CLINICAL MICROBIOLOGY

Chronic bacteraemia due to *Staphylococcus epidermidis* after bone marrow transplantation

B. LINA‡, F. FOREY*, J. D. TIGAUD† and J. FLEURETTE*

*Département de Recherche en Bactériologie Médicale (EA 1655), Faculté de Médecine Alexis Carrel, rue Guillaume Paradis, 69372 Lyon Cedex 08 and † Service d’HématoLOGie, Pavillon E, Hôpital Edouard Herriot, 69437 Lyon Cedex 03, France

Summary. A chronic bacteraemia due to *Staphylococcus epidermidis* occurred in a patient undergoing allogeneic bone marrow transplantation. Forty-two *S. epidermidis* isolates were obtained from blood cultures over a period of 5 months. Isolates were separated into three groups by *SmaI* macrorestriction characterisation with pulsed-field gel electrophoresis (PFE-1, one isolate; PFE-2, 32 isolates; PFE-3, nine isolates). Differences were detected in antimicrobial susceptibility patterns among isolates belonging to group PFE-2. The two strains, PFE-2 and PFE-3, were both responsible for the chronic bacteraemia and were isolated during different admissions to the hospital. A central venous catheter was the portal of entry for PFE-2. DNA macro-restriction analysis with pulsed-field gel electrophoresis was helpful in the epidemiological investigation of this *S. epidermidis* chronic bacteraemia.

Introduction

Coagulase-negative staphylococci (CNS) are the organisms most frequently isolated from blood cultures, and although the majority of isolations represent contamination, clinically important infections do occur and these organisms represent a major cause of septicaemia or severe diseases related to intra-vascular catheter colonisation or to severe immunodepression. Because of the ubiquitous nature of *Staphylococcus epidermidis* and of its pathogenic potential, strain characterisation of this organism has become increasingly important.

The usual approach to determine the clinical relevance of *S. epidermidis* isolates is (i) to require repeated isolation from independent specimens from the patient, and (ii) that the isolates should share identical typing characteristics. Unfortunately, no single typing method is entirely satisfactory for *S. epidermidis*. Among recently developed epidemiological fingerprinting techniques, pulsed-field gel electrophoresis (PFGE) has been described as useful in strain characterisation of *S. epidermidis*, yielding stable and discriminatory restriction patterns.

In this study, 42 isolates of *S. epidermidis* from 41 blood cultures performed over a 5-month period in a patient undergoing allogeneic bone marrow transplantation (BMT) were compared by analysis of the *SmaI* macro-restriction patterns in PFGE.

Patient, materials and methods

Case report

A 33-year-old man was admitted to the hospital for treatment of chronic myeloid leukaemia. After the course of chemotherapy, the patient received an allogeneic BMT from his HLA Dr-identical brother. After transplantation, two routine blood cultures were performed daily through the catheter. On day 10, one blood culture yielded *S. epidermidis* (antibiotic-resistance type B). Engraftment was achieved on day 17. The patient developed a chronic graft versus host disease (GVHD) on day 20. Between day 27 and day 55, 16 blood cultures were positive with *S. epidermidis* (AtB-type B). Isolates were from blood cultures performed through the catheter and in peripheral samples. Changes were observed in antimicrobial resistance of one isolate (AtB-type C). Despite intravenous vancomycin (15 mg/kg daily) and oral pristinamycin (3 g daily), the patient remained febrile until the central venous line was removed on day 58. A *S. epidermidis* isolate with identical antibiogram to those of the blood cultures was obtained from this catheter (AtB-type B).
A second catheter was implanted and the antibiotic therapy was stopped. Of the 61 blood cultures performed through the second catheter before the first discharge, only three yielded *S. epidermidis* isolates, presenting different antimicrobial resistance patterns (AtB-types C and E). The patient was discharged afebrile on day 89.

