The in Vitro Cytopathology of a Porcine and the Simian (SA-11) Strains of Rotavirus

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Rotaviruses have been implicated as the major causal agents of acute diarrhoea in mammals and fowls. Experimental rotavirus infection have been associated to a series of sub-cellular pathologic alterations leading to cell lysis which may represent key functions in the pathogenesis of the diarrhoeic disease. The current work describes the cytopathic changes in cultured MA-104 cells infected by a simian (SA-11) and a porcine (1154) rotavirus strains. Trypan blue exclusion staining showed increased cell permeability after infection by both strains, as demonstrated by cell viability. This effect was confirmed by the leakage of infected cells evaluated by chromium release. Nuclear fragmentation was observed by acridine orange and Wright staining but specific DNA cleavage was not detected. Ultrastructural changes, such as chromatin condensation, cytoplasm vacuolisation, and loss of intercellular contact were shown in infected cells for both strains. In situ terminal deoxynucleotidyl transferase (Tunel) assay did not show positive result. In conclusion, we demonstrated that both strains of rotavirus induced necrosis as the major degenerative effect.

Key words: rotavirus - cytopathology - necrosis - apoptosis - cell culture

Rotaviruses are the main cause of acute diarrhoea in newborn and young mammals and some avian species (Kapikian & Chanock 1996). The genus rotavirus, within the family Reoviridae, consists of viruses that present eleven segments of double-stranded RNA and 70 nm in diameter (Shaw et al. 1995). The virions are icosahedral and non-enveloped with a concentric triple-layered double-shelled capsid. The outer layer is made of VP4 and VP7 proteins, the intermediate layer constituted of VP6 and the inner layer represented by VP2. VP1-VP3, and the genome form the virion core (Patton 1995).

The viruses infect villous enterocytes of the small intestine (Blacklow & Greenberg 1991) causing cell lysis and sloughing of the epithelium cells. The mechanism of rotavirus-induced diarrhoea is multifactorial and among the proposed causes is the secretory effect attributed to nonstructural virus protein (NSP4) and/or to the stimulation of enteric nervous system (Lundgren & Svensson 2001). Although many pathways of rotavirus morphogenesis and several cytopathic effect features are established, virus-induced cell death remains uncertain. It was shown that experimental rotavirus infection promoted a progressive change in membrane permeability to cations (Michelangel et al. 1995). Additionally, Perez et al. (1998) also showed cell rounding and loss of membrane integrity demonstrated by the increase of membrane permeability to macromolecules and release of cellular contents. All these features point at necrosis as the outcome of rotavirus infection. However, changes such as peripheral condensation of the chromatin and fragmentation of the nucleus were found as suggestive of apoptosis induced by the virus in HT29 cell culture (Superti et al. 1996).

Necrosis is a non-specific form of cell death following injury from an external agent. It is characterized by cell swelling, loss of plasma membrane integrity, and leakage of the cell contents after rupture of the cell (Majno & Joris 1995).

Apoptosis, on the other hand, is a process of physiological cellular death in that cells are submitted to death in response to a variety of stimuli (Teodoro & Branton 1997). Morphologically it is characterized by the reduction of cellular volume, loss of intercellular contact, condensation of the chromatin at the nuclear periphery, vacuolisation of the cytoplasm, cell membrane “blebbing”, and formation of apoptotic bodies. In some cases activation of endogenous endonucleases occurs and results in cellular DNA cleavage in oligonucleosomal fragments of multiples of 180 to 200 pb, differently to the random DNA fragmentation observed in necrosis (Wyllie et al. 1984).

It has been shown that apoptosis induced by several virus infections contribute directly to the cytopathology and pathogenesis of those infections (Alonso et al. 1998, Oura et al. 1998). Evidences also show that many viruses encode specific proteins that arrest apoptosis (Tolskaya et al. 1995, Teodoro & Branton 1997, Derfuss et al. 1998).

