Evidence for Two Mechanisms of Plasmid Transfer in Mixed Cultures of *Staphylococcus aureus*

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Non-lysogenic *Staphylococcus aureus* strain 1030 was lysogenized with 12 different bacteriophages. Lysogeny was associated with acquisition of phage inducibility by mitomycin C treatment or ultraviolet irradiation, with the presence of plaque-forming phage in culture supernatants and with considerable narrowing in susceptibility to the typing bacteriophages and also with increased sensitivity to trimethoprim and sulphadiazine. The presence of prophages in the donor and/or the recipient could either promote or inhibit transfer of plasmids between mixed cultures. Transfer frequencies in mixed culture after 18 h incubation could be as high as $7.0 \times 10^{-1}$ resistant recipients/final donor, and evidence was adduced for a mechanism distinct from transfer by spontaneous transduction. It is suggested that this method of gene transfer be described as ‘phage-mediated conjugation’. Chromosomal genes were not transferred by this method. Two similar plasmids cp-2 and cp-3 were able to promote their own transfer through clones of 1030; plasmids coding for resistance to either neomycin or tetracycline could be transferred to a recipient by the presence of cp-2 or cp-3 simultaneously in the donor. The presence of plasmids in 1030 was associated with a small increase in sensitivity to trimethoprim or sulphadiazine.

**INTRODUCTION**

Bacteriophages extracted from cultures of *Staphylococcus aureus* are well known to mediate generalized transduction within this species (Pattee & Baldwin, 1961; Novick, 1963). The possibility that antibiotic resistance in *S. aureus* might spread between isolates in nature by transduction was strengthened by the construction of an element that contained DNA of both plasmid and phage origin (Novick, 1967). This element could pass spontaneously between cells in mixed cultures *in vitro* at frequencies of about $10^{-4}$, and at rather lower frequencies following inoculation of the cultures into mice (Novick & Morse, 1967). However, a specific helper phage in the donor is necessary for the production of transferring particles, and the role that such an element can play in the spread of antibiotic resistance in staphylococci in general is therefore uncertain. Spread of resistance by generalized transduction in nature has also been considered unlikely to occur commonly because of the low frequencies attainable *in vitro*, typically $10^{-6}$ to $10^{-8}$ (Richmond, 1969). However, resistance to several antibiotics can be transferred spontaneously from unmanipulated cultures to appropriate recipients *in vitro*, and experimentally on the skin surface (Lacey, 1971a, b, 1975). This transfer requires between 0.001 and 0.01 M-Ca²⁺ or Mg²⁺, is abolished by sodium citrate and requires the use of a phage-resistant recipient; however, the frequency increases with high cell density, and transferring particles may not be detectable in the donor culture supernatant (Lacey, 1971a; Lacey & Stokes, 1979). Whilst such transfer might be mediated via generalized transduction, with the phages being either cell-bound or...
translated recipient strain, whilst a cell-free extract containing ‘transducing’ phages cannot mediate transfer to such a culture. (2) Lantos (1977) has shown that a single resistance determinant is transferred from a donor to a recipient by conventional transduction, but following incubation of whole cells of the donor with the recipient, three resistance determinants are transferred simultaneously. (3) Naidoo & Noble (1978) have shown that gentamicin resistance can be transferred between strains when they were inoculated on to the skin of human volunteers or mice; however, transfer of gentamicin resistance between these cultures was not detected when they were cultured in nutrient broth. (4) The frequency of transfer of drug resistance between strains in mixed culture may increase when the recipient (1030) is lysogenic for a prophage (Lacey, 1979).

The role of transformation in mediating gene transfer between staphylococci in vivo is not clear because to obtain the optimal frequency of transformation a concentration of 0-1 m-Ca²⁺ is required and extracellular DNAase must be inhibited. Bacteriophage gene(s) in the recipient appear to be essential for transformation (Lindberg & Novick, 1973).

Thus, bacteriophages appear to be closely involved in gene transfer in S. aureus in vitro, although their precise role(s) is uncertain in vivo. This paper describes experiments which show that bacteriophage carriage in S. aureus 1030 can either promote or inhibit plasmid transfer.

**METHODS**

Media. Nutrient broth (Oxoid no. 2) contained additional 0-01 m-CaCl₂ unless otherwise stated. Nutrient agar was obtained from Mast Laboratories, Liverpool, and diagnostic sensitivity test (DST) agar from Oxoid. Milk agar was obtained by the addition of ‘Ideal’ milk to Mast agar to a final concentration of 30 % (v/v).

