A particle counting immunoassay for the direct detection of *Clostridium difficile* serogroup specific antigen in faecal specimens

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Summary. The potential of a particle counting immunoassay (PACIA) for the direct detection of *Clostridium difficile* serogroup G specific antigen in faecal specimens was evaluated. F(ab')₂ fragments from a rabbit anti-serogroup G antiserum were covalently coupled to carboxylated latex beads. This reagent was mixed with acid extracts of faecal specimens and the reaction was assayed with an optical counter which discriminated unagglutinated from agglutinated latex particles. Culture for *C. difficile*, faecal cytotoxin detection, PACIA and serogrouping of *C. difficile* isolates were performed on 249 stools. Of the 71 culture-negative specimens, none gave a positive result in the cytotoxin assay or in PACIA. Faecal cytotoxin was detected in 100 of the 178 culture-positive specimens. PACIA was positive for 63 of the 71 faecal specimens that yielded serogroup G *C. difficile* on culture. PACIA gave negative results for all other culture-positive stools tested with one exception, from which a serogroup A7 *C. difficile* strain was isolated. PACIA detection of serogroup G antigen in faecal specimens showed a sensitivity of 88.7%, a specificity of 99.7%, a predictive value of a positive culture with a serogroup G strain of 98.4%, and a predictive value for specimens that were culture-negative for a serogroup G strain of 95.6%. The results indicate that PACIA with specific antiserum is a rapid and reliable method for detecting serogroup specific antigens of *C. difficile* in faecal specimens. Clinical applications of the method are discussed.

Introduction

Serogrouping of *Clostridium difficile* by slide agglutination with rabbit antisera allows the differentiation of 10 serogroups. The epidemiology of *C. difficile* infection, particularly nosocomial transmission in hospital outbreaks, has been studied by this and other typing methods such as polyacrylamide gel electrophoresis (PAGE), immunoblotting, plasmid profiles or restriction endonuclease profiles. Several correlations between serogroup, toxigenicity and clinical course have been observed. Toxigenic isolates of serogroups C, H and K are usually implicated in colitis in adults, but isolates of serogroups B, F and X are isolated almost exclusively from asymptomatic children. Serogroup F strains produce a cytotoxin but no toxin A and are unable to induce colitis in animal models, and strains of serogroups B, D and I are non-toxigenic.

Correlations between serogroups and PAGE were also found. Specific profiles were associated with each of the 10 serogroups, except serogroup A, for which 12 different profiles were observed, designated A1–A12. Later we showed that the cross-agglutination among serogroup A strains was due to a common flagellar antigen, and removal of the flagella allowed specific slide agglutination of the 12 PAGE subgroups of A with 12 corresponding antisera. A specific somatic antigenic determinant was identified for each serogroup and the corresponding protein band was located on PAGE profiles.

Here, we describe the direct detection of serogroup G specific *C. difficile* antigen in faecal specimens by a particle counting immunoassay (PACIA). This method is a sensitive homogeneous non-isotopic latex agglutination assay in which the extent of the immune reaction is assayed by counting unagglutinated latex particles after the suspension of antibody-coated particles has been incubated with the antigen. Materials and methods

Faecal specimens

Stool samples from patients at the St-Luc University Hospital of Brussels suspected of having *C. difficile*-associated disease were submitted to the routine laboratory. Those examined in this study comprised:
135 consecutive culture-positive stools; 71 randomly selected culture-negative stools; and 43 additional faecal specimens from which serogroup G C. difficile strains had been isolated. All specimens were cultured on cycloserine cefoxitin fructose agar (CCFA) for C. difficile, and assayed directly for cytotoxin on HeLa cells with specific neutralisation by C. sordellii antiserum. When not processed immediately for antigen detection by PACIA, faeces were stored at −80°C. C. difficile isolates were identified as described previously. They were serotyped by slide agglutination, and serogroup A strains were further differentiated on the basis of PAGE profiles. Isolates of C. difficile were tested for cytotoxin production in the specific neutralisation HeLa cell assay, as for faeces, but with the filtrate of a 5-day culture in anaerobic liquid medium (LM) as described previously.

**Antiserum preparation**

A rabbit antiserum against the serogroup specific antigenic determinant was prepared with ATCC 43599, the serogroup G reference strain, as described previously. Briefly, cells from an 18-h culture in LM were sonicated and centrifuged. Two volumes of the supernate were mixed with one volume of SDS buffer (25 mM Tris·hydrochloride with 2-mercaptoethanol 15% v/v, glycine 30% v/v, sodium dodecyl sulphate 10% w/v and bromophenol blue 0.1%). After heating at 100°C for 5 min, 1400 μl of the homogenate were loaded on to a 20 cm × 25 cm polyacrylamide 10% gel in single 190-mm wide wells, and electrophoresed at room temperature for 15 h at 60 V. Low mol.-wt standards were from BioRad Laboratories. Gels were stained with Coomassie Blue. The band corresponding to the specific antigen was excised and electro-eluted in dialysis bags with Tris-glycine buffer, pH 8.3, containing SDS 0.01%. After centrifugation, the liquid was dialysed against ammonium bicarbonate buffer 1%, pH 7.8, and concentrated by lyophilisation. Antiserum was raised in rabbits by repeated intradermal injection with 2 ml of a 1:1 emulsion of purified serogroup G protein solution (0.5 mg/ml) in incomplete Freund’s adjuvant. The rabbits were given weekly injections for 6 weeks, then monthly for 2 months, and then bled.

