ECA-immunogenicity of Proteus mirabilis strains

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Abstract

Introduction: Bacteria of the genus Proteus are opportunistic pathogens and cause mainly urinary tract infections. They also play a role in the pathogenesis of reactive arthritis (RA). Patients suffering from Yersinia-triggered RA often carry high titers of antibodies specific to enterobacterial common antigen (ECA). The immunogenicity of ECA has not received much attention thus far and studies have focused mainly on the ECA of Escherichia coli and Yersinia enterocolitica. In this paper the ECA-immunogenicity of Proteus mirabilis is elucidated using two wild-type strains (S1959 and O28) as well as their rough (R) derivative strains R110/1959, which expresses lipopolysaccharide (LPS) with a full core, and R4/O28, which expresses LPS with only an inner core.

Materials and Methods: Rabbit polyclonal antisera were produced by immunization with boiled suspensions of the four P. mirabilis strains. The antisera were tested for the presence of antibodies specific to ECA by Western blotting using glycerocephospholipid-linked ECA (ECAPG) of Salmonella montevideo as antigen. Lipopolysaccharide (LPS) was isolated from the four strains by the hot phenol/water procedure in which ECA PG is co-extracted with LPS and by the phenol/chloroform/petroleum ether extraction that results in the isolation of LPS and/or LPS-linked ECA (ECA LPS) free of ECAPG. The LPS preparations were tested for the presence of ECA by Western blotting using ECA-specific antibodies.

Results: The results demonstrated that all four P. mirabilis strains were ECA immunogenic. The rabbit antisera immunized by the four strains all contained ECA-specific antibodies. Analysis of the LPS preparations demonstrated that the P. mirabilis wild-type strains O28 and S1959 and the Ra mutant strain R110/1959 expressed ECA_LPS, suggesting that it induced the anti-ECA antibody responses. Only the presence of ECA_PEG could be demonstrated in the Rc mutant strain R4/O28.

Conclusions: These results therefore suggest that, similar to E. coli, LPS with a full core is also required as the acceptor of ECA for P. mirabilis strains to produce ECA_LPS. Since ECAPG is not immunogenic unless combined with some proteins, it is likely that ECAPG-protein complexes formed during the intravenous immunization with the Rc mutant strain R4/O28.

Key words: Proteus mirabilis, S and R strains, enterobacterial common antigen, lipopolysaccharide, ECA immunogenicity.

Abbreviations: ECA – enterobacterial common antigen, LPS – lipopolysaccharide, PAGE – polyacrylamide gel electrophoresis, PEG – phenol/chloroform/petroleum ether procedure, Ph/W – phenol/water procedure.

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INTRODUCTION

The genus Proteus contains five named species: P. mirabilis, P. penneri, P. vulgaris, P. myxofaciens, and P. hauseri, and three unnamed genomospecies 4, 5, and 6 (O’Hara et al. 2000a; O’Hara et al. 2000b; Róžalski 2002). These Gram-negative rods are widespread in the environment, for example in soil, water, and sewage, and, with the exception of P. myxofaciens, are common human intestinal and urinary tract pathogens (Róžalski et al. 1997). Under favorable conditions, Proteus infections may lead to serious complications, such as acute or chronic pyelonephritis, the formation of bladder and kidney stones, and catheter obstruction (Róžalski 2002). Evidence was recently presented that P. mirabilis is an etiopathogenic agent in rheumatoid arthritis, a chronic inflammatory polyarthritis disease (Rashid and Ebringer 2007).

Several virulence factors have been identified in Proteus strains. The most important are fimbriae-medi-
ated adherence, glycolcalyx formation, lipopolysaccharide (LPS), and probably also enterobacterial common antigen (ECA). Interestingly, antibodies against ECA are present in the serum of patients suffering from reactive arthritis (Granfors et al. 1989; Rastawicki 2007).

