Typing of strains of *Staphylococcus aureus* by Western Blot analysis of culture supernates

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**Summary.** Extracellular proteins produced by *Staphylococcus aureus* strains were examined by Western Blot analysis with blood donor plasma as a source of antibodies. Comparison of epidemiologically related strains showed strong concordance between plot pattern and phage type.

**Introduction**

Bacteriophage typing of *Staphylococcus aureus* isolates is a valuable tool in the study of staphylococcal disease in man (Parker, 1983). The occurrence of a significant number of non-typable strains, and uncertainties arising from the knowledge that changes in the susceptibility to bacteriophages may be brought about by lysogenization (Rountree, 1959), make alternative typing methods desirable. Serotyping schemes based on surface antigens have been developed and as many as 30 antigenic factors can be recognised (Oeding, 1978).

Polyacrylamide-gel electrophoresis (PAGE) of bacterial proteins has been used extensively as a taxonomic tool, and, more recently, for typing (Poxton et al., 1984; Tabaqchali et al., 1984). Sacks et al. (1969) used PAGE of cell proteins to classify Enterobacteriaceae; they included *S. aureus* in their study and showed that the polypeptide profiles of this organism were very different from those of the Enterobacteriaceae examined.

In this study we have compared different strains of *S. aureus* by PAGE of whole cell extracts, and by Western blotting of extracellular proteins with human plasma as a source of anti-staphylococcal antibodies.

**Materials and methods**

*S. aureus* strains

*S. aureus* strains were obtained from nasal swabs from a group of men living in complete isolation for a year in Antarctica. Isolates from six subjects were used in this study. They were isolated from swabs taken over a period of c. 40 weeks. The table shows details of the strains. They were identified as *S. aureus* in Antarctica by colonial morphology, Gram's stain and coagulase testing. Cultures were stored on nutrient-agar slopes in 1-ml screw-capped bottles held at 4°C until the cultures were revived at the Staphylococcal Reference Laboratory, Colindale, where phage typing was performed. Fresh storage cultures were prepared as before, and transferred to the Department of Bacteriology, University of Aberdeen for further analysis.

**Preparation of material for electrophoresis**

In Aberdeen the organisms were plated out on nutrient agar plates which were then incubated at 37°C for 24 h. A sweep of colonies from these plates was inoculated into 10 ml of nutrient broth in a 20-ml screw-capped container which was incubated overnight at 37°C with shaking in an orbital incubator. The resulting broth culture was centrifuged at 2500 g for 15 min and the upper 2 ml of the supernate removed for gel analysis. The remainder of the supernate was discarded and a concentrated suspension of the cell pellet placed in a Microfuge (Beckman B) tube and centrifuged for 5 min. The pellet thus produced was washed in an equal volume of distilled water and disrupted for 5 min with an ultrasound probe (Rapidis 150, Ultrasonics Ltd, Shipley, W. Yorks) emitting 45W at maximum power. Protein concentrations were estimated by the method of Bradford (1976).

**Polyacrylamide-gel electrophoresis (PAGE)**

A modified version of the SDS-PAGE technique of Laemmli (1970) was used. Specimens were prepared for PAGE by boiling for 5 min in sodium dodecyl sulphate (SDS) 2% w/v, 2-mercaptoethanol 5% v/v, glycerol 10% v/v and 0.05 M Tris-HCl, pH 6.8. They were applied to wells in a 3-6% acrylamide stacking gel over a separating gel of 10% acrylamide. Gels were run in a Mk 1 Protein Cell (Bio-Rad) at 60 mA, with the buffers described by Laemmli, until the bromophenol blue marker had reached the bottom of the gel; this occurred after c. 660 V-h. Proteins were either stained with Coomassie Blue or transferred to nitrocellulose paper.
Table Phage types of isolates from each subject

<table>
<thead>
<tr>
<th>Subject</th>
<th>Phage typing results</th>
<th>Experimental phages</th>
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<tr>
<td></td>
<td>RTD</td>
<td>100 × RTD</td>
</tr>
<tr>
<td>9</td>
<td>94/96</td>
<td>94/96</td>
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<tr>
<td>12</td>
<td>94/96</td>
<td>94/96</td>
</tr>
<tr>
<td>16</td>
<td>29/52 +</td>
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<tr>
<td>21</td>
<td>NT</td>
<td>83A/85 +</td>
</tr>
<tr>
<td>23</td>
<td>6/47/53/54</td>
<td>6/47/53/54/75/77</td>
</tr>
<tr>
<td>28</td>
<td>NT</td>
<td>29/47/54/75/77/85/81</td>
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Blotting

