Detection and Identification of Yersinia pseudotuberculosis and Pathogenic Yersinia enterocolitica by an Improved Polymerase Chain Reaction Method

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We developed a polymerase chain reaction method in order to detect and identify both Yersinia pseudotuberculosis and pathogenic Yersinia enterocolitica. Polymerase chain reaction was performed by using a mixture of primers against the inv gene from Y. pseudotuberculosis and the ail gene from pathogenic Y. enterocolitica. Further addition of primers against the plasmid-coded virF gene from Y. enterocolitica made it possible to detect a virulence-associated gene of both species at the same time. This method was proved to be an adequate and convenient procedure for routine detection and identification of these bacilli.

Recently, a method for the rapid detection of pathogenic Yersinia species, Y. pseudotuberculosis and Y. enterocolitica, by the polymerase chain reaction (PCR) technique was reported by Wren and Tabaqchali (8) and Fenwick and Murray (2). They used primers based on the vir F gene and the *ail* gene, respectively. By these methods, it was impossible to differentiate between the two bacilli, because DNA fragments of the same size were amplified by virF primers and a product from the ail gene was detected only from pathogenic Y. enterocolitica and not from Y. pseudotuberculosis. In the case of Yersinia infection, however, Y pseudotuberculosis and Y. enterocolitica cause almost the same symptoms, so it is impossible to differentiate between the bacilli on the basis of the symptoms that result from them. Therefore, we developed ^a rapid PCR method for the detection and differentiation of both Yersinia species and the detection of their virulence-associated genes directly from their cultures. The sensitivity of PCR detection was also examined.

The bacterial strains used in this study are listed in Table 1. These included 25 strains of virulence plasmid-harboring $(p⁺)$ *Y. pseudotuberculosis* belonging to each serotype from patients and a wild animal (raccoon dog), 6 strains of virulence plasmidless (p^-) *Y. pseudotuberculosis* (serotypes 2a, 2b, 4a, 5a, and 6) from river water, ¹² strains of Y enterocolitica (p⁺; serotypes O:3 and O:8) from patients, 2 patient-derived strains of Y. enterocolitica of which the virulence plasmids were lost after long storage $(p^-;$ serotypes $O:3$ and $O:8$), 13 strains of nonpathogenic \overline{Y} . enterocolitica from river water, 11 strains of Yersinia frederiksenii, 17 strains of Yersinia intermedia from a wild animal (raccoon dog), and other isolates from patients (4 isolates of Salmonella spp., 1 isolate of Campylobacter jejuni, 1 isolate of enteroinvasive Escherichia coli, 2 isolates of enterotoxigenic E. coli producing heat-stable or heat-labile enterotoxin, and 4 isolates of enteropathogenic E. coli with the following serotypes: 055:H7, 0111:H21, 0126:H27, and 0127:H21). Plasmids were isolated by the method of Kado and Liu (6) and detected on agarose gel. For PCR, we selected a 295-bp

region of the inv gene coding for 594 to 692 amino acid residues of invasin protein on the chromosome of Y pseudotuberculosis (5) as ^a target DNA. Since this region lacks homology with Y . enterocolitica (9) , we could synthesize primers to amplify a 295-bp DNA fragment only from Y. pseudotuberculosis; the primers were 5'-TAAGGGTAC TATCGCGGCGGA-3' and 5'-CGTGAAATTAACCGTCA CACT-3'. For the purpose of detecting pathogenic Y. enterocolitica, the 664- to 833-nucleotide region of the ail (7) gene related to its cell adhesion was selected as a target, and a 170-bp product was expected to be amplified by the following primers: 5'-ACTCGATGATAACTGGGGAG-3' and ⁵'- CCCCCAGTAATCCATAAAGG-3'. The primers for the $virF$ (1) gene used by Wren and Tabaqchali (8) to detect pathogenic Yersinia spp. harboring a 67- to 72-kb virulence plasmid (3, 4) were used for amplification of a 591-bp product as previously described (8). Each primer set for the inv or ail gene was used for the detection of each gene, and a mixture of primers against the inv, ail, and vir F genes was used for the detection of one or more genes among the inv, ail, and virF genes. Template DNAs for PCR were prepared as follows. (i) From colonies, each isolate was suspended in autoclaved distilled water to achieve a concentration of 108 CFU/ml and boiled for 10 min, and 5 μ l of each sample (10⁵) CFU) was applied to PCR. (ii) In order to examine the detection limit for PCR, ^a series of 10-fold dilutions of Y pseudotuberculosis or Y. enterocolitica strains with river water or autoclaved water was made. Cells of each dilution were collected by centrifugation and suspended in 20 μ l of proteinase K buffer (50 mM Tris-HCl [pH 7.5], 5 mM CaCl₂). After being freeze-thawed three times, $0.5 \mu l$ of proteinase K (5 mg/ml in 10 mM $CaCl₂$ solution) was added to the suspension, which was incubated at 45°C for ¹ h, and PCR was performed with the suspension. DNA samples were amplified in a total of 50 μ l of the following reaction mixture: 50 mM KCl-10 mM Tris-HCl (pH 8.3)-1.5 mM $MgCl_{2}$ - 0.001% (wt/vol) gelatin; 100 μ M each dATP, dCTP, dGTP, and dTTP (Perkin-Elmer Cetus); $0.05 \mu M$ inv primers, 0.1 μ M each ail and virF primers; 0.5 μ g of RNase A (Sigma); and 0.5 U of AmpliTaq DNA polymerase (Perkin-Elmer Cetus). The samples were overlaid with a drop of mineral oil and amplified by PCR with the Thermal Sequencer

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 $a^a p⁺$, virulence plasmid harboring.