**PACIA latex reagents**

Two reagents were prepared, one from the serum of rabbits immunised as described, and the other, control, from unimmunised animals. The IgG fraction was isolated by treatment with 6,9-diamino-acridine lactate (Rivanol 0.4%), pH 8.0, followed by precipitation with 4·1 M ammonium sulphate, pH 7.5. F(ab')2 fragments were prepared by peptic digestion of purified IgG in 0·1 M acetate buffer, pH 4·5, for 20 h at 37°C with an enzyme:protein ratio of 1:50 w:w. The reaction was stopped by increasing the pH to 7 with 1 M Tris. F(ab')2 fragments were purified by chromatography on Ultrogel AcA 4-4 and covalently coupled to carboxylated 0.8-μm diameter latex beads (Estafor K 150, Rhône-Poulenc, Courbevoie, France) by the carbodiimide method. The coated particles were stored at −20°C in small volumes. Before use, the latex particles were diluted 200-fold with GBS (0·1 M glycine, 0·17 M NaCl, pH 9.2) containing bovine serum albumin 10 g/L, and sodium azide 40 mg/L, and sonicated for 10 s.

**PACIA method**

One ml of faeces was homogenised with 2 ml of saline. The suspension was adjusted to pH 2 with HCl, shaken at room temperature for 30 min and centrifuged at 5000 g for 30 min. The supernate was adjusted to pH 7 with 1 M Tris buffer. For cultures of C. difficile, 1 ml of a 24-h culture on LM was used rather than 1 ml of stool sample. Twenty-five μl of this acid extract was mixed with 25 μl of the first additive (3 M NaCl with Dextran T 500 20 g/L; Pharmacia, Uppsala, Sweden) and 25 μl of the second additive (GBS with normal rabbit serum 100 ml/L and F(ab')2 fragments of IgG from non-immunised rabbits 25 g/L) and incubated at 37°C for 40 min. Twenty-five μl of antibody-coated latex suspension were added and the mixture was vortex mixed at 37°C for 40 min in a purpose-built incubator-agitator. After stopping the reaction by the addition of 0·75 ml of GBS, unagglutinated particles were counted in an optical counter which discriminated the unagglutinated from the agglutinated latex particles. The concentration of antigen was inversely proportional to the number of unagglutinated particles.

**Results**

The PACIA was calibrated with 10-fold dilutions of the supernate from a 2-day culture of ATCC 43599 in LM centrifuged at 5000 g for 30 min (figure). Uninoculated broth controls gave a peak count considered as 100%, corresponding to the absence of specific agglutination. A peak count < 90% of the control was considered to be a positive result for serogroup G.
Table. Results of PACIA for the various serogroups isolated

<table>
<thead>
<tr>
<th>Serogroup</th>
<th>Number of strains (%) isolated</th>
<th>Number of faeces (%) cytotoxin positive</th>
<th>PACIA positive</th>
</tr>
</thead>
<tbody>
<tr>
<td>G</td>
<td>71 68 (96)</td>
<td>43 (60)</td>
<td>63 (89)</td>
</tr>
<tr>
<td>A</td>
<td>20 16 (80)</td>
<td>14 (70)</td>
<td>1 (5)</td>
</tr>
<tr>
<td>C</td>
<td>42 32 (76)</td>
<td>27 (64)</td>
<td>0</td>
</tr>
<tr>
<td>D</td>
<td>4 0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>H</td>
<td>25 19 (76)</td>
<td>11 (44)</td>
<td>0</td>
</tr>
<tr>
<td>F</td>
<td>1 1 (100)</td>
<td>1 (100)</td>
<td>0</td>
</tr>
<tr>
<td>K</td>
<td>12 5 (45)</td>
<td>4 (33)</td>
<td>0</td>
</tr>
<tr>
<td>X</td>
<td>3 0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Totals</td>
<td>178 141 (79)</td>
<td>100 (56)</td>
<td>64 (36)</td>
</tr>
</tbody>
</table>

antigen. Antigen was detectable in the culture supernate up to the 10^4 dilution (figure).

Of the 71 culture-negative specimens, none gave a positive result in the cytotoxin assay or in PACIA. From the 178 culture-positive specimens, C. difficile strains were isolated, serogrouped and investigated for cytotoxin production in vitro. All isolates were typable by slide agglutination, and were of serogroups A, C, D, F, G, H, K or X. PAGE subtypes of the 20 isolates of serogroup A were: nine in A1, three each in A10, A8 and A5; one in A7; and one with a previously undocumented PAGE profile.

