ISOLATION OF ROTAVIRUSES FROM CALVES WITH ACUTE ENTERITIS AND THEIR CULTIVATION IN VITRO

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Abstract


Five bovine rotavirus strains were isolated in monolayers of MA-104 cell culture from faeces of 3-to 11-day-old calves suffering from gastroenteritis. Cytopathic effect, accompanied by the release of cells from the glass surface of cultivation flasks was observed from the 5th to the 6th passage. After the stabilization of the cytopathic effect, the highest virus concentration was observed in cell cultures that had been frozen 18–36 hrs after inoculation. Electron microscopy revealed particles with typical rotavirus morphology. Antigenic relationship of the isolates with the reference strain Lincoln was confirmed by ELISA. The isolates were classified as members of the serological group A by ELISA and immunofluorescence assay. The identity of the rotavirus isolates was also confirmed by electrophoresis in agarose gel. The assay confirmed that their genome consists of segmented viral RNA, which produced in the electric field migration patterns, electrophorogrammes, typical of rotaviruses. In 4 isolates the migration speed of the segments in the electric field was identical with that of the reference strain. A different electropherotype was identified in the remaining isolate.

Rotaviruses are at present considered as important causal agents of diarrheic infections in many domestic and wild ruminants, swine, monkey, horse, cat, dog, mouse, rabbit, poultry, fish, and also man (Wyatt et al. 1974; Flewett and Woode 1978; Fulton et al. 1981; Hoshino et al. 1981; Theil et al. 1985; Thouless et al. 1986; Korych 1987; Marshall 1990). They are found especially in newborn farm animals (neonatal diarrhoea) and children. Rotaviruses have been detected in various climatic zones, mostly in connection with an outbreak of gastrointestinal infections with various intensity of clinical manifestation, ranging from asymptomatic infection to severe, even lethal cases (Dea et al. 1985; Korych 1987; Ruuska and Vesikari 1990).

Causal relation of rotaviruses to gastroenteritis in newborn calves was first described by Mebus et al. (1969), who induced gastroenteritis in calves by peroral administration of bacteria-free filtrates of faeces from scouring animals. Reovirus-like agents were detected in the gut contents of infected calves (Fernelius et al. 1972). After a series of unsuccessful attempts, the virus was isolated and adapted to the growth in cultures of bovine embryonic kidney cells (Fernelius et al. 1972). The isolate was identified as bovine rotavirus, different from reovirus (Flewett and Woode 1978). The new information concerning the aetiology of scour was confirmed by several authors in the U.S.A. (Eckern et al. 1981), United Kingdom (Woode et al. 1974; McNulty et al. 1977) and other countries including Czechoslovakia (Scherrer et al. 1976; Marsolais et al. 1978; Wyn-Jones et al. 1978; Köves 1979; Šmid et al. 1980; Castrucci et al. 1983; Barboi et al. 1987). Most studies, however, focused on the methods of rotavirus demonstration in faeces of calves suffering from gastroenteritis (Rodger and Holmes 1979; Šmid et al. 1980; Eckern et al. 1981). Early attempts to isolate rotaviruses in cell or organ cultures derived from various animal species were unsuccessful. Studies with established cell lines of various origins were carried out with the aim of selecting suitable ones for the growth of rotavirus strains...
and for the diagnosis of rotavirus infection (Bryden et al. 1977; Butchaiah et al. 1984). Good results were obtained by using the cell line MA-104, derived from Maccacus rhesus foetal kidney (Castrucci et al. 1983; Butchaiah et al. 1984; Thouless et al. 1986). The virulence increased after repeated passages in the gastrointestinal tract of sensitive animals and the following transfer of the obtained virus suspension to a suitable cell culture (Birch et al. 1983; Korych 1987).

More effective adsorption of virus particles on cell monolayers was obtained by a short low-speed centrifugation performed immediately after inoculation (Korych 1987). Positive effect on the culture was also obtained using the roller culture technique (McNulty et al. 1977; Thouless et al. 1986). But most important finding was treatment of rotavirus suspension with proteases (trypsin is the most frequently used treatment) results in a marked increase of virus infectivity (Almeida et al. 1978; Babiuk and Mohammed 1978).

These findings facilitated the isolation of new rotavirus strains from clinical samples and their cultivation in vitro, which is necessary for further studies of rotaviral antigenic structure, serotype identification as well as for the development and improvement of diagnostic methods and immunoprophylactic measures. For this reason, our experiments were concentrated on isolation of bovine strains occurring in cattle herds in the Czech Republic, the investigation of their growth properties and optimal culture conditions.