 $b \, \bar{p}$, virulence plasmidless.

 c p⁻ strains on account of long storage.

ST, heat-stable enterotoxin.

LT, heat-labile enterotoxin.

(IWAKI), which consisted of the following: predenaturation at 94°C for 1 min, 25 cycles of denaturation at 94°C for 0.5 min, primer annealing at 55°C for 1 min, extension at 70'C for 2 min, and further extension at 70°C for 5 min. In the case of the examination of the detection limit, 30 PCR cycles were repeated after the addition of AmpliTaq DNA polymerase. After PCR amplification, $10 \mu l$ of each PCR product was run on ^a 1.5% agarose gel at ^a constant voltage of ¹⁰⁰ V in TAE buffer (40 mM Tris-acetate, ¹ mM EDTA [pH 8.0]). The gel was stained with ethidium bromide and photographed under UV light.

Each primer against the inv or ail gene was confirmed to amplify a 295-bp product from a Y. pseudotuberculosis (p^+) strain or a 170-bp product from a *Y. enterocolitica* (p^+) strain (Fig. 1, lanes 1 and 2, respectively). With a mixture of inv, ail, and virF primers, PCR could differentiate Y. pseudotuberculosis (p^{\dagger}) from Y. enterocolitica (p^+) in suspensions containing only one or both species; 591- and 295-bp products were amplified from a Y. pseudotuberculosis (p^+) strain, while 591- and 170-bp products from Y. enterocolitica $(p⁺)$ were amplified (Fig. 2 [lanes 1 and 3] and 1 [lane 3]). The results of PCR for Yersinia species and other isolates are shown in Fig. 2 and Table 1. All of the strains of a variety of serotypes of Y. pseudotuberculosis (p^+) from patients and raccoon dogs gave a positive reaction for the *inv* gene and also for the *virF* gene, while *Y. pseudotuberculosis* strains (p^-) ; serotypes 2a, 2b, 4a, 5a, and 6) from water samples were positive only for the inv gene. Y. enterocolitica strains $(p^+;$ serotypes O:3 and O:8) from patients gave positive reactions for both the ail and the vir F genes but a negative reaction for the inv gene, and virulence plasmidless Y. enterocolitica strains (p^- ; serotypes O:3 and O:8) from patients were positive only for the ail gene. Avirulent Yersinia strains from wild animals and water samples (Y. enterocolitica, Y. frederiksenii, and Y. intermedia) and other

FIG. 1. Specific amplification of 591-, 295-, and 170-bp products by PCR. Lanes: 1, 295-bp product from a Y. pseudotuberculosis strain (p^+) with inv primers; 2, 170-bp product from a pathogenic Y. enterocolitica (p^+) strain by with ail primers; 3, 591-, 295-, and 170-bp products from a suspension carrying both strains used above with ^a mixture of primers; M, 123-bp DNA ladder marker.

isolates from patients (Salmonella spp., C. jejuni, enteroinvasive E. coli, enterotoxigenic E. coli, and enteropathogenic $E.$ coli) gave negative reactions for the inv, ail, and vir F genes. These results showed that Y. pseudotuberculosis, pathogenic Y. enterocolitica, and other isolates could be differentiated by this PCR method. Therefore, we highly recommend this assay as a method of rapid identification and differentiation of Y . pseudotuberculosis and pathogenic Y . enterocolitica strains. Furthermore, the detection limit of PCR for *Y. pseudotuberculosis* and pathogenic *Y. enteroco*litica was examined by using the *inv* and *ail* genes, respectively. The result showed that more than 10^3 to 10^4 CFU of bacteria in a suspension of river water was necessary for the detection of ^a product amplified after 30 PCR cycles (Fig. 3). However, fewer than 10 bacteria were detectable by the

FIG. 2. Differentiation among Yersinia bacilli by amplified PCR products with a mixture of inv, ail, and virF primers. Lanes: 1, Y. p seudotuberculosis $(p^+); 2, Y$. pseudotuberculosis $(p^-); 3, Y$. enterocolitica $(p^+); 4$, nonpathogenic Y. enterocolitica; 5, Y. enterocolitica (p⁻); 6, Y. frederiksenii; 7, Y. intermedia; M, 123-bp DNA ladder marker.

FIG. 3. Detection limit of PCR with inv and ail primers. Template DNAs were prepared from river water containing 10^5 , 10^4 , 10^3 , 10^2 , and 10^1 CFU of \hat{Y} . *pseudotuberculosis* (A) and 10^6 , 10^5 , 10^4 , 10^3 , 10^2 , and 10^1 CFU of pathogenic Y. enterocolitica (B) and used for PCR.

repetition of 30 PCR cycles after the addition of AmpliTaq DNA polymerase (data not shown). We got ^a similar result by using autoclaved water as a suspension medium (data not shown). Therefore, it is also suggested that this method is applicable for the detection of pathogenic Yersinia spp. from water samples for the purpose of epidemiological research.

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