The current work was proposed to evaluate the outcome of a porcine and simian rotavirus infection underlying their in vitro cytopathology.
MATERIALS AND METHODS

Virus and cell culture - Simian (SA-11) and porcine (strain 1154) (Ramos et al. 2000) strains of rotavirus were grown in MA-104 cell cultures (monkey kidney cells) maintained in Dulbecco’s Modified Eagle Medium free of foetal bovine serum (FBS), with antibiotics, and 10 µg/ml of crystalline trypsin used throughout, unless otherwise indicated. Virus inocula were previously treated with 30 µg/ml of trypsin for 50 min at 37°C, and for the experiments indicated. Virus inocula were previously treated with 30 µg/ml of crystalline trypsin used throughout, unless otherwise indicated. Bovine serum (FBS), with antibiotics, and 10 µg/ml of crystalline trypsin were maintained in Dulbecco’s Modified Eagle Medium free of foetal bovine serum (FBS), with antibiotics, and 10 µg/ml of crystalline trypsin used throughout, unless otherwise indicated.

Trypan blue exclusion staining (TB) - Coverslips with virus-infected cell cultures, as well as uninfected cultures, were harvested at 0, 2, 4, 6, 8, 10, and 12 h p.i. and stained with 1% TB. Results are presented as the percentage of viable cell, calculated by X = 100 – (A/B x 100), where A is the number of infected (death) cells and B is the number of live (control) cells. The number of cells (A and B) is the average of three independent counts.

Chromium ($^{51}$Cr) release assay (CR) - Briefly, 48 h confluent cell cultures were dispersed with 0.02% EDTA, centrifuged, and the pellet was resuspended in 5 ml of culture medium containing 100 µCi of Na$_2$[Cu]$^{51}$CrO$_4$ (3.6 µg Cr/ml) (Amersham Pharm. Byotech, São Paulo, Brazil) and incubated for 45 min at 37°C. Cell suspension was washed twice with warm medium and the resultant pellet resuspended in 5 ml medium, and split into two equal volumes. The cells were infected with virus, previously treated with 30 µg/ml of trypsin. After adsorption and cell pelleting, fluid medium containing viral inocula was discarded, cells resuspended in fresh culture medium containing 5% FBS and 10 µg/ml trypsin, and 1 ml-aliquot of cell suspension placed in culture tubes. Mock-infected cultures were prepared similarly. At time intervals of 0, 2, 4, 6, 8, 10, and 12 h p.i., cultures were centrifuged and the supernatant submitted to radioactivity counts in gamma counter (Polos & Gallerah 1981). The percentage of chromium release was calculated by X = C/D x 100, where C is cpm (count per minute) in the supernatant of infected cells and D is cpm in the supernatant of non-infected cells. Counts of C and D are the average of three independent counts.

Acridine orange staining (AO) - Briefly, cell cultures grown in coverslips and infected with virus were harvested at 24 h and 48 h p.i., and fixed with Carnoy’s fixative for 5 min followed by hydration with decreasing concentrations of ethanol. Cells were washed in McIlvaine’s buffer stained with 4.3 µM AO for 5 min, and observed under UV light (Guimarães & Nozawa 1990).

Wright staining (WS) - Infected cultures were harvested at 24 h and 48 h p.i., washed with PBS, fixed with methanol for 1 h, and overlaid with 0.3% Wright stain for 4 min. Cultures were washed with phosphate buffer, pH 6.5, mounted in slides, and observed.

Transmission electron microscopy - Infected cells were scrapped off from the tissue culture bottles at 12 h and 24 h p.i., washed by centrifugation with PBS, and fixed with 2.5% glutaraldehyde in 0.1 M sodium cacodylate/0.2 M sucrose buffer, pH 7.2. One percent osmium tetroxide post-fixation was followed by 1% uranyl acetate treatment. Dehydration was done with acetone followed by embedding in epon. Ultra-thin sections were stained with lead citrate and uranyl acetate (Uehara et al. 1992).