He was admitted to the hospital on day 115 with a macroscopic haematuria due to thrombocytopenia, and diarrhoea due to acute GVHD. He received platelet transfusion in association with high doses of methylprednisolone and oral vancomycin. On day 123, he developed fever and two blood cultures drawn through the catheter yielded *S. epidermidis* (AtB-types B and E). Despite a course of intravenous vancomycin started on the same day (15 mg/kg daily), both peripheral and catheter blood cultures were positive daily with *S. epidermidis* between days 135 and 144 (AtB-types B and D). Two vanco-locks were then performed through the catheter on day 145, allowing the blood cultures to remain sterile for a week before re-appearance of *S. epidermidis* (AtB-types B and E). The catheter was removed on day 158 and the administration of intravenous vancomycin was discontinued. No *S. epidermidis* was isolated from the culture of the catheter nor from the blood-culture samples. The patient was discharged afebrile.

Bacterial strains and antibiotic susceptibility testing

All strains were identified by the ID-32-Staph gallery (bioMérieux, Marcy-l’Etoile, France). Antimicrobial susceptibility was determined by the ATB-Staph gallery, as recommended by the manufacturer (bioMérieux) (table).

Culture of the catheter tips

The removed catheters were cultured according to the method described by Brun-Buisson et al. Briefly, the catheter tip was immersed in 1 ml of Brain Heart Infusion Broth (bioMérieux) and shaken vigorously for 1 min. Then, 0.1 ml of the suspension was inoculated on to sheep blood agar (bioMérieux) allowing the number of cfu to be counted. Concomitantly, the catheter tip was cultured in the brain heart infusion broth.

Chromosomal analysis by PFGE

Chromosomal DNA was extracted as described previously. Intact DNA was digested with 30 U of *SmaI*, as recommended by the manufacturer (Boehringer Mannheim, Meylan, France). PFGE was performed with a contour-clamped homogeneous electric field apparatus (CHEF-DR II; BioRad, Ivry-sur-Seine, France) at 150 V for 20 h in cooled TBE buffer, with a constant pulse time of 20 s. After the run, gels were stained for 15 min in ethidium bromide solution 10 mg/ml and photographed with a Polaroid camera under UV light.

Results

Three clearly different restriction pulsed patterns were observed (figs. 1 and 2). The first isolate (1) presented its own unique PFGE pattern (PFE-1) while the 41 remaining isolates were gathered into two groups of, respectively, 32 (group PFE-2) and nine isolates (group PFE-3); one blood culture yielded both strains (no. 37 in fig. 2).

Except for isolate no. 1, all isolates obtained before the first catheter was removed belonged to group PFE-2 (figs. 1 and 3). This strain was also isolated from the first catheter. After implantation of the second catheter and before discharge, the first two isolates belonged to group PFE-3 (figs. 1 and 3). Then,
Fig. 2. *SmaI* pulsed-field patterns of isolates nos. 22-42. L, concatemers of phage λ as molecular size marker (kb); RN, *S. aureus* RN4220, used as a control strain. Isolates from blood culture no. 37 displayed a combination of patterns PFE-2 and PFE-3; both strains were isolated from the blood sample.

Fig. 3. Schematic representation of the correlation between AtB-types and PFE-types. AtB-B, C and D isolates shared the same PFE-Type (PFE-2). Blood culture no. 37 yielded both PFE-2 and -3 strains. Arrows indicate catheter removal; G, episodes of acute GVHD; VL, Vanco-locks, ----, discharge.

Table. Correlation between antibiotypes (AtB-types) and DNA restriction pulsed-field patterns (PFE-types) of the 42 *S. epidermidis* isolates

<table>
<thead>
<tr>
<th>Number of isolates</th>
<th>Antimicrobial resistance pattern</th>
<th>AtB-type</th>
<th>PFE-type</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>pen</td>
<td>A</td>
<td>1</td>
</tr>
<tr>
<td>28</td>
<td>pen/met/gen/pef/sxt</td>
<td>B</td>
<td>2</td>
</tr>
<tr>
<td>3</td>
<td>pen/met/gen/ery/rif/pef/sxt/fos/fus</td>
<td>C</td>
<td>2</td>
</tr>
<tr>
<td>1</td>
<td>pen/met/gen/pef/sxt/fos</td>
<td>D</td>
<td>2</td>
</tr>
<tr>
<td>9</td>
<td>pen/met/gen/ery/prt/rif/pef/sxt/fos/fus</td>
<td>E</td>
<td>3</td>
</tr>
</tbody>
</table>

Pen, penicillin; met, methicillin; gen, gentamicin; ery, erythromycin; prrt, pristinamycin; rif, rifampicin; pef, pefloxacin; sxt, sulphanmethoxazole + trimethoprim; fos, fosfomycin; fus, fusidic acid.

