes, such as P[8]G5, P[6]G2, G8, G10, mixed infections, and a significant number of untypeable strains, have been previously reported (14, 20). Serotype G9 RV was first described in the United States (4) and later in different localities, including Japan (15), Thailand (22), India (18), Italy (1), Bangladesh (21), and Malawi (6). Griffin et al. (12) detected and characterized G9 RV strains in 10 of 12 cities in the United States and suggested that this strain could be emerging as significant, reaching epidemic proportions. Other recent studies support the increasing importance of serotype G9 as a cause of severe diarrhea in children around the world (2, 5, 16, 17). Therefore, serotype G9 must be taken into consideration for prospective RV vaccine studies, since this serotype is not constituted in any of the preceding RV vaccines.

Considering the large diversity of RV infecting Brazilian children, we report strain genotyping results from 53 positive RV fecal specimens collected from hospitalized children less than 3 years old with acute gastroenteritis at Hospital Municipal Jesus (Rio de Janeiro, RJ, Brazil), a reference center for children care. The samples were collected between January 1997 and December 1998 during the first 8 h of hospitalization, and the children took medical care for at least 2 days. Approximately 10% (wt/vol) stool suspensions were prepared in Tris-HCl Ca2+ (0.01 M) (pH 7.2), and an enzyme immunoassay for RV and adenovirus antigen detection (EIARA) was carried out. Concomitantly, stool suspensions were used for dsRNA extraction by the glass powder method (3), followed by a polyacrylamide gel electrophoresis (PAGE). Electrophoresed dsRNA segments were visualized by silver staining. The EIARA, PAGE, and silver staining procedures were performed as previously described (14).

The viral dsRNA, extracted from RV-positive clarified stool supernatants by the glass powder method (3), was first reverse transcribed (RT) and amplified by PCR (first amplification step) with a pair of consensus primers corresponding to a conserved nucleotide sequence of the VP7 (7, 11) or VP4 (10) genes. The DNA fragment obtained of 904 bp (VP7) or 876 bp (VP4) was then used as a template in a second PCR, carried out by using 1 µl of the first amplicon and a pool of genotype-specific primers complementary to variable regions of the VP7 (11) or VP4 (10) gene. Temperature and time conditions for PCR amplifications were performed as originally described (10, 11). Distilled Milli-Q water was used as a negative control in all techniques, and recommended manipulations for PCR procedures were carried out as a precaution to avoid false-positive results.

The P[4]G2 genotype was the most common strain detected in 21% of these isolates, followed by P[8]G2 (17%), P[8]G1 (11%), P[8]G3 (6%), P[8]G4 (6%), P[8]G10 (6%), P[6]G4 (4%), and one mixed infection by P[8]+G3 (2%). In addition, we detected seven G9 strains: five P[8]G9 (9%), one P[6]G9 (2%), and one P[4]G9 (2%). As shown in Fig. 1, DNA bands corresponding to amplified segments of 110 bp for genotype G9 were visualized. The G9 strains were confirmed using Southern hybridization and chemiluminescent detection (data not shown), performed as previously described (14). The 5's end-labeled digoxigenin-oligonucleotide probe for the G9 gene strain US1205 (5' GC ATC AAC TCA AAT TGG AGA T; VP7 gene, nucleotides 313 to 333, plus-sense) was designed to
be homologous to internal regions of the consensus G9 first-round RT-PCR product produced by primers 9con1 and 9con2. In addition, we sequenced the G9 genotype-specific PCR products with the Big Dye Terminator (PE Biosystem Inc.) on an automated sequencer, ABI 377. The sequenced Brazilian strains were very similar to United States and Malawi G9 strains (data not shown), suggesting that they belong to type G9. The nucleotide and amino acid homology between Brazilian strain 1527 and other G serotypes showed very high homology to prototype G9 strain US1205 (Table 1), confirming that these strains belong to serotype G9.

Analysis of dsRNA by PAGE and silver staining revealed that all P[8] and P[4] strains had long electropherotypes, while the P[6] strain had a short profile. Samples that could not be characterized for G type (4 samples), P type (1 sample), or G and P types (3 samples) represented 14% of the strains. Despite the methodologies used in this study, some strains remain untypeable, raising the possibility that additional genotypes may be present in the population. It is important to characterize these strains, since they may emerge as being predominant in the future, as type G9 did between March and May of 1998 in the present work. The present study reports for the first time G9 strains infecting humans in Brazil; thus, these findings have significant implications, since this serotype has recently emerged in Australia (17), France (2), England (5), Ireland (16), and the United States (12), raising the possibility that RV type G9 may represent a fifth globally important serotype. The identification of novel P and G combinations of RV in Rio de Janeiro emphasizes the ability of this segmented virus to form reassortants (21) which may result in the establishment of new important strains.

**Nucleotide sequence accession number.** The VP7 gene sequence of Brazilian RV strain 1527 has been assigned the accession number AJ279082.

This work was supported by FIOCRUZ, COLAB/MS, and CNPq. We thank members of the WHO/PAHO Rotavirus Collaborating Center in the Viral Gastroenteritis Section of the Centers for Disease Control and Prevention (Dixie Griffin and Jon Gentsch) for assistance with this investigation and for critical review of the manuscript (Jon Gentsch).

**ADDENDUM IN PROOF**

Since the submission of this report, N. Santos et al. (J. Clin. Microbiol. 39:1157–1160, 2001) reported rotavirus serotype G9 in the state of Rio de Janeiro from 1997 to 1999, and the present study reinforces the previous description with the identification of G9 strains infecting hospitalized children (inpatient).

**REFERENCES**


