Molecular analysis of the microflora in chronic venous leg ulceration

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INTRODUCTION

Chronic venous leg ulcers (CVLU) affect approximately 1% of the UK population and are debilitating and painful. The aetiology of these wounds is not well understood, but it is likely to be multi-factorial. CVLU support a diverse microbrial flora, and there is evidence that micro-organisms in this biofilm contribute to the non-healing phenotype. Moreover, these wounds occur in individuals and tissues that are at increased risk of bacterial invasion due to poor vascular supply and systemic factors (Falanga, 1993). Based as it is on an unclear understanding of the aetiology, management is currently unsatisfactory and expensive, with costs of over £1 billion per annum in the UK.

Cultural analyses have documented that staphylococci, streptococci, enterococci and facultative Gram-negative bacilli are the bacterial groups most frequently recovered from CVLU (Hansson et al., 1995). However, more recently, the importance of fastidious and slow-growing species such as anaerobes is being established. With the employment of strict anaerobic isolation techniques and prolonged incubation times, almost 60% of CVLU have been shown to harbour anaerobic bacteria such as Fusobacterium spp. and peptostreptococci (Halbert et al., 1992; Murdoch et al., 1994). However, it is becoming increasingly clear that there are considerable limitations to cultivation-dependent methods in determining the composition of complex microbial communities and that culture alone is unlikely to characterize fully the microflora present in chronic wounds. In a limited number of microbial systems, considerably greater diversity of the microflora has been revealed by molecular analysis, in particular the retrieval of rRNA gene sequences by PCR. In addition, molecular analysis is increasingly being employed as an adjunct to conventional cultural techniques in the analysis of diseases with complex microflora (Wilson et al., 1997). Sequences so retrieved are cloned and sequenced and the sequences compared to sequence databases for identification or phylogenetic characterization. For example, the study of the oral microflora associated with periodontal disease and purulent infection, in which it has been estimated that only 30–50% of microbial species are amenable to culture, has been significantly advanced by the application of such molecular tools (Kroes et al., 1999; Dymock et al., 1996).

In this study, we have applied both molecular and enhanced cultural techniques in the analysis of the microbial population within a single, chronic, non-healing, venous leg ulcer wound.
METHODS

Patient details. Following ethical approval, a typical patient (i.e. elderly, obese, female) at the Wound Healing Research Unit, University Hospital of Wales, Cardiff, receiving standard compression therapy for a clinically non-infected wound, was selected for study with informed consent. Additional patient characteristics include: age, 59; height, 165 cm; weight, 107 kg; duration of wound, 2 years; wound size, 1310 mm². There was no presence of systemic disease and the patient had not had systemic or topical antimicrobial therapy in the previous month.

Sampling of the wound. The wound surface was irrigated with sterile saline and the wound bed sampled via a surface swab and a 6 mm punch biopsy. The swab and tissue were transferred in separate vials of (2 ml) reduced transport medium (TM) (Bowden & Hardie, 1971) to the laboratory for processing within 1 h of collection. The tissue was divided aseptically into two portions using a sterile scalpel, one for cultural analysis and the other for molecular analysis.

Cultural analysis of wound microflora. The swab sample was vortex-mixed in TM for 5 min. The tissue for cultural analysis was cut up finely with a sterile scalpel and also vortex-mixed in TM for 5 min. Serial dilutions in TM were plated onto the following media (from LabM): blood agar (BA), fastidious anaerobe agar (FAA), both supplemented with 5 % (v/v) horse blood, MacConkey no. 3 and Sabouraud’s medium. All plates except FAA were incubated aerobically at 37 °C. FAA plates were incubated anaerobically (10 % CO₂, 10 % H₂, 80 % N₂) at 37 °C. Prolonged incubation of the macerated tissue was undertaken in fastidious anaerobe broth for 7 days prior to plating on FAA to allow recovery of fastidious and slow-growing anaerobic species. Primary isolation plates were initially examined after 48 h and then incubated for at least 10 days. Identification of bacteria followed standard microbiological schemes by examination of a range of phenotypic properties (staining reactions, colonial morphology, carbohydrate fermentation patterns) and, where appropriate, commercial identification kits. In this way, it was possible to identify each isolate to species level.