Antibiotic sensitivity. Sensitivity was determined by the method of Lacey (1979) except that sulphadiazine and trimethoprim were incorporated into DST agar at the following concentrations: 0-05, 0-075, 0-10, 0-15, 0-20, 0-25, 0-30, 0-35, 0-4, 0-5, 1-0 mg trimethoprim l⁻¹; 5, 7-5, 10, 20, 25, 30, 40, 50 mg sulphadiazine l⁻¹.

Bacteriophage typing, mitomycin C induction, ultraviolet light induction, and transduction. These methods have been described previously (Lacey, 1971a, 1979). The optimal mitomycin C concentration for induction of lysogenic derivatives of strain 1030 was 1-0 mg l⁻¹. Accordingly, to test for inducibility of derivatives of strain 1030, they were treated with either 1-0 or 3-0 mg mitomycin C l⁻¹. Induction with either mitomycin C or ultraviolet light was assessed as clearing relative to that of untreated cultures: + + denoted complete clearing, + a reduction in turbidity to about 0-2, and ± a reduction in turbidity to between 0-2 and 0-5. During transduction, care was taken to protect recipients from becoming lysogenic by using a phage to cell ratio of 0-01, and after allowing 15 min for phage adsorption in the presence of 0-01 m-calcium chloride, all subsequent procedures were done in the presence of 0-02 m-sodium citrate. Phages were titrated on culture 1030, unless otherwise stated. Transduction frequencies were expressed as the ratio of the number of transductants to the number of cells in the donor culture at the time of addition of mitomycin C. Phages were checked for sterility following filtration through 0-45 μm Millipore filters and titrated on strain 1030 wild.

*Staphylococcus aureus strain*. Strain 1030 was obtained in 1970 from Dr R. P. Novick, New York, and maintained on nutrient agar slopes at room temperature for 8 years with one subculture. This culture is described as non-lysogenic (Novick, 1967), is lysed at varying efficiencies by each of the currently used typing phages and is non-inducible with ultraviolet light or mitomycin C. Strain 1030 was lysogenized by several typing phages and one of the prophages harboured by strain 13136 (Lacey, 1972). The latter is designated C. Lysogens were studied for sensitivity to trimethoprim and sulphadiazine because thymine (or thymidine) starvation invoked by these agents results in induction of lysogenic cultures (Melechen, 1962; Devoret & Blanco, 1970). Filtrates of this culture that had been treated with 1-0 or 3-0 mg mitomycin C l⁻¹, together with a lysogenic culture derived from this [1030(C)], were examined under the electron microscope by Evelyn Lewis and Dr P. Bennett by the method of Robinson et al. (1977). The 1030(C) lysozyme contained phage particles, but neither preparation of 1030 wild strain contained either intact phage particles or structures resembling phage components. Filtrates of culture 1030 were also studied for plaque-forming activity on 50 freshly isolated clinical strains of *S. aureus*. No plaques were seen. It is inferred that culture 1030 is extremely unlikely to carry complete prophages. Since culture 1030 was to be manipulated, a chromosomal mutant (resistant to 100 mg rifampicin l⁻¹) was selected initially; all derivatives of 1030 also showed a
Table 1. Staphylococcal plasmids

<table>
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<tr>
<th>Plasmid</th>
<th>Resistance(s) coded for</th>
<th>Reference</th>
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<tbody>
<tr>
<td>tet-r</td>
<td>Tetracycline</td>
<td>Lacey &amp; Grinsted (1973)</td>
</tr>
<tr>
<td>neo-r</td>
<td>Neomycin</td>
<td>Chopra et al. (1973)</td>
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<tr>
<td>chl-r</td>
<td>Chloramphenicol</td>
<td>Chopra et al. (1973)</td>
</tr>
<tr>
<td>pen-r</td>
<td>Penicillin, cadmium, arsenate and mercury</td>
<td>Lacey &amp; Grinsted (1973)</td>
</tr>
<tr>
<td>cad-r</td>
<td>Cadmium, arsenate and mercury</td>
<td>Chopra et al. (1973)</td>
</tr>
<tr>
<td>cpns</td>
<td>Penicillin, cadmium, neomycin, streptomycin</td>
<td>Lacey (1979)</td>
</tr>
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<td>cp-1</td>
<td>Penicillin and cadmium</td>
<td>Lacey (1979)</td>
</tr>
<tr>
<td>cp-2</td>
<td>Penicillin and cadmium</td>
<td>Lacey (1979)</td>
</tr>
<tr>
<td>cp-3</td>
<td>Penicillin and cadmium</td>
<td>Lacey (1979)</td>
</tr>
<tr>
<td>sn</td>
<td>Streptomycin and neomycin</td>
<td>Lacey (1979)</td>
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</table>

characteristic very pale buff pigmentation on milk agar and produced deoxyribonuclease (tested on Oxoid DNAase agar).

