Comparison of ELISA, SPACE, and electron microscopy for the routine diagnosis of rotavirus infection

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SUMMARY Previous studies on the serological diagnosis of rotavirus infection have utilised locally produced antibodies. In this study we have compared two commercially produced assays, an ELISA (Rotazyme, Abbott) and a newly developed assay—solid phase aggregation of coupled erythrocytes (SPACE) (Wellcome Research Laboratories), with electron microscopy (EM).

The SPACE test appeared less sensitive than EM. The ELISA was shown to be as sensitive as EM but more versatile. Our experience suggests that the ELISA could be successfully incorporated into the routine of any diagnostic laboratory.

Rotaviruses are now accepted as the most important known aetiological agents in acute non-bacterial gastroenteritis in infancy and childhood. Since the first recognition of the virus in human stools, electronmicroscopy (EM) has been the major method by which diagnosis has been established. Because of the limitations of EM however, other diagnostic techniques have been developed. While cell culture has been achieved, it lacks routine practicality, and the alternative diagnostic approach is serological. A variety of techniques have been studied, including complement fixation, counter current immunoelectrophoresis, radioimmunoassay (RIA) and enzyme linked immunosorbent assay (ELISA). Of these RIA and ELISA are considered to achieve or exceed the sensitivity of EM.

More recently the technique known as solid phase aggregation of coupled erythrocytes (SPACE) has been described for detecting rotavirus. The technique is similar to solid phase RIA and ELISA but utilises antibody-coated red blood cells as the indicator system instead of radioisotopes or enzymes.

Until recently however none of these serological tests has been available commercially, thus limiting their widespread application. An ELISA for rotavirus is now marketed (Rotazyme, Abbott) and a SPACE test may be marketed in the future.

The current study was therefore undertaken to compare EM, ELISA and SPACE for detecting rotavirus and to assess the use of these techniques in a routine diagnostic laboratory.

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Material and methods

Eighty-nine consecutive stool samples submitted to the Department of Clinical Microbiology, UCH, were evaluated. Each sample was coded and tested without knowledge of the results obtained by other tests.

Electron microscopy

Approximately 1 ml of stool was diluted 1/10 in distilled water and shaken. The suspension was then mixed with an equal volume of arcton 113 (ICI) centrifuged at 3000 rpm in a bench centrifuge for 30 min. The supernatant was filtered through a millipore SA and the filtrate spun at 35000 rpm (139000 g) for one hour in an OTD-50 ultracentrifuge (Swingout Rotor AH650).

The resultant pellet was resuspended in 0-1 ml of distilled water, stained with 2% phosphotungstic acid, and examined in a Hitachi H500 electron microscope.

SPACE

SPACE was carried out as previously described using materials donated by Wellcome Laboratories (Dr J Almeida). Flexible mitrotitre plates with U-shaped wells were coated with 100 µl guinea pig antibody in 0-01 M carbonate buffer pH 9-5, per well, by incubation at 37°C for 2 h in an airtight plastic box. The plates were washed three times in 0-05 Tween 20 in phosphate-buffered saline (PBS).
Suspensions (1/10 and 1/100) of each stool sample were prepared in PBS. Quantities (50 μl) of suspension were added per well, the plates covered, and incubated overnight at 4°C in an airtight box. Faecal suspensions were evacuated into Chloros using a vacuum pump. The plates were washed twice in PBS Tween and once in PBS. A further 50 μl of a 0.5% suspension in PBS of goat red blood cells (RBC) coated with bovine antitrovirus antibody were added to each well. The plates were left for two hours at room temperature in a vibration free atmosphere to allow settling of the RBC. Positive results were recognised by forming a shield of cells, negative results by forming a button. Positive and negative controls were included on each plate.

ELISA
ELISA was carried out using Rotazyme (Abbott) according to the manufacturer's instructions.

An approximately 40% stool suspension was prepared in PBS and spun to clarify at 3000 rpm in a bench centrifuge for 5 min. The supernatant (200 μl) was pipetted into one well of a reaction tray to which a bead coated with guinea pig antitrovirus antibody was added, and incubated overnight at room temperature. Each bead was then washed 4 times with distilled water. In addition, 200 μl of rabbit antitrovirus antibody conjugated to horseradish peroxidase were added and incubated for one hour at 45°C in a water bath.