Molecular analysis of wound microflora. Molecular analysis was done on the tissue sample. DNA was extracted by standard techniques (Kay et al., 1998) with the addition of three freezes–thaw steps (each consisting of 2 min in liquid nitrogen and 2 min at 65 °C). DNA was extracted directly from the tissue sample and also from individual microbial species cultured from the wound. PCR was performed as described previously (Marchesi et al., 1998; Lane, 1991; Carroll et al., 2000; Klaußegger et al., 1999) using three sets of universal primers. These were designated according to Esherichia coli 16S rDNA numbering as follows: 63f/1387r (Marchesi et al., 1998), 27f/1492r and 27f/1525r (Lane, 1991). Two sets of primers designed primarily to amplify Gram-positive bacterial species were also used in a nested PCR using primers 27f/1492r in the first round of PCR followed by a second PCR with primers 712f/1067r (Carroll et al., 2000) and 1185f/1540r (Klaussegger et al., 1999). PCR was followed by ligation into the pCR2.1 Topo vector (Invitrogen) and transformation into Top10 competent E. coli cells (Invitrogen). Blue/white screening of transformants was done on Law–Rosenberg agar (Sambrook et al., 1989) containing 50 mg ampicillin ml⁻¹) and top-spread with 40 ml X-Gal solution (20 mg ml⁻¹). Clones amplified from tissue were screened by RFLP analysis of M13-amplified PCR products using the restriction enzymes CfoI, RsaI and HaeIII (Promega). DNA for sequencing was prepared from clones using Wizard Plus SV minipreps (Promega). Clones with unique RFLP patterns and two clones from each cultured organism were sequenced on an automated laser fluorescence sequencer (ABI 377; PE Applied Biosystems) by using primers M13f and M13r, giving double coverage of > 1 kbp of 16S rDNA for the universal primer products for phylogenetic analysis. Gram-positive primer products were considerably smaller, 355 and 365 bp (Carroll et al., 2000; Klaußegger et al., 1999), for which 100 % double coverage was achieved. Sequences obtained were compared to database sequences and phylogenetic trees were constructed as described previously (Hill et al., 1999).

RESULTS

Cultural analysis revealed that Acinetobacter spp. and Staphylococcus epidermidis were found in large numbers (respectively 6·0 × 10⁴ and 2·0 × 10⁳ c.f.u. ml⁻¹; Table 1) in the wound tissue and that no additional cultural isolates were obtained after prolonged anaerobic incubation of the sample. Distinct differences were observed between sample sites (Table 1). Two species, Staphylococcus epidermidis and Proteus spp., were isolated from the tissue and the swab, respectively, but not from both sample sites, and Acinetobacter spp. were found in large numbers in both the tissue and the swab.

The results of the molecular analysis are shown in the dendrogram in Fig. 1 and in Table 2. In total, 26 clones amplified using five different primer sets were sequenced. It was possible by PCR to amplify sequences from bacteria found in the tissue and, in addition, sequences representing bacteria cultured from the swab only, e.g. Proteus spp., were detected. Fig. 1 shows that the sequences retrieved varied to some extent with the primer pairs used. For example, clones 3_3 and 3_1, showing greatest similarity to Acinetobacter baumannii and Bacteroides ureolyticus, respectively, were only amplified using primers 27f/1525r. Furthermore, other clones obtained using the other two universal primer pairs were not found in this part of the tree. Likewise, Gram-positive species were only amplified when using the Gram-positive-specific primers 712f/1067r and 1185f/1540r in a nested PCR and not when using any of the three universal primer sets (Table 2). The latter sequences were unsuitable for inclusion in the dendrogram due to a lack of overlap between the two primer pairs and also because of their short length (355 and 365 bp), which would have compromised the robustness of the tree.