**Plasmids.** Except for plasmid cpns and four derivatives, cp-1, cp-2, cp-3 and sn (Lacey, 1979), the plasmids used have been fully characterized (Table 1).

**Nomenclature.** Prophage carriage is denoted by the corresponding arabic number in parenthesis, genes of plasmid origin in italics, and mutants selected for chromosomal resistance by capital letters (S, streptomycin; F, fusidic acid). All cultures were rifampicin-resistant; this is not specified for each derivative. Culture 1030(55)tet-r is 1030 lysogenic for typing phage 55 and contains a plasmid determining tetracycline resistance, and 1030(C)S is lysogenic for phage C (from strain 13136) and possesses chromosomal resistance to streptomycin (100 mg l⁻¹).

**Transfer of resistance in mixed cultures.** This was done by the method of Lacey (1971a); modifications are described where appropriate. Unless otherwise stated, the frequency of transfer is defined as the ratio of the number of resistant recipients to the final number of donors after 18 h incubation. From the detailed study of transfer of tet-r to and from clones of 1030(55) (see below) it is assumed that the transfer frequencies represent predominantly transfer from the putative donor to recipient.

**Resistance transfer.** This was detected by the incorporation of either 4 mg tetracycline l⁻¹, 10⁻⁴ M-cadmium acetate, 5 mg neomycin l⁻¹ or 25 mg chloramphenicol base l⁻¹ with either 100 or 400 mg streptomycin l⁻¹ or 10 mg fusidic acid l⁻¹. Coincidental transfer of two resistance markers was studied using nutrient agar containing agents at 70% of these concentrations or by replica-plating of colonies selected for acquisition of a single resistance.

**Storage of cultures.** Purified clones of each derivative were stored on the surface of nutrient agar plates at 4°C for up to 2 months when each was subcultured.

**Transfer experiments under anaerobic conditions.** Broth cultures were mixed on the surface of nutrient agar containing 0.01 M-Ca²⁺ that had been cooled to 4°C and were incubated in a Baird & Tatlock jar in an atmosphere of 10% CO₂ and 90% H₂. Controls included known obligate aerobic and anaerobic cultures in each jar. Some of the plates inoculated with the two cultures were sampled after 1 h incubation to exclude the possibility that transfer had occurred during the seeding of the plates. After incubation for 20 h at 37°C, the plates were withdrawn from the jar and flooded with 2:0 ml 0:05 M-sodium citrate, and bacteria were emulsified and diluted in water containing 0:02 M-sodium citrate, before viable counts were performed on appropriate media also containing 0:02 M-sodium citrate.

**Chemicals.** Bovine pancreatic deoxyribonuclease I was added to cultures at 1 to 100 mg l⁻¹, bovine spleen deoxyribonuclease II at 10 mg to 1 g l⁻¹, and micrococcal nuclease at 0:1 to 10 mg l⁻¹. These enzymes and mitomycin C were obtained from Sigma.

**RESULTS**

**Effect of plasmid carriage on the properties of strain 1030**

Plasmid genes were transduced individually into strain 1030 wild using mitomycin C-induced lysates of donor cultures; during the construction of multiresistant clones of strain 1030 by successive transduction there was no evidence for incompatibility between the plasmids. All the plasmid-containing derivatives had the same bacteriophage typing pattern as that of strain 1030 wild, and all except 1030cp-2, and 1030cp-3 were non-inducible with mitomycin C or ultraviolet light. However, broth culture filtrates of 1030cp-2 and 1030cp-3...
Table 2. Sensitivity of strain 1030 and derivatives to sulphadiazine and trimethoprim

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<tr>
<th>Strain</th>
<th>Sulphadiazine</th>
<th>Trimethoprim</th>
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<tbody>
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</tr>
<tr>
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<tr>
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<td>1030(88)</td>
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<td>0.15</td>
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<tr>
<td>1030(96)</td>
<td>15</td>
<td>0.25</td>
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* Similar results were obtained on at least three occasions.

did not contain any plaque-forming units (p.f.u.), and neither did the other derivatives. Thus, except for culture 1030cp-2 and 1030cp-3 (and see below), there was no evidence for lysogeny in these cultures. However, all were reproducibly slightly more sensitive to trimethoprim and sulphadiazine than culture 1030 wild (Table 2).