Each bead was again washed and transferred to a 3 × ½" plastic tube to which was added 200 μl o-phenylenediamine 2 HCl (OPD) substrate. The tubes were incubated in the dark for 15 min and the enzyme substrate reaction stopped by adding 1 ml N HCl. Results were read either in a spectrophotometer at 492 nm or by comparison with a completely colourless negative control.

Neutralisation
Stool suspension was incubated for 1 hour at 37°C with an equal volume of high titre antitrovirus antibody (human convalescent serum). The ELISA procedure was then repeated as above and the result compared with the stool suspension diluted 1/2 with PBS. Neutralisation was considered to have occurred if either complete or 50% reduction in absorbance resulted.

Results
Eighty-nine stool samples were tested by each method and the overall results are presented in the Table. Forty-nine samples were negative by all tests. Thirty-seven samples were positive by EM. All these were also positive by ELISA, but only 33 positive by SPACE. Of these 33, 32 were positive by both 1/10 and 1/100 dilution and 1 at 1/10 dilution only. Of the four negative by SPACE, three were weakly positive only, by EM and ELISA, suggesting diminished sensitivity of the SPACE test.

One stool was positive by ELISA and negative by other tests. This sample and a random selection of other ELISA positive samples were retested after prior incubation with high titre rotavirus antibody. All such samples then gave negative results by ELISA suggesting that specific antigenic reactivity had been neutralised.

Two stools were initially positive by SPACE and negative by other tests. One stool on retesting was negative by all tests, and the initial positive result was considered to result from difficulty in interpretation of the pattern of the RBC. The second stool was repeatedly weakly positive by SPACE alone. EM revealed small cubic viral particles against which calf serum was found to have antibodies. The guinea pig serum was shown to complex the particles (Dr J Almeida, personal communication, 1980) and the positive result was therefore likely to have resulted from this second antigen-antibody reaction.

Discussion
Since rotaviruses were first incriminated as important causes of gastrointestinal illness, EM has been the mainstay of diagnosis and the yardstick by which other diagnostic methods are judged. EM is, however, time-consuming, costly in terms of both equipment and labour, and not always readily accessible. Recent experience with handling an outbreak of rotavirus infection has confirmed the difficulties of processing a large number of specimens in a short period.9

Of the many techniques available for detecting microbial antigens, RIA and ELISA are considered the most sensitive. As a routine diagnostic tool RIA has well recognised disadvantages.10

ELISA techniques have been widely used for detecting rotavirus but all such publications have concerned locally produced antibodies. Access to these materials is not possible for the majority of diagnostic laboratories; which generally must rely on commercial kits.
This study was therefore designed to evaluate assays which are, or may become, commercially available. Each depends on absorption of rotavirus antigen from faecal suspensions on to solid phase, with subsequent recognition of antigen by an enzyme-labelled antibody or red cell antibody indicator system. In terms of practical laboratory use, both were similar permitting results to be obtained the day after receipt of samples. Each can be adapted to provide results in one working day but overnight incubation of sample with solid phase is convenient for batching samples and may enhance sensitivity.

SPACE, however, appeared less sensitive. Results were also sometimes difficult to read, because the settling patterns of the RBC were not clear cut.

The full significance of the false positive reactions is difficult to evaluate on the number of specimens examined.

Rotazyme had the sensitivity of EM. On one occasion ELISA was positive and EM negative. This may reflect increased sensitivity, or that EM will detect only intact viral particles while immunological techniques are capable of detecting antigenic fractions. The third possibility of a false positive reaction was ruled out following successful neutralisation with specific rotavirus antibody. The problem of false positive results does suggest however that the specificity of a positive result should be confirmed by a blocking technique. Regrettably the necessary antibody is not an integral part of the kit. The colour reactions are nevertheless easy to read even without the aid of a spectrophotometer, and the current kit enables single assays to be performed. This study suggests that Rotazyme is a sensitive practical test which would allow the diagnosis of rotavirus shedding to be undertaken by any laboratory.

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References


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