ECA, a cell surface lipoglycan, is present in all bacteria of the family Enterobacteriaceae, with the exception of the plant pathogenic bacterium Erwinia chrysanthemi (Kuhn et al. 1987; Rinno et al. 1980). ECA is a heteropolymer built of →3)-α-D-Fucp4NAc-(1→4)-β-D-ManpNAcA-(1→4)-α-D-GlcNAc(1→ trisaccharide repeating units (Ługowski et al. 1983). It occurs in three structurally different forms: (i) ECAPG, which is anchored in the outer membrane of the bacterial cell wall via L-glycerophospholipid, (ii) ECA_LPS, in which lipid A functions as a membrane-anchor and to which ECA is linked via the core oligosaccharide, and (iii) ECA_Cyo, which is deprived of a lipid carrier and is located in the periplasm (Kajimura et al. 2005; Kuhn et al. 1988). The ECAPG form is not immunogenic unless bound to a protein, which could occur in animal serum after intravenous injection. ECA_LPS, in contrast, is immunogenic. Presence of ECA_LPS has been demonstrated in E. coli Ra mutants possessing the full core type of coli R1, R4, and K-12 in their LPS (Kuhn et al. 1988) and also in E. coli Ra mutants with the full core type of coli R2 or R3 (Kasperkiewicz et al. 2004). We have also shown that Yersinia enterocolitica O:3 R mutants with a full core (chemotype Ra) as well as those with a truncated outer core or only an inner core (chemotype Rc) in their LPS are ECA immunogenic. In all of them, ECA was linked to the inner core of the LPS (Duda 2007a; Duda 2007b; Radziejewska-Lebrecht et al. 2003). Moreover, the Y. enterocolitica O:3 wild-type strain Ye75S was ECA immunogenic in rabbits and contained O-chain polysaccharide as well as ECA bound to the inner core of its LPS (Radziejewska-Lebrecht et al. 1998). In this study we wanted to examine whether ECA_LPS was also present in Proteus mirabilis, another species of Enterobacteriaceae.

MATERIALS AND METHODS

Bacterial strains

P. mirabilis strains O28 (wild type), S1959 (wild type), R4/O28 (rough, Rc chemotype), and R110/1959 (rough, Ra chemotype), described earlier (Radziejewska-Lebrecht et al. 1980; Radziejewska-Lebrecht and Mayer 1989; Vinogradov and Radziejewska-Lebrecht 2000; Vinogradov et al. 2000), were obtained from the collection of the Institute of Microbiology and Immunology in Łódź.

Antigens and serological assays

LPS preparations (LPS Ph/W) of P. mirabilis O28 (Vinogradov and Radziejewska-Lebrecht 2000) and S1959 (Vinogradov et al. 2000) were obtained by the hot phenol/water procedure of Westphal (Westphal and Jann 1965). LPS Ph/W was then subjected to the phenol/chloroform/light petroleum (PCP) procedure of Galanos (Galanos et al. 1969) to obtain LPS PCP. Only the LPS PCP preparation was obtained from the Ra mutant strain R110/1959 by straight PCP extraction of the dry bacterial mass (Radziejewska-Lebrecht and Mayer 1989). For the Rc mutant strain R4/O28, both the LPS Ph/W and LPS PCP preparations were acquired directly from dry bacterial mass (Radziejewska-Lebrecht et al. 1988). The LPS Ph/W preparations may contain ECA_PG, which co-extracts with LPS in the hot phenol/water procedure, whereas LPS PCP preparations are deprived of ECA_PG and contain LPS and, if present, also ECA_LPS. The LPS preparations and the control ECA_PG preparation from S. montevideo SH94 (kind gift from Drs. H. Mayer and H.-M. Kuhn, Max-Planck-Institute, Freiburg, Germany) were used as antigens in SDS/polyacrylamide gel electrophoresis (SDS/PAGE) and Western blotting as described (Radziejewska-Lebrecht et al. 2003). The experiments were repeated five times and representative results are shown in the figures.

Antibodies and antisera

Monoclonal antibody specific to ECA, mAb 898, recognizing the ManpNAcA epitope (Meier-Dieter et al. 1989) was a kind gift from Prof. D. Bitter-Suermann (Institute of Medical Microbiology, Hannover, Germany).

The monovalent antiserum against the ECA of E. coli O14:K7 was prepared by the adsorption of rabbit antiserum against E. coli O14:K7 by isogenic ECA-negative mutant bacteria. The antiserum, enriched in anti-ECA antibodies, was received from Dr. H. Mayer from the Max-Planck-Institute, Freiburg, Germany (Männel and Mayer 1978). Alkaline phosphatase-conjugated goat anti-mouse and goat anti-rabbit immunoglobulin antibodies (D0486 and D0487, respectively, DAKO) were used in a 1:2000 dilution as the secondary antibodies in Western blotting.