Transfer of proteins was achieved by the technique of Towbin et al. (1979), with Burnette’s (1981) addition of filter paper to the sandwich and with a Bio-Rad Transblot cell. Transfer was effected overnight at 70 mA, with the buffer described by Burnette. The paper was washed in blocking buffer (phosphate-buffered saline, pH 7.4, containing newborn calf serum 10% v/v) and Triton X-100 0.2% v/v for 30 min, followed by treatment with human plasma diluted 1 in 50 in blocking buffer for 1 h. Preliminary study by Western Blot analysis of whole-cell extracts had shown that plasma from this donor contained antibodies against a large number of staphylococcal polypeptides. The paper was then washed three times for 15 min each in blocking buffer followed by 45 min in a solution of peroxidase-linked goat anti-human serum (Miles-Yeda, Israel) diluted 1 in 500. It was then washed another three times for 15 min each in blocking buffer, rinsed in phosphate-buffered-saline and stained with diaminobenzidine tetrahydrochloride 0.05% w/v in 0.1 M Tris (pH 7.6) containing hydrogen peroxide 0.01% v/v.

Results

Figs. A and B show the results obtained with isolates from six subjects. Fig. A shows a polyacrylamide gel, stained with Coomassie Blue, showing the results obtained with whole-cell extracts. Apart from the absence of a single protein band in tracks N, O and P, all the isolates were virtually indistinguishable. Fig. 1B shows a Western Blot of supernate from the same isolates in the same order as fig. A. The isolates from five individuals can be grouped according to their pattern of bands, isolates from an individual resembled each other closely, and isolates from different individuals were easily distinguished from each other. The only exception to this was with the isolates from subjects 9 and 12 which were indistinguishable. Isolates were grouped after visual inspection of Western-Blot band patterns and on the basis of (i) possession of bands unique to a group, (ii) band intensity, and (iii) the presence or absence of bands common to several groups. Comparison of grouping by Western-Blot band pattern and phage-typing results showed complete concordance (table); isolates from subjects 16, 21, 23 and 28 were distinguishable from

Figure (A) Polyacrylamide-gel electropherogram of whole-cell extracts, stained with Coomassie Blue. (B) Western Blot analysis of culture supernatants from the same strains. Track A, isolate from subject 9; tracks B–E, isolates from subject 12; tracks F–I isolates from subject 16; tracks J–M, isolates from subject 21; tracks N–P, isolates from subject 23, tracks Q–T, isolates from subject 28. Numbers refer to the week on which the isolate was obtained.
each other and from the isolates from subjects 9 and 12, whereas strains from the latter pair of subjects could not be distinguished from each other either by Western-blot-band pattern or with the phages in the International Basic Set. The pattern produced by a particular strain isolated repeatedly from a subject was constant over a period of several months and did not change after 10 subcultures in the laboratory (results not shown).

Discussion

Each of the six subjects in this study carried a *S. aureus* of different phage type throughout the year. The phage types are shown in the table. Because the subjects were in total isolation and under intensive microbiological surveillance throughout the period during which the samples were obtained it is felt that strains obtained were particularly suitable epidemiologically for a study of this nature. The Western Blots clearly show that isolates from a single individual produced very similar bands, whereas, with the exception of the strains from subjects 9 and 12, isolates from different individuals produced different patterns. As a typing technique, this method appears to be comparable in reproducibility to phage typing and it is interesting to note the similarity between the two strains of phage-group V (subjects 9 and 12).

The whole-cell-protein profiles in Coomassie Blue-stained gels were very similar and the majority of isolates were virtually indistinguishable.

This study was limited to strains from healthy individuals without clinical manifestations of staphylococcal infection during the study. We plan to study the extracellular products of strains associated with clinical infection.

Analysis by Western blotting was chosen because it allows detection of products present in amounts too small to be detected by conventional staining methods and because the identification and characterisation of extracellular products which elicit antibodies in human subjects may be of value in furthering an understanding of the pathogenesis of staphylococcal infections.

By obtaining a time-expired donation of plasma from the Blood Transfusion Service, sufficient material for many typing studies was ensured, particularly in view of the 1 in 50 working dilution used. Further studies are in progress in which Western-blot patterns shown by other human sera are being compared to allow an assessment of the general usefulness of such an antibody source for typing staphylococci with this method.

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REFERENCES