All of the sequences recovered from bacteria cultivated from the wound tissue were >98 % identical to sequences within the public databases. In contrast, the 16S rRNA sequences recovered by direct amplification from tissue showed a greater degree of variability. For example, clones of the amplified sequences in cluster A, namely clones 0_16, 0_11 and 0_10, showed more than 5 % dissimilarity from any

Table 1. Cultural analysis of swab and tissue samples

<table>
<thead>
<tr>
<th>Species</th>
<th>Swab</th>
<th>Tissue</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acinetobacter spp.</td>
<td>5·48 × 10⁴</td>
<td>6·0 × 10⁴</td>
</tr>
<tr>
<td>Staphylococcus epidermidis</td>
<td>ND</td>
<td>2·0 × 10⁴</td>
</tr>
<tr>
<td>Proteus spp.</td>
<td>3·6 × 10⁴</td>
<td>ND</td>
</tr>
<tr>
<td>Candida tropicalis (yeast)</td>
<td>4</td>
<td>ND</td>
</tr>
</tbody>
</table>

Values are c.f.u. ml⁻¹. ND, Not detected.
known sequence, a level of variation that is often found between members of different genera. None of the cultivated sequences exhibited this degree of sequence dissimilarity. Significantly, three clones, 3_1, 5_63 and 5_69, showed very low similarity (90.2%) to all database sequences (most similar sequences from Bacteroides ureolyticus, Lactosphaera pasteurii and Acinetobacter lwoffi respectively) and were not detected by culture and may therefore represent novel, previously uncultured species (Fig. 1; Table 2). In addition, nine clones in cluster B (Fig. 1) and clone 5_75 (Table 2) appeared to have high similarity, >95% (clones 0_183, 0_8) and >98% (clones 0_4 to 1_1), to a sequence from Morganella morganii ATCC 25830T, again a species that was not cultured from this wound.

**DISCUSSION**

All previous studies to enumerate and identify the microflora of venous leg ulcers have been conducted by bacterial culture alone (Bowler, 1998). This is, to our knowledge, the first such study to use molecular techniques, particularly 16S rRNA sequencing, to identify microorganisms in venous leg ulcers. The study also highlighted the importance of culture-independent methods in microbiology, as many of the sequences were novel and not found in the database.

**Table 2. 16S rRNA sequencing of five clones amplified using Gram-positive-bacteria-specific primers**

<table>
<thead>
<tr>
<th>Clone</th>
<th>Primers</th>
<th>16S rRNA species identification</th>
<th>Sequence identity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>4_38</td>
<td>712f/1067r</td>
<td>Enterococcus faecalis</td>
<td>99-1</td>
</tr>
<tr>
<td>5_60</td>
<td>1185f/1540r</td>
<td>Peptostreptococcus octavius</td>
<td>97</td>
</tr>
<tr>
<td>5_75</td>
<td>1185f/1540r</td>
<td>Morganella morganii</td>
<td>95</td>
</tr>
<tr>
<td>5_69</td>
<td>1185f/1540r</td>
<td>Acinetobacter lwoffi</td>
<td>89-8*</td>
</tr>
<tr>
<td>5_63</td>
<td>1185f/1540r</td>
<td>Lactosphaera pasteurii</td>
<td>89-3*</td>
</tr>
</tbody>
</table>

*Very low identity to database sequences indicates that these clones may represent novel phylotypes.
study applying both a cultural and a molecular approach to chronic wounds. Indeed, few medically important microbial communities have been studied extensively in this way. The microflora associated with the oral cavity, including the human subgingival crevice (Kroes et al., 1999), endodontic infections (Rolph et al., 2001) and childhood caries (Becker et al., 2002), has been studied. Importantly, all of these studies have shown the importance of the molecular approach in revealing greater species diversity. Similarly, our data suggest that cultivation alone underrepresents the true extent of bacterial diversity within the chronic wound and that a more comprehensive analysis is possible with the application of molecular techniques (Hugenholtz & Pace, 1996). A number of clones closely related to cultured microorganisms such as Acinetobacter baumannii and Proteus mirabilis were detected. More importantly, however, clones most closely related to Morganella morganii, the anaerobe Bacteroides ureolyticus and the Gram-positive species Enterococcus faecalis, Peptostreptococcus octavius and Lactocphaera pasteurii were amplified directly from tissue, although these species had not been cultured from this wound.