DNA fragmentation - Infected cells harvested at time intervals of 8, 24, 48, and 96 h p.i. were treated with 500 µl of lysis buffer (10 mM Tris-HCl; 10 mM EDTA; 0.5% SDS, pH 8.0) at 56°C for 24 h and extracted with phenol/chloroform. Extracted DNA was precipitated overnight with 5 M NaCl and isopropanol at –20°C, resuspended in 30 µl Tris-EDTA buffer (10 mM Tris-HCl; 1 mM EDTA, pH 8.0) and submitted to electrophoresis either in 1.5% agarose (AGE) or 7% polyacrylamide (PAGE) gels (McDonell et al. 1977, Herring et al. 1982).

In situ terminal deoxytransferase-mediated dUTP nick end-labelling (Tunel) - Twenty-four and 48 h p.i. cultures were treated with 2 µg/ml of proteinase K followed by 1 h incubation at 37°C with hybridization solution [100 mM sodium cacodilate, pH 6.5; 2 mM cobalt chloride; 50 µg/ml bovine serum albumin; 30 U terminal deoxynucleotidyl transferase (Tdt); 20 µM 5-BrdU; 4 µM dGTP, dCTP, and dATP; 30 mM Tris-HCl]. Cells were washed with 50% formamide followed by three washings with citrate-saline (0.3 M NaCl; 0.03 M sodium citrate) at 45°C and twice with PN buffer (0.05% Nonidet P-40 in PBS, pH 8.0) at 30°C. This was followed by incubation with mouse anti-BrdU antibodies (Sigma Chem.Co., MO, US) diluted to 1:200 in PBS for 30 min at 37°C, and incubated with FITC conjugate secondary goat antibodies (Sigma) diluted to 1:100. Preparations were observed under UV (Ansari et al. 1993). Poliovirus-infected HEp-2 cell cultures were used as Tunel positive control.

RESULTS

Rotavirus-infected cell viability assayed by TB exclusion staining is shown in Fig. 1, at indicated period of time. The percentage of cell viability varied from 10.1% at time zero to 21.5% at 12 h p.i. for porcine strain. For simian strain it varied from 10% at time zero to 18% at 12 h p.i.
Infected cells submitted to CR test, at the same time intervals, demonstrated that for simian strain percentages of chromium release from 2% at time zero to 24% at 12 h p.i. were observed, and for porcine strain 4% at time zero to 27.5% at 12 h p.i. (Fig. 2). Concerning nuclear fragmentation monitored by AO and Wright staining no difference was detected among infected and control cells 24 h p.i., for both strains. However, at 48 h p.i. fragmentation was observed in approximately 2 and 4% of infected cells, respectively for simian and porcine strains (Fig. 3).

Thin sections of porcine virus-infected cell cultures, harvested at 12 h p.i., revealed a decrease in the ratio of cellular and nuclear volumes. Condensation and marginalization of the chromatin, increase of the nucleoli, and vacuolisation of the cytoplasm were also observed. Loss of intercellular contacts was demonstrated (Fig. 4A). Enveloped and non-enveloped virus particles were also detected within dilated cistern (Fig. 4A, B). Similar results were also found in simian virus-infected cells (not shown).

DNA of infected cultures, collected at 8, 24, 48, and 96 h p.i., extracted and submitted to AGE and PAGE showed no cleavage as to form fragments of 180-200 pb. However, it was demonstrated that from 24 h p.i. onward virus yields increased significantly as shown by the optical density of viral RNA segments in AGE (not shown). Tunel technique revealed negative results for both strains (not shown).

