An Outbreak of Gastroenteritis Associated with Astrovirus Serotype 1 in a Day Care Center, in Rio de Janeiro, Brazil

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Between June 4th and June 20th 1996 rotavirus, adenovirus, and astrovirus (HAstrV) were investigated in fecal samples from 27 children under three years old with acute diarrhea, attending the Bertha Lutz day care center, in Rio de Janeiro. All fecal samples were analyzed by polyacrylamide gel electrophoresis (PAGE), reverse transcriptase polymerase chain reaction (RT-PCR), enzyme immunoassays (EIA), and electron microscopy (EM). Nine of them (33%) showed positive results for HAstrV by at least one of the employed methodologies. Eight were positive by RT-PCR and EIA, and six by EM. All positive samples were inoculated onto HT-29 (human colon adenocarcinoma) cultured cells for HAstrV isolation and seven were positive after three passages. The sequencing analysis of eight RT-PCR products (449 bp) from gene that codifies VP2 protein, showed a total nucleotide identity among them and 98% with HAstrV-1 (strain Oxford type 1). This is the first report of a gastroenteritis outbreak associated with HAstrV-1 in a day care center in Rio de Janeiro and it reinforces the importance of this virus in association with infantile acute gastroenteritis.

Key words: astrovirus - gastroenteritis - outbreak - Rio de Janeiro - Brazil

Astroviruses are plus-sense single-stranded RNA (ssRNA) viruses with 28 nm in diameter without a phospholipid envelope. Some viral particles show a star-like morphology by electron microscopy (EM) and recently were classified as belonging to the Astroviridae family (Monroe et al. 1994). Astroviruses have been isolated from human as well as from animal species. Eight human serotypes of astroviruses (HAstrV 1-8) have been described and the HAstrV-1 is the most prevalent strain (Kakizawa et al. 1997).

The medical importance of HAstrV was established when they were found to be the second most common cause of viral diarrhea in young children (Matsui & Greenberg 1996). HAstrV infection induces a mild watery diarrhea during two or three days, associated with other symptoms like vomiting, fever, anorexia and abdominal pain. Usually HAstrV infects young children, although gastroenteritis outbreaks associated with HAstrV involving older children, elderly people, and immunocompromised patients have been described (Matsui & Greenberg 1996).

In Brazil, there are few reports describing HAstrV infections (Nozawa et al. 1985, Leite et al. 1991, Stewien et al. 1991, Timenetsky et al. 1993, Tanaka et al. 1994) and all of them have used EM only for virus detection.

In this report, we describe for the first time an outbreak of gastroenteritis associated with HAstrV-1 in a day care center, in Rio de Janeiro, using different methodologies for virus characterization.

MATERIALS AND METHODS

Fecal specimens - Fecal specimens were obtained from 27 children under three years old with acute diarrhea. These specimens were collected at Bertha Lutz day care center, Oswaldo Cruz Foundation (Fiocruz), Rio de Janeiro, between June 4th-20th, 1996. Fecal suspensions (10%) were prepared in 10mM Tris, 1mM Ca++, pH 7.2, homogenized by vortexing, clarified at 3,000 x g for 10 min at 4°C and stored at -20°C until analysis.

Enzyme immunoassays - Initially, all fecal suspensions were screened for rotavirus and adenovirus using an enzyme immunoassay for group A rotavirus and adenovirus (EIARA) (Pereira et al. 1985). The test was carried out according to the manufacturer’s protocol (Bio-Manguinhos, Fiocruz, Ministry of Health, Brazil).

A commercial qualitative enzyme immunoassay (ELISA) (IDELIA™ Astrovirus) for detection of HAstrV from fecal suspension was performed following the recommendations of the manufacturer.
A non-commercial enzyme immunoassay (EIA), previously described by Noel et al. (1995), was performed to detect HAsTrV from supernatants of the infected HT-29 (human colon adenocarcinoma) cells (3rd passage).

**RNA extraction** - Five hundred µl of 10% fecal suspensions were submitted to RNA extraction using the glass powder methodology described by Boom et al. (1990).

**Polyacrilamide gel electrophoresis** - The polyacrilamide gel electrophoresis (PAGE) and silver staining methodologies were used to investigate the presence of segmented dsRNA viruses as previously described by Pereira et al. (1983).