Proteus spp. were cultured only from the swab, perhaps indicating that it was only a surface colonizer, although, conversely, clones similar to Proteus spp. were detected by molecular means from the tissue. Notably, we have observed little variation in sampling within a biopsy specimen when it was divided into quarters prior to cultural analysis, isolating identical bacterial species and at similar bacterial counts from each biopsy segment (unpublished observations). It is therefore unclear why we were unable to detect Proteus in the tissue by cultural means, although it is possible that cells able to invade the deep tissue may change their phenotype to a semi-dormant state (viable but not culturable), making them difficult to revive by standard culture. Staphylococcus epidermidis was not detected in the swab, presumably as it had colonized the deep tissues and not the wound surface. Surprisingly, Staphylococcus spp. were not amplified by PCR, although clearly present in the tissue from the cultural analysis.

Along with other researchers, we have highlighted the importance of analysis with multiple primer sets in order to obtain the widest spectrum of phylogenetic groups and to reduce the effect of primer biases in such studies (Kroes et al., 1999; Paster et al., 2001). All primers used by us and others have been validated extensively against a wide range of bacteria yet, despite this, biases are apparent. It is possible that the primers behave differently, perhaps annealing less efficiently, when applied in mixed culture. Certainly, in this study, despite the validated universal nature of a number of the primers employed, quite different species were detected depending on the primer set used. Moreover, the Gram-positive-specific primers were shown to be capable of detecting sequences representative of Gram-negative species from this wound. This selective amplification reflects a pitfall in the PCR, where primer annealing of one DNA target can be biased, regardless of the initial proportions of the templates (Suzuki & Giovannoni, 1996). Notwithstanding these biases, however, it is apparent that greater understanding of these biases is needed and that the molecular approach is essential to begin to reveal the true extent of bacterial diversity.

Clones 3_1 and 0_16 to 0_10 (as they appear in the dendrogram in Fig. 1) are most interesting, in that they represent novel phylotypes, a phylotype being defined as a cluster that differs from known species by ~2 % and where members are at least 99 % similar to other members of the same cluster (Paster et al., 2001). Importantly, phylotypes represented only by clones (e.g. clones 3_1, 5_63 and 5_69) may represent currently uncultured or unrecognized potential pathogens. Sequences from the uncultured organisms can be used to design PCR primers for subsequent application to larger numbers of clinical samples, for example via DNA microarrays or using quantitative PCR, hence facilitating the identification of potential markers of disease. Ten clones were found to be most closely related to Morganella morganii (clones 3_75 and 0_16 to 0_10), which is amenable to culture and has previously been detected in chronic wounds (Bowler & Davies, 1999), but was not cultured in this study. Hence, there appears to be a bias in the cultural analysis. It is possible that less numerous or less vigorous species may have been swamped by the predominant ones and, hence, that they were out-competed on the non-selective media used for culture. One species commonly found in venous ulceration, Pseudomonas aeruginosa, was not detected in this patient.

The debate still exists as to whether it is relevant to treat venous leg ulcers routinely with antibiotic (topical and systemic) or anti-septic (topical) therapies. Clearly, the patient in this study had a colonized ulcer, but had no apparent signs of clinical infection, so it is debatable whether they would have benefited in the long term from such treatment. Antibiotic therapy (usually broad-spectrum) is effective in reducing the bacterial burden of such wounds in the short term, i.e. for the duration of the antibiotic application. However, once stopped, the microflora is quickly able to re-establish itself. This means that routine microbiological sampling of such wounds is therefore often pointless. Clearly, appropriate systemic antibiotics are essential for the treatment of deteriorating, clinically infected wounds characterized by severe pain, oedema, erythema and strong odour. In such cases, microbiological sampling of the wound can be an important aid to appropriate antibiotic therapy.

This study demonstrates that molecular techniques can detect the presence of bacteria in chronic wounds when culture techniques yield a negative result and can be used to identify a wider range of chronic wound-related bacteria, including the presence of previously unidentified or unculturable bacteria. The clinical relevance of the organisms detected in this study remains unclear. However, the comprehensive characterization of the microflora in this way is a prerequisite for the elucidation of associations between bacterial carriage, clinical outcome and effective treatment of this important condition.
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REFERENCES


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