Materials and Methods

Cell cultures and media

The established line of foetal monkey kidney cells MA-104 was grown in Eagle's minimum essential medium (MEM) containing 10% foetal calf serum.

Reference virus

Reference strain Lincoln, obtained from the Collection of zoopathogenic microorganisms in the Veterinary Research Institute, Brno, was used in the experiments.

Faecal samples

Samples of faeces were collected during 1989—1990 in rearing premises where frequent scours were recorded in up to 3-week-old calves. The samples were collected into plastic bottles, frozen on dry ice, transported to the laboratory and stored at −22°C until processing. Eleven faecal samples, in which rotaviruses were demonstrated to be present by electron microscopy and ELISA, were used for the isolation.

Inoculum preparation

Inoculum for cell cultures was prepared from approximately 20% suspension of faeces in PBS. The suspensions were centrifugated at 7,000 g for 20 minutes and supernatants were filtered through a 0.22 μm Millipore filter. Infectivity of rotaviruses in virus suspensions was enhanced by the addition of trypsin to a final concentration of 10 μg per 1 ml. The suspensions were incubated at 37°C for 30 minutes and then used as the inoculum.

Isolation and adaptation of strains to cell cultures

Complete monolayers of MA-104 cells grown in Mueller's flasks were washed 3 times with serum-free Eagle's medium and inoculated with 0.5 ml of trypsin-treated virus suspension per flask. After 1 hour incubation at 37°C with an occasional pouring of the inoculum over the monolayer surface, 5 ml of calf serum-free Eagle's maintenance medium, supplemented with trypsin to the final concentration 1 μg per 1 ml of medium, were pipetted into each flask. The cultures were incubated at 37°C until a distinct cytopathic effect became apparent. If no CPE was evident, the incubation proceeded for 7 days at least. Then the cultures were frozen and stored at −22°C.

Enzymoimmunoanalysis (ELISA)

The direct sandwich method was used. Aliquots of series of two-fold dilutions ranged from 1 : 2 to 1 : 256. After washing, the antitrovirus antibodies conjugated with peroxidase were added. The reaction was visualized by the addition of a substrate consisting of hydrogen peroxide and 5-aminosalicylic acid and measured spectrophotometrically at 492 nm. The results were compared with those of positive and negative controls, which were included into each series of examinations.

Electron microscopy

Electron microscopic identification of rotavirus was done as described by Šmid et al. (1980). Approx. 20% suspensions of samples in PBS were prepared and centrifugated at 7,000 g for 20 min. A drop of each sample was used for negative staining with 2% phospho-tungstic acid and examined by electron microscope Tesla BS 513.
Electrophoresis of viral RNA in agarose gel
Horizontal electrophoresis in agarose gel, as described by Chudzio et al. (1989) and Pšikal et al. (1991) was used for the demonstration of virus RNA. The gel was prepared as a 5-mm-thick layer of 1.5% agarose boiled in 0.09M Tris-borate buffer, pH 8.2. The liquid gel was poured into a 8.5 x 8 cm mould, into which a comb had been placed to form starts. After solidification, the mould was put into an electrophoretic vessel and poured over with Tris-borate buffer. The processed samples were put onto starts by a micropipette and after connection to a direct current generator, the electrophoresis was left to run at 100 V for 2 hrs. After switching off, the gel was stained in ethidium bromide solution (1 µg/ml) for 10 min. The stained segments of RNA were viewed under ultraviolet light at 312 nm.

Immunofluorescence test
Monolayers of MA-104 cells, grown on slides 0.8 x 3.0 cm were washed to remove the growth medium, placed into polystyrene Petri dishes (diameter 6.0 cm), inoculated with 0.5 ml of examined virus and poured over with Eagle's maintenance medium. After 24 hrs incubation at 37°C in an atmosphere containing 5% CO₂, the slides with monolayers were washed in PBS, fixed with chilled acetone (−22°C) and dried at laboratory temperature. The monolayers were then stained with the direct conjugate to bovine rotavirus for 45 min. After washing in PBS, slides were mounted into glycerol solution in PBS and viewed under a fluorescence microscope.