**Effect of lysogenization on the properties of strain 1030**

Twelve lysogenic derivatives of strain 1030 were established; these were separately lysogenic for phage C (from strain 13136) and typing phages 52, 79, 81, 55, 71, 95, 42E, 54, 77, 88 and 96. Each lysogen showed a reduction in susceptibility to several typing phages (Table 3) and increased sensitivity to trimethoprim and sulphadiazine (Table 2). All except strain 1030(52) were fully inducible [i.e. cultures completely cleared and contained >10^5 p.f.u. ml^{-1}] with both mitomycin C and ultraviolet light induction. Non-inducible cultures contained <10^4 p.f.u. ml^{-1}. Culture 1030(52) cleared slightly (±) after mitomycin C treatment, but not after treatment with ultraviolet light. However, the culture supernatants of purified 1030(52) contained plaque-forming agents; lysogeny in this derivative is therefore established. A proportion (17%) of 1030(55), 1030(88) and 1030(52) that had been selected with low concentrations of trimethoprim (0.25 to 0.30 mg l^{-1}) had regained the complete typing pattern of 1030 wild. It is inferred that exposure of lysogenic cultures to levels of trimethoprim just above those that are inhibitory to the majority of the culture selects cells that are non-lysogenic.

**Transduction of tetracycline resistance and penicillinase production from strain 13136 to 1030**

A mitomycin C-induced lysate of 13136 was used to transduce both tetracycline resistance and penicillinase production to culture 1030 wild and its lysogenic derivatives. Transduction was successful to each recipient except 1030(52). Replica-plating of at least 50 transductants obtained from each recipient showed no cotransduction of pen^-r and tet^-r. Mixed culture
### Table 3. Reaction of strain 1030 and derivatives to typing phages

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<th>79</th>
<th>80</th>
<th>3A</th>
<th>3C</th>
<th>55</th>
<th>71</th>
<th>95</th>
<th>42E</th>
<th>47</th>
<th>53</th>
<th>54</th>
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</table>

* ++ denotes > 50 plaques, + 20 to 50 plaques and ± < 20 plaques at routine test dilution; – denotes no reaction.
incubation also showed transfer of either pen-r or tet-r separately to each recipient except to 1030(52)S (Table 4). Failure to transfer pen-r or tet-r to 1030(52) might be due to failure of phage adsorption to this strain since this culture is resistant to lysis by phage C. To test for this, 10^6 p.f.u. ml^-1 of phages C, 29, 55 and 77 were incubated with 1030 wild and 1030(52) for 60 min and then the cultures were filtered. Culture 1030 wild adsorbed more than 99% of each phage; 1030(52) adsorbed phage 77 by more than 99%, but failed to adsorb phages C, 29 and 55 (the titre reducing by less than 10% in each case). Possibly lysogeny with phage 52 prevents phage adsorption, although it could be that 1030(52) is both lysogenic and a phage-resistant mutant.

Thus, the properties of transfer of pen-r and tet-r from strain 13136 to 1030 by transduction and mixed culture incubation were similar – the mechanism of mixed culture transfer between 13136 and 1030 is therefore probably by spontaneous transduction. Consistent with this was the observation that when the transductants, obtained after mixed culture incubation, were subcultured to nutrient agar containing 0.01 M-CaCl₂, spontaneous phage activity was seen in seven resistant recipients obtained by either method. In contrast, spontaneous phage activity was uncommon following transfer between donors and recipients both derived from 1030 (see below). Culture filtrates of 13136 were also able to transfer tet-r [1·5 x 10^4 resistant recipients (ml filtrate)^-1] and pen-r [1·0 x 10^2 l^-1] to 1030 wild.

These findings also show that except for 1030(52), which was not evaluated, prophage carriage does not affect the establishment of either the pen-r or tet-r plasmids individually.