Polyclonal antiserum against P. mirabilis strains O28, S1959, and R110/1959 as well as antiserum against Staphylococcus albus (used as a negative control as it lacks cross-reactivity with LPS and ECA, which are typical antigens of Gram-negative bacteria) were obtained by immunization of New Zealand white rabbits with heat-killed bacterial suspensions. Immunization was carried out with the bacterial suspensions (1.5 × 10¹⁰ CFU/ml) at doses of 0.25, 0.5, and 1 ml of each over a period of 10 days. The antiserum was stored at –20°C. The antiserum was diluted in sterile 0.85% NaCl over a period of 51 days. The antiserum used in the last injection was stored at –20°C.
RESULTS AND DISCUSSION

In the present study, two smooth (serogroup O:28 and O:3) and two rough P. mirabilis strains were tested for their ECA-immunogenicity. The results (Fig. 1, panel A) demonstrate that ECA-specific antibodies were present in all the tested antisera. The ladder-like banding profile characteristic of ECA was present in the 10- to 50-kDa region, similar to that in the control reaction with ECA-specific mAb 898 (Fig. 1, panel B). The antisera against the Rc mutant R4/O28 and the Ra mutant R110/1959 (the ladder-like profile for this sample is only faintly visible) contained higher titers of anti-ECA antibodies than the antisera against the wild-type strains O28 and S1959. Although there is a possibility of variation among individual rabbits, it is likely that the differences in titers are due to small amounts of immunogenic ECA in the latter bacteria (see below) or to the possibility that in the wild-type strains the ECA chains could be masked by the abundant O-polysaccharides chains on the bacterial cell wall. However, the latter alternative might not be valid as the bacteria are most likely degraded and processed by rabbit antigen-presenting cells during immunization. ACKER et al. reported that on whole bacteria, ECA is better exposed on the cell surface of Rc mutant Ye75R bacteria (ACKER et al. 1981). In conclusion, the obtained results suggest that the ECA-immunogenicity of the smooth strains O28 and S1959 and of the Ra mutant R110/1959 was due to the presence of ECA_{LPS} whereas that of the Rc mutant R4/O28 was due to the presence of ECA_{PG}. As mentioned, the latter co-extracts with LPS in the Ph/W procedure.

Our previous studies on R mutants of Y. enterocolitica O:3 showed that monovalent antiserum against E. coli O14:K7 enriched with ECA-specific antibodies was a more sensitive tool in the detection of ECA than mAb 898, perhaps reflecting a lower affinity of the monoclonal antibody (Duda 2007a). In line with this, the immunostainings with the monovalent antiserum (Fig. 3, panel A) were stronger than with mAb 898 (Fig. 2, panel A). All the P. mirabilis LPS preparations gave positive reactions in the 10- to 50-kDa region corresponding to ECA compared with the control reaction with ECA_{PG} (Fig. 3, panel A), with the exception of the LPS PCP of R4/O28. As above, the gel was silver-stained after transfer of the samples to the nitrocellulose membrane (Fig. 3, panel B) and this confirmed the proper loading of the samples onto the gel. At the same time it indicated that the lack of reactivity of the R4/O28 LPS PCP with mAb 898 was likely due to the absence of ECA_{LPS} in this preparation. Of note, ECA preparations cannot be silver-stained as there are no vicinal hydroxyl groups in the ECA repeat unit required for the oxidation step in the staining; thus it is not visible in Fig. 2, panel B.

In conclusion, the obtained results suggest that the ECA-immunogenicity of the smooth strains O28 and S1959 and of the Rc mutant R110/1959 was due to the presence of ECA_{LPS} whereas that of the Rc mutant R4/O28 was due to the presence of ECA_{PG}. As mentioned, the latter co-extracts with LPS in the Ph/W procedure.

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a carrier, for example proteins (Kuhn et al. 1988), and this is the likely explanation for the ECA immunogenicity of the Rc mutant R4/O28.

The expression of ECA\textsubscript{LPS} by the Ra mutant R110/1959 with a full core region and not by the Rc mutant R4/O28 with a truncated core region suggests that in \textit{P. mirabilis}, similarly to \textit{E. coli}, the complete core region of LPS is a prerequisite for the linking of ECA to LPS. This conclusion is additionally supported by the following two observations. (i) The presence in the 1- to 5-kDa region of the membrane of the two clearly separated ECA-positive bands in the LPS PCP preparation of strain O28 and several similar bands in the LPS PCP of the Ra mutant R110/1959; these bands correspond to the migration of full core molecules (Fig. 3, panel A). (ii) The lack of ECA-specific immunostaining in the 1- to 5-kDa region of the LPS PCP preparation of the Rc mutant R4/O28, which possesses only the inner core in its LPS.

In contrast to previous observations (Kuhn et al. 1988), it is very likely that ECA\textsubscript{LPS} occurs abundantly in nature and one can envision that in wild-type strains...
under certain environmental conditions, a higher proportion of LPS molecules could be unsubstituted by O-polysaccharide and they could then become ECA-substituted. Our results therefore warrant further studies to find out whether other members of Enterobacteriaceae in addition to E. coli, Y. enterocolitica, and P. mirabilis express ECA-immunogenic ECA-LPS.

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