Evaluation of virus growth in cell cultures
Cell cultures MA-104 grown in Mueller's flasks were inoculated with the same quantity of rotavirus 24 hrs after the seeding. ELISA was used for the detection of virus concentrations in the samples obtained by freezing of the cell cultures in different periods following inoculation.

Results

Virus isolation in cell cultures and characterization of cytopathic lesions
No cytopathic changes were observed in any of the isolates in the 1st passage after 7 days of incubation. The first evident cytopathic changes developed in the 2nd or 3rd passage. They were noticed in cell cultures infected with five faecal samples (BR 14, BR 197, BR 207, BR 224 and BR 256) and were characterized by the formation of rounded cells on the surface of almost intact monolayers. In the 4th passage, the cytopathic lesions were already observed on 3rd day p. i. The number of degenerated cells, located on the surface of monolayer, or released into the medium, grew, and the first signs of impaired compactness of the cell layer became evident. CPE stabilized during the subsequent passages, corresponding to that of the reference strain Lincoln as for its dynamics and morphology. The first signs of cell degeneration appeared 18 to 24 hours after inoculation. The number of degenerated cells grew rapidly, and the proceeding destruction of the cell layer resulted in the release of cells from the glass surface (Plate XVIII., Fig. 3). CPE culminated 36 hours p. i. when 80 to 90 percent of the monolayer was destroyed. No CPE was observed during 3 passages in cell cultures inoculated with any of the remaining six samples.

Immunofluorescent investigations of the isolated strains
Although fluorescence was already recorded in the first two passages of 9 rotavirus suspensions, it was limited to single cells disseminated irregularly in the monolayer. Groups of cells, showing mostly granular, perinuclear fluorescence were observed in the 3rd passage in those cell cultures, from which cytopathogenic rotavirus strains were isolated. The first positive cells appeared 18 to 24 hours p. i. Fluorescence in approx. 5% of the cells was already detectable 6 hours p. i. from the 4th passage on. The number of cells grew rapidly and 20 to 25% of cells were positive 12 to 18 hours p. i. The fluorescence was located in the cyto-
plasm and became diffuse. 40 to 50% of cells were positive 18 to 36 hours p. i. (Fig. 4). A similar type and extent of fluorescence was observed in cell cultures inoculated with the reference strain Lincoln.

**Virus multiplication in cell cultures**

As confirmed by ELISA, the rise of rotavirus concentration in inoculated cell cultures was proportional to the development of CPE and fluorescence. The first new virus was already detectable 6 hours p. i. and peak concentrations were reached 18 to 36 hours p. i.

**Electron and immunoelectron microscopy**

Virus particles, corresponding in morphology to rotaviruses, were seen in negatively stained preparations of the 5th passage of the cytopathogenic isolates (Fig. 5). The particles were aggregated into immune complexes by antibodies to the strain Lincoln. No other virus particles were observed in the preparations.

**Agarose gel electrophoresis of viral RNA**

Segments of RNA, belonging to four classes relative to their migration properties, were obtained by agarose gel electrophoresis of nucleic acid of the isolated rotavirus strains. The migration patterns were characteristic for rotaviruses. They can be used for direct species identification, and analysis of migration properties of certain segments enables to distinguish various electrophoretic types of the virus. As can be seen in Fig. 2, four of the isolates which yielded electrophorograms were identical with that of the strain Lincoln. A markedly different position of the segment 11, located between the segment 6 and the segment triad 7, 8, 9, was evident in the electrophorogramme of the strain BR 207. A similar arrangement was found in the rotavirus strain present in the sample BR 125 which failed to adapt to cell cultures. The segment 11 was located between the segment 6 and segment triad 7, 8, 9, but in close vicinity of the segment 7 (Fig. 1). The two electropherotypes, different from the strain Lincoln, could be distinguished despite the limited resolution capacity of agarose gel, which did not allow a complete separation of all 11 segments. The segments 2 and 3 migrated together producing one broader and brighter band under UV light. Also the segments 7, 8 and 9 migrated together and thus the number of bands, visible under UV light, was reduced from 11 to 8 (Psikal et al. 1991). No genome reassortment was observed in any of the isolates during the passages in cell cultures, as demonstrated by the comparison of electrophorogrammes of the final passages with genomic profiles of rotavirus strains present in the faecal samples.

**Discussion**

Isolation of bovine rotaviruses from faeces of calves affected with acute gastro-enteritis, and their propagation in cell cultures are described. Faecal samples, in which high concentrations of rotaviruses had been confirmed by several methods, were used in isolation attempts. The samples were collected in several farms, and the rotaviruses present in them belonged to three different electropherotypes.