Gene transfer mediated by the addition of typing phages in mixed culture experiment

The prospective donor, 1030tet-r, appears to be non-lysogenic and did not transfer tetracycline resistance to recipient 1030S in mixed broth cultures. Mixed cultures were then prepared, with appropriate controls, containing 10^7 each of donor and recipient cells ml^-1 and 10^6 p.f.u. ml^-1 of one of the typing phages or phage C. After 24 h incubation, the mixtures were diluted and inoculated on to antibiotic-containing nutrient agar plates to
Plasmid transfer in *S. aureus*

Table 5. Transfer of resistance between cultures 1030tet-r and 1030S mediated by bacteriophages

Note that the direction of transfer is not established.

<table>
<thead>
<tr>
<th>Phage added</th>
<th>No. of cells ml⁻¹ having phenotype 1030Stet-r</th>
<th>Phage added</th>
<th>No. of cells ml⁻¹ having phenotype 1030Stet-r</th>
</tr>
</thead>
<tbody>
<tr>
<td>29</td>
<td>8.4 x 10⁴</td>
<td>47</td>
<td>0</td>
</tr>
<tr>
<td>52</td>
<td>0 *</td>
<td>53</td>
<td>0</td>
</tr>
<tr>
<td>52A</td>
<td>3.0 x 10⁵</td>
<td>54</td>
<td>0</td>
</tr>
<tr>
<td>79</td>
<td>2.0 x 10⁵</td>
<td>75</td>
<td>0</td>
</tr>
<tr>
<td>80</td>
<td>0</td>
<td>77</td>
<td>0</td>
</tr>
<tr>
<td>3A</td>
<td>0</td>
<td>83A</td>
<td>5.4 x 10²</td>
</tr>
<tr>
<td>3C</td>
<td>0</td>
<td>84</td>
<td>0</td>
</tr>
<tr>
<td>55</td>
<td>7.2 x 10³</td>
<td>85</td>
<td>5.9 x 10⁸</td>
</tr>
<tr>
<td>71</td>
<td>7.5 x 10⁴</td>
<td>88</td>
<td>3.0 x 10⁴</td>
</tr>
<tr>
<td>95</td>
<td>4.2 x 10⁴</td>
<td>81</td>
<td>0</td>
</tr>
<tr>
<td>6</td>
<td>0</td>
<td>94</td>
<td>0</td>
</tr>
<tr>
<td>42E</td>
<td>0</td>
<td>96</td>
<td>8.6 x 10⁴</td>
</tr>
</tbody>
</table>

* < 10¹ ml⁻¹; the mutation frequency of 1030tet-r to 1030Stet-r was always < 10⁻⁹, i.e. there were always < 10⁶ such derivatives in control cultures.

Select for transfer of resistance. Derivatives resistant to both tetracycline and streptomycin were obtained from cultures to which the following phages had been added: 29, 52A, 79, 55, 71, 95, 83A, 85, 88 and 96 (Table 5). The frequency of acquired resistance/total bacteria varied from about 10⁻³ to 10⁻⁶. These results indicate that many phages can be involved in gene transfer, although it is not possible to ascertain the mechanism from these experiments.

Effect of prophage carriage in donor and recipient on transfer of tetracycline resistance in mixed culture

Culture 1030 and nine lysogenic derivatives each containing a plasmid coding for tetracycline resistance were incubated with each of the 13 recipients, 12 being lysogenic. Transfer of resistance occurred in most mixtures, but at variable frequencies. Typical results are shown in Table 6. The presence of phage C or 52 in the recipient was usually associated with a low frequency of transfer (about 10⁻⁷) whilst little transfer was seen when 1030(42E)-tet-r was incubated with each recipient. Although the direction of the plasmid transfer was not absolutely certain, these results suggest that prophage in either the donor or the recipient may permit transfer.

The effect of lysogeny in the donor was first studied in more detail. Donor cultures 1030tet-r, 1030neo-r, 1030chl-r and nine cultures of each lysogenic derivative of 1030 were incubated singly with the non-lysogenic recipient (1030S). As expected, no transfer occurred from 1030tet-r, 1030neo-r or 1030chl-r to non-lysogenic 1030S. Following incubation of the lysogenic donors with the putative recipients, plasmid transfer varied from about 10⁻⁸, for the putative donors lysogenic for phage 42E, to 10⁻³, for cultures lysogenic for phage 95. Examination of a proportion of the resistant recipients obtained after mixed culture incubation revealed that all still gave the intact typing pattern of 1030 wild and their culture filtrates did not produce plaques on culture 1030 wild. Thus, there was no evidence for plasmid transfer mediated by first the passage of the phage from the donor to the recipient, followed by transfer of the plasmid coding for antibiotic resistance from the donor to a lysogenic recipient. It is probable, however, that the direction of plasmid transfer was from the putative donors to the recipients.