**Reverse transcription followed by polymerase chain reaction** - The RT-PCR for HAsTrV was carried out as previously described by Noel et al. (1995). Distilled milli-Q water was used as negative control in ssRNA extraction and in RT-PCR procedure. HAsTrV-2 used as positive control was kindly provided by Dr SS Monroe, Viral Gastroenteritis Section, Enteric Respiratory Virus Branch, Centers for Disease Control and Prevention, Atlanta, GA, USA.

All manipulation recommended for PCR procedures were carried out as a precaution to avoid false positive results (Kwok & Higuchi 1989).

**Electron microscopy** - Fecal suspensions (10%) were treated with 2.45 M ammonium sulfate (v/w) and negatively stained with 2% phosphotungstic acid (pH 7.2) as described by Barth (1998). The grids were observed in a Zeiss EM-900 electron microscope, at magnification of 50,000X.

**Isolation of HAsTrV in HT-29 cells** - HAsTrV isolation was attempted from all positive samples by EM, RT-PCR and/or EIA by inoculation in HT-29 cells, which were kindly supplied by Dr Francisco Candal, Cell Cultural Development Section, Biological Products Branch, Centers for Disease Control and Prevention, Atlanta, GA, USA.

The cells were propagated in Dulbecco’s Medium supplemented with antibiotics (penicillin 100 U/ml and streptomycin 100 µg/ml), 2.5 µg/ml fungizone (DMS), and 10% fetal bovine serum (FBS) (Life Technologies, Inc. Gaithersburg, MD, USA). Two hundred-fifty µl of 10% fecal suspensions were treated with 20 µl antibiotics for 30 min at room temperature, homogenized by vortexing and clarified at 16.000X g for 5 min at 4°C. Two hundred ml of treated supernatants were mixed with equal volume of DMS containing 5 µg/ml of trypsin (Sigma-Aldrich Co. St. Louis, MO, USA) and inoculated onto confluent HT-29 monolayer cells, previously washed twice with 3 ml of DMS without FBS. The adsorption was carried out for 60 min at 37°C, then the supernatants were discarded and 5ml of DMS with 2% of FBS were added. The cells were incubated at 37°C for 18 h and a new fresh DMS containing 10 µg/ml of trypsin were added. After that, monolayer were incubated for five days at 37°C. Subsequently, cultures were harvested by freezing/thawing (-70/37°C) three times and submitted a two consecutive passages.

**Sequencing of RT-PCR astrovirus amplicons** - The RT-PCR products of eight positive samples were sequenced using the Prism Ready Reaction Dye Deoxy Terminator Cycle Sequencing Kit (Applied Biosystems, Inc, USA) following the protocol described by Noel et al. (1995) and an ABI 377 automatic sequencer (Applied Biosystems, Inc., USA). Sequence data from both strands of the RT-PCR products were aligned and compared using the GCG program of the Staden sequence analysis package (Dear & Staden 1991). The data were compared with the other HAsTrV sequences by using the University of Wisconsin, Genetics Computer package of programs (Devereux et al. 1984).

**Bacteriological and parasitological investigations** - The bacteriological and parasitological investigations were carried out following the protocols described by Edwards and Ewing (1996), Neves (1995), and Lima (1995), respectively.

**RESULTS**

**Laboratorial findings** - From 27 fecal samples analyzed by EM, RT-PCR and IDEIA™ Astrovirus, nine (33%) were positive by at least one methodology and five presented positive results for all applied methodologies (Table). No other enterophagogen was detected by the methods routinely used for investigation of bacteria, parasites, and rotavirus and adenovirus, characterizing an outbreak of acute gastroenteritis associated with HAsTrV.

In a group of nine positive samples, eight were detected by RT-PCR showing the expected 449bp fragment (Fig. 1). One sample was positive by IDEIA™ Astrovirus only (Table). As shown in Fig. 2, typical HAsTrV particles were visualized in 6 out of 27 samples (22%) analyzed by EM.

Nine positive samples were inoculated onto HT-29 cells and seven of them (78%) were positive by a non-commercial EIA, after the third passage.

The sequencing analysis of the RT-PCR products (nt 4526 to nt 4974, VP2 region) from eight isolates showed a nucleotide identity of 100% among them and 98% with HAsTrV-1 (strain Oxford type 1) (data not shown).