The cell line MA-104 which is used for the propagation of rotaviruses most frequently, was selected for the isolation attempts. Only 5 of the 11 rotavirus suspensions were adapted to cell cultures and could be passaged in vitro serially. This poor isolation success was apparently due to the fact that not all effective culture methods (roller cultures, e. g.) were applied.
The typical cytopathic effect became stabilized in higher passages only and therefore the propagation of rotaviruses was checked by the fluorescent antibody technique. This technique is reliable enough and can be used for that purpose in virus strains propagating without characteristic CPE (Birch et al. 1983). Some authors reported the appearance of fluorescence 2 to 8 hours after inoculation, i.e. at the time when no CPE is detectable in infected monolayers. In our experiments, specific fluorescence became visible 6 hours after inoculation. No specific fluorescence was detectable in the 2nd and 3rd subsequent passages of the rotavirus suspensions which failed to adapt to the growth in vitro.

Electron microscopic examinations of five cytopathogenic isolates demonstrated the presence of virus particles corresponding morphologically to either the one- or two layer capsid type of rotavirus. Immunoelectron microscopy confirmed the results of fluorescent antibody technique and ELISA, classifying all isolated strains as members of the serological group A as defined by Estes and Cohen (1989).

Genomic analysis by agarose gel electrophoresis of RNA allowed a further differentiation of the isolated strains. While the migration patterns of four of them were identical with that of the strain Lincoln, the fifth - BR 207 - belonged to another electropherotype, different from them in the position of segment 11. Passage of isolated strains was not associated with any changes of their migration profiles.

Our results have confirmed the finding of Pšikal et al. (1991), that rotaviruses circulating in the local cattle population cannot be regarded as an entirely homogeneous set of strains. This finding should be considered in the ongoing updating of diagnostic methods and development of biological products for prophylactic application.

Izolace rotavirů z telat s akutní enteritidou a jejich kultivace in vitro

Выделение ротавирусов из телят с острым гастроэнтеритом и их культивирование ин витро

Из пометов 3 - 11 денных телят заболеваемых гастроэнтеритом выделили в монослоях клеточной культуры МА-104 пять штаммов бычьего ротавируса. Их рост происходил от 6-го пассажа развития характерных изменений, соединенных с отпадением клеток от поверхности стекла. После стабилизации цитопатических эффектов, т. е. от 5-6-го пассажа обнаружили самую большую концентрацию вируса в клеточных культурах замороженных 18-36 часов после инфекции. Помощью электронного микроскопа доказали что морфология выделенных штаммов сходится формой и размером с ротавирусом. Иммуноферментный анализ (ЕЛИСА) подтвердил антигенный родство выделений с эталонным штаммом Линкольн. Результаты иммуноферментного анализа и иммуноперфосцентного анализа предполагают отношение выделенных штаммов в серогруппе А. Идентичность выделений с ротавирусами подтверждена электрофоэреом в агарозовом геле. Этот анализ подтвердил, что их геном состоит из сегментов вирусной РНК, которые образовали в электрическом поле миграционные формулы - электрофоретограммы - типичные для ротавирусов. Миграционная активность этих сегментов в электрическом поле была у четырех выделенных штаммов идентична с миграционной активностью эталонного штамма. Один из выделенных штаммов отличался электрофоретикопо.

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References


Fig. 1: Electrophoresis of rotavirus RNA extracted from diarrhoeic calves in 1.5% agarose gel stained with ethidium-bromide. Lanes: A - BR 207, B - BR 197, C - BR 237, D - Br 125, E - BR 198, F - BR 224, G - BR 144, H - BR 256, I - BR 14.

Fig. 2: Electrophoresis of RNA extracted from rotavirus strain grown in cell culture MA - 104. Lanes: A - rotavirus-positive faecal sample (internal laboratory standard), B - BR 14, C - BR 197, D - BR 224, E - BR 207, F - strain Lincoln, G - BR 256.
Fig. 3: Cytopathic effect induced by rotavirus strain BR 207 at the fifth passage in MA - 104 cells. A: 24 hours post inoculation, B: uninfected control culture.

Fig. 4: Specific immunofluorescence in the cytoplasm of MA - 104 cells. A: strain BR 207, B: strain BR 224.

Fig. 5: Viral particle of isolated rotavirus strain BR 207 (negative staining).