Faecal cytotoxin was detected in 100 of the 178 culture-positive specimens, including 43 of the 71 stool samples that yielded serogroup G strains. Of the 178 isolates of C. difficile, 141 produced cytotoxin in vitro. The results obtained by PACIA are compared with isolate serogroup in the table. PACIA gave positive results in 63 of the 71 faecal specimens from which a strain of serogroup G was isolated. PACIA gave negative results for all other stool samples, with one exception, from which a strain of serogroup A7 was isolated. The performance of PACIA in the detection of serogroup G antigen in faecal specimens was: sensitivity, 88.7%; specificity, 99.7%; predictive value of a positive reaction, 98.4%; predictive value of a negative reaction, 95.6%.

The eight serogroup G isolates from faecal specimens that gave negative results by PACIA were grown in LM which was then tested by PACIA. All gave a positive result. By contrast, the serogroup A7 strain from the stool that was positive in PACIA gave a negative PACIA result.

Discussion

Detection of bacterial antigen in biological specimens has the obvious advantage of reducing the diagnostic delay, provided that it is specific and sensitive. An early latex agglutination assay for C. difficile toxin A in faeces was later demonstrated to be specific not for the toxin but for an enzyme. The test is still used in some laboratories for screening. More recently, several commercial kits for the detection of toxin A or for toxins A and B have yielded promising results. By contrast, our approach, having previously identified serogroup-specific antigens and established correlations between these serogroups and pathogenicity, has been to investigate methods that detect serogroup-specific antigens in stool samples.

PACIA is an immunological method which has been exploited in many diagnostic applications, and which can attain a sensitivity in the femtogram/ml range. This study demonstrates that PACIA is able to detect serogroup G-specific antigen in faecal specimens with a sensitivity of 88.7% and a specificity of 99.4%. We believe that the acid extraction of antigens from faeces described here may be, in part, responsible for the excellent results obtained. Other extraction procedures, such as simple sonication, yielded much less convincing results (unpublished data).

There were eight false negative and one false positive PACIA results in this series. The false negatives indicate a problem with assay sensitivity rather than specificity, as culture supernates from each of the eight isolates from these specimens gave positive PACIA results. The single false positive result was probably not due to antigenic cross-reaction with the serogroup A7 C. difficile isolate from this specimen, as PACIA results on culture supernates of this isolate were negative. This apparently anomalous positive reaction was more probably due to the additional presence of a serogroup G strain in such low relative numbers that it remained unrecognised as a distinct isolate. The concomitant presence of two distinct strains of C. difficile in the same specimen has been reported. Unfortunately we did not have the opportunity to re-culture the stool to check this possibility.

The typing aspect of serogrouping, and the correlation between serogroup and virulence of C. difficile, make the PACIA approach well suited to clinical practice. Within a few hours, we were able to detect not only the presence of C. difficile in stools, but also to give an indication of the likely virulence of the organism and to specify a serogroup for epidemiological tracing. The main interest of rapid typing in routine practice is the ability to identify patients who have been colonised during hospital outbreaks. Nosocomial transmission of C. difficile has been well demonstrated. Severe outbreaks caused by a single epidemic strain have been reported, and some serogroups are thought to show particular predilection to cause outbreaks. Toxigenic strains of serogroup C, which are more resistant to antibiotics than other serogroups, are most often implicated in outbreaks. Conventional typing of isolates, by immunoblotting, PAGE profile or slide serogrouping, is much slower than the PACIA method described here. During an outbreak, the early PACIA detection of patients colonised with the epidemic strain could allow more effective preventive measures.

Equally, serogrouping results may be important in the detection of colonisation with strains that protect
against symptomatic infection. The protective role of colonisation with non-toxigenic strains that prevent subsequent colonisation by toxigenic strains has been demonstrated in animal models. Borriello and Barclay showed that prior colonisation of hamsters with non-pathogenic strains protects the animals against ileo-caecitis when challenged with a toxigenic strain. 26 Similarly, Depiret et al. showed that mice colonised by a serogroup F strain that produces a cytotoxin but no toxin A do not develop intestinal disease and are protected when challenged subsequently with a serogroup C strain which would otherwise kill the animals. 28 In man, administration of a non-toxigenic isolate to patients suffering from C. difficile-associated colitis at the end of vancomycin therapy can prevent relapse. 29 Some serogroups (B, D and I) do not produce either of the toxins implicated in disease, and another (F) produces only the cytotoxin; isolates of these serogroups are not pathogenic. 30 Rapid detection and serogrouping in patients colonised with these serogroups could contra-indicate antibiotic therapy.

We regard the work described here as a preliminary step. Our choice of serogroup G as a proving system was guided by the early availability of a specific antiserum, rather than particular clinical interest. Although c. 25 serogroups, including the PAGE subgroups within serogroup A, are known, in our experience of serogrouping > 6000 isolates, the eight most frequent serogroups cover about 80% of all isolates (unpublished data). Moreover, the more clinically significant serogroups are restricted to six: C, G, H, K, A1 and A8. The problems of the multiplicity of reagents required for a workable rather than a comprehensive PACIA are not as formidable as might appear at first sight. The fact that some serogroups encompass toxigenic and non-toxigenic strains might be overcome by testing in PACIA and with rapid, direct toxin detection kits. Such a system would offer a combination of rapid detection of the organism and the toxins, and typing information that would be valuable in diagnosis, treatment and cross-infection control.

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