On the second admission to hospital, two PFE-2 isolates were obtained, followed by three PFE-3 isolates (figs. 2 and 3). Then PFE-2 isolates persisted until the vanco-locks were performed. Between the vanco-locks and the second catheter removal, PFE-2 and PFE-3 isolates were recovered alternately (figs. 2 and 3). “Isolate no. 37” displayed a pattern combining PFE-2 and PFE-3 patterns; both strains were isolated from the blood culture.

Changes were detected in antibiotic susceptibility of four isolates belonging to group PFE-2 (table). All PFE-3 isolates had the same antibiogram.
Discussion

*S. epidermidis* is recognised as an important nosocomial pathogen in immunocompromised patients. The conventional approach to determine the clinical relevance of *S. epidermidis* isolates in a patient is to require repeated isolation from independent specimens of isolates that share typing characteristics. PFGE has been described as effective and discriminating in strain comparison, since it can provide easy-to-compare and unequivocal DNA restriction patterns.\(^{12-15}\)

In the case described above, *S. epidermidis* isolates from blood cultures yielded three different pulsed field patterns, PFE-1, PFE-2 and PFE-3 (figs. 1 and 2). Type PFE-2 and PFE-3 gathered, respectively, 32 and nine isolates (blood culture no. 37 yielded both strains) with identical *Smal* restriction patterns (figs. 1 and 2). Type PFE-2 isolates were obtained from blood samples before the first catheter was removed, from the first catheter tip, and from blood cultures after the catheter removal. Isolation from the culture of the catheter indicated that it was the portal of entry, but as the strain was isolated from blood cultures 28 days after the catheter removal, it is possible that the catheter was endogenously colonised from another site. One major reservoir of *S. epidermidis* is the gastrointestinal tract.\(^{17}\) The PFE-2 strain could have been present in the intestinal flora and been translocating chronically during the chronic intestinal GVHD developed in the patient. Unfortunately, *S. epidermidis* was not isolated from the faeces because the patient received an oral vancomycin regimen. A second explanation for this late recurrence may have been a possible metastatic localisation of the PFE-2 strain. There was no clinical evidence for any putative metastatic site, and except for the stool surveillance, no investigation was performed to assess its existence (for example, on skin, oral mucosa or catheter hub cultures).

Changes in antimicrobial susceptibility were detected in four PFE-2 isolates (acquisition of resistance to fosfomycin in isolate no. 25 and to erythromycin, rifampicin, fosfomycin and fusidic acid in isolates nos. 7, 20 and 21). These changes lead the clinician to consider those isolates as different. Such switches in antibiotic susceptibility are known to occur in vivo, mostly due to the acquisition of resistance genes mediated by plasmids or transposons from reservoir strains.\(^{18}\) In the four observed variants, comparison of the restriction pulsed-field patterns demonstrated clonality of these variants with the other PFE-2 isolates. This emphasises the need for considerable care when assessing differences in isolates with minor differences in antimicrobial resistance patterns.

The pathogenic role of the second strain (PFE-3) was not clear. This pristinamycin-resistant strain may have emerged during the course of antimicrobial therapy and been responsible for bacteraemia during eradication of PFE-2 (fig. 3). However, as PFE-3 isolates were obtained during three short periods (fig. 3), this strain could also be considered as chronically contaminating blood cultures and not being of clinical relevance. No portal of entry was determined for PFE-3. Nevertheless, as the blood samples were sterile after removal of the second catheter, it is likely that the catheter tip was the source of PFE-3, even if the strain was not isolated from the culture of this catheter. We assume that the vanco-locks were responsible for eradication of this PFE-3 strain from the catheter.

This observation emphasises that the pathogenic role of *S. epidermidis* remains difficult to analyse and that antimicrobial susceptibility patterns may lead clinicians to wrongly define two isolates as different. The use of PFGE will be of help in the understanding of both mechanisms of transmission and pathogenicity of coagulase-negative staphylococci, as well as in the epidemiological analysis of such recurrent bacteraemia.

References