**DISCUSSION**

Rotavirus infection in vitro and in vivo is associated to several sub-cellular pathological changes that ultimately culminate with cell lysis. The mechanism that rotaviruses use to interact with host cell and results in cytopathetic effect is not clearly defined. It is thought that these

![Fig. 2: percentage of chromium (51Cr) release in rotavirus-infected MA-104 cell cultures. Simian (●) and porcine (□) strain-infected cell cultures at a multiplicity of infection of approximately 1 at the indicated time intervals.](image)

![Fig. 3: nuclear fragmentation in simian (S) and porcine (P) rotavirus-infected MA-104 cell cultures. Cell cultures infected at a multiplicity of infection of approximately 1, 48 h p.i. stained with acridine orange (AO) (S1 and P1, 200X) and Wright stain (W) (S2 and P2, 400X) showing nuclear fragmentation (arrow). Control non-infected cell cultures stained with AO (C1, 200X) and W (C2, 400X).](image)
changes may represent a key role in the pathogenesis of diarrhoeic disease caused by rotaviruses.

In the current work, the analysis of cell viability after infection with simian and porcine strains of rotavirus demonstrated that in the absence of cellular morphological changes, respectively, 18 to 22% of the cells presented altered membrane permeability 12 h p.i., as shown by TB uptake. CR assayed at the same time intervals demonstrated that over 23 and 27% of the radioactivity were released, respectively, for simian and porcine strains by 12 h p.i. Death of rotavirus-infected cells has been established as being caused by lytic process (Estes 1996). However, it was suggested that some cytopathic events observed in rotavirus-infected cells, including chromatin clumping could be attributed to apoptosis induced by the virus (Superti et al. 1996). In our experiments necrosis takes place as shown by membrane permeability derangement. Also infected cells presented nuclear fragmentation later in infection, observed after WS and AO staining. We suggested that death of rotavirus-infected cells occur mainly due to necrosis, at least for the virus strains we used. However, a small percentage of infected cells showed evidence of apoptosis as demonstrated by the nuclear fragmentation, although Superti et al. (1996) suggested that apoptosis was the prevalent process in the cytopathology of rotavirus. Nuclear fragmentation was demonstrated in our experiments but we could not detect DNA ladder internucleosomal fragments, as shown by Pérez et al. (1998) with OSU strain of porcine rotavirus, but this may not be mandatory in rotavirus-induced apoptosis (Superti et al. 1996). Cohen et al. (1992) and Oberhammer et al. (1993) showed apoptotic ultrastructural changes in the absence of internucleosomal cleavage. As far as ultrastructural changes are concerned we detected some changes that could be attributed to apoptosis in infected cells in comparison to controls. High molecular weight DNA fragments (300-700 bp and 50 Kbp) were described to occur previously to internucleosomal cleavage (Brown et al. 1993, Zhivotovsky et al. 1994) suggested as an irreversible signal of apoptosis (Orrenius 1995). Yet again our results did not demonstrate high molecular weight DNA fragments either.

Although histochemical staining have shown nuclear fragmentation we could not demonstrate Tunel positive reaction. It is possible that either AGE and PAGE are not enough sensitive to detect low scale DNA fragmentation or alternatively there could be different endonucleases involved, not allowing Tunel reaction (Desjardins & MacManus 1995). These findings may suggest that overall lytic alterations observed in rotavirus-infected cells should be attributed to cytotoxic origin. This may be explained by the accumulation of viral proteins as suggested by Michelangeli et al. (1995) and loss of cytoplasmic membrane integrity even early in infection, as we showed by TB uptake and CR. In corroborations, NSP4 increases cytoplasmic membrane permeability being, therefore, implicated in rotavirus cytopathic effect due to changes of Ca2+ permeability in the endoplasmic reticulum (ER) (Newton et al. 1997). This type of trans-ER resident glycoprotein has been suggested to act, among other functions, as diarrhoea inducing enterotoxin in mice and in the reorganisation of cellular Ca2+ (Ball et al. 1996, Newton et al. 1997, Mirazimi et al. 1998). In fact, Exton (1994) demonstrated that the increase of intracellular Ca2+ in response to NSP4 could alter cell metabolism in activating enzymes and ultimately leading to necrosis.

In conclusion, we suggested that both species of rotavirus studied interact with cultured cells overall as a lytic process and necrosis seems to be the basis of their cytopathology.

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