**Epidemiological and clinical findings** - HAsTrV were detected only in children under two years old (5 to 20 months), although the 27 fecal samples were obtained from children ranged between 5 to 32 months.
DISCUSSION

The development of specific assays with higher sensitivity have been established the role of HAsV as agent of gastroenteritis in young children, as well as in both elderly and immunocompromised patients (Mitchel et al. 1993, Jonassen et al. 1995, Mustafa et al. 2000). In order to characterize the etiological agent responsible for the outbreak of gastroenteritis at Bertha Lutz day care, different methodologies for virus, bacteria, and parasite detection, were employed. HAsV-1 was the only enterophatogen detected in this outbreak, with a positive rate of 33%. This is the first gastroenteritis outbreak associated with HAsV-1 in a day care center reported in Rio de Janeiro, Brazil. Other outbreaks in the same day care center, occurring during 1996 and 1997, were investigated, but HAsV were not detected. During the same period, HAsV were detected in sporadic infections among hospitalized children, demonstrating the circulation of those viruses in Rio de Janeiro (Silva et al. unpublished observations).

RT-PCR and ELISA (IDEIA™ Astrovirus) were equivalents to detected HAsV. However, the positivity of one sample just by IDEIA™ Astrovirus, suggests: (i) the presence of defective particles in a lower concentration than EM sensitivity; (ii) a false positive result. Despite the small number of samples, all methods used in that investigation showed to be satisfactory to detect HAsV. As suggested by Glass et al. (1996), the RT-PCR and EIA should be the methods of choice for HAsV detection, since RT-PCR have been increased the viral detection in stool samples.

Although outbreaks associated with several HAsV serotypes have been described (Oishi et al. 1994, Belliot et al. 1997), HAsV-1 is the most prevalent serotype circulating worldwide (Noel et al. 1995, Shastri et al. 1998, Medina et al. 2000).

In Brazil, reports have demonstrated the occurrence of HAsV at low frequencies (2-5%). The use of the EM as the only methodology for virus detection showed to be satisfactory to detect HAsV. RT-PCR and ELISA (IDEIA™ Astrovirus) were equivalents to detected HAsV. However, the positivity of one sample just by IDEIA™ Astrovirus, suggests: (i) the presence of defective particles in a lower concentration than EM sensitivity; (ii) a false positive result. Despite the small number of samples, all methods used in that investigation showed to be satisfactory to detect HAsV. As suggested by Glass et al. (1996), the RT-PCR and EIA should be the methods of choice for HAsV detection, since RT-PCR have been increased the viral detection in stool samples.

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Fig. 1: detection of genomic RNA segment of astrovirus (HAsV) by RT-PCR from fecal samples. Line 1: MW 123bp Ladder (Life Technologies, Inc); lines 2-5, 7, 10: negative stool samples; lines 6, 8, 9, 11, 12: positive stool samples; lane 13: negative control; lane 14: HAsV-2 positive control.

Fig. 2: electron micrograph of astrovirus (HAsV) particles from sample LVC426. Arrows are indicating typical morphology of HAsV. Bar = 100 nm.

The major clinical symptoms observed during the outbreak were: vomiting, fever (37.5-38°C), abdominal pain, inappetence, cough, coryza and mucus on feces. The symptoms associated with diarrhea persisted for three to six days.
To supply IDEIA™ Astrovirus kit. To the staff of Bertha Lutz day care center all positive samples were detected from children between 5 and 20 months. Our data corroborate the findings reported by Mendes (1997) during a sera survey conducted in Belém, Brazil, when it was demonstrated that HAdV seropositivity rates increased from 13.9%, at 6 months of age to 64.9% in children up to 2 years old, suggesting that HAdV infection is more frequent than the previously reported (Nozawa et al. 1985, Leite et al. 1991, Stewien et al. unpublished observations).

At Bertha Lutz day care center all positive samples were detected from children between 5 and 20 months. Our data corroborate the findings reported by Mendes (1997) during a sera survey conducted in Belém, Brazil, when it was demonstrated that HAdV seropositivity rates increased from 13.9%, at 6 months of age to 64.9% in children up to 2 years old, suggesting that HAdV infection is more frequent than the previously reported (Nozawa et al. 1985, Leite et al. 1991, Stewien et al. unpublished observations).

The molecular epidemiology of HAdV remains to be elucidated in Brazil and further investigations should provide the answer as to the role of individual serotypes in outbreaks in day care centers, elderly houses, and inpatients clinics. This would be achieved by performing molecular epidemiology of both common and uncommon serotypes.

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