Transfer of tetracycline resistance from strain 1030tet-r to 1030(55)SF

The effect of lysogeny in the recipient 1030(55) was studied in detail since it had apparently been found to acquire resistance at high frequency (Table 6). To establish that the direction
Table 6. Transfer of tetracycline resistance between derivatives of strain 1030 in mixed culture

<table>
<thead>
<tr>
<th>Putative donor</th>
<th>1030S</th>
<th>1030(C)S</th>
<th>1030(52)S</th>
<th>1030(79)S</th>
<th>1030(55)S</th>
<th>1030(71)S</th>
<th>1030(95)S</th>
</tr>
</thead>
<tbody>
<tr>
<td>1030tet-r (non-lysogenic)</td>
<td>0†</td>
<td>0</td>
<td>0</td>
<td>3.3 x 10^-2</td>
<td>5.6 x 10^-4</td>
<td>1.9 x 10^-2</td>
<td>2.6 x 10^-4</td>
</tr>
<tr>
<td>1030(C)tet-r</td>
<td>4.8 x 10^-8</td>
<td>4.8 x 10^-4</td>
<td>2.2 x 10^-5</td>
<td>5.5 x 10^-5</td>
<td>8.9 x 10^-4</td>
<td>1.9 x 10^-2</td>
<td>8.5 x 10^-4</td>
</tr>
<tr>
<td>1030(71)tet-r</td>
<td>4.0 x 10^-8</td>
<td>3.3 x 10^-8</td>
<td>2.9 x 10^-5</td>
<td>5.9 x 10^-5</td>
<td>7.2 x 10^-5</td>
<td>1.5 x 10^-4</td>
<td>9.2 x 10^-4</td>
</tr>
<tr>
<td>1030(55)tet-r</td>
<td>3.5 x 10^-6</td>
<td>5.4 x 10^-5</td>
<td>5.5 x 10^-5</td>
<td>8.9 x 10^-5</td>
<td>1.1 x 10^-2</td>
<td>1.5 x 10^-4</td>
<td>7.3 x 10^-4</td>
</tr>
<tr>
<td>1030(95)tet-r</td>
<td>1.5 x 10^-7</td>
<td>1.8 x 10^-7</td>
<td>3.2 x 10^-5</td>
<td>8.7 x 10^-5</td>
<td>1.4 x 10^-2</td>
<td>3.2 x 10^-4</td>
<td>2.6 x 10^-2</td>
</tr>
<tr>
<td>1030(42E)tet-r</td>
<td>5.0 x 10^-8</td>
<td>0.0</td>
<td>3.3 x 10^-5</td>
<td>7.2 x 10^-5</td>
<td>4.8 x 10^-5</td>
<td>1.1 x 10^-3</td>
<td>2.8 x 10^-4</td>
</tr>
<tr>
<td>1030(54)tet-r</td>
<td>5.6 x 10^-2</td>
<td>6.0 x 10^-4</td>
<td>8.3 x 10^-4</td>
<td>7.0 x 10^-5</td>
<td>1.0 x 10^-4</td>
<td>7.2 x 10^-5</td>
<td>5.3 x 10^-3</td>
</tr>
<tr>
<td>1030(77)tet-r</td>
<td>0.0</td>
<td>1.8 x 10^-7</td>
<td>0.0</td>
<td>6.3 x 10^-4</td>
<td>2.6 x 10^-3</td>
<td>1.9 x 10^-5</td>
<td>9.1 x 10^-3</td>
</tr>
<tr>
<td>1030(88)tet-r</td>
<td>4.0 x 10^-8</td>
<td>3.3 x 10^-8</td>
<td>5.5 x 10^-8</td>
<td>5.5 x 10^-3</td>
<td>5.5 x 10^-3</td>
<td>5.5 x 10^-3</td>
<td>4.5 x 10^-3</td>
</tr>
<tr>
<td>1030(96)tet-r</td>
<td>2.1 x 10^-5</td>
<td>2.1 x 10^-5</td>
<td>1.0 x 10^-5</td>
<td>2.9 x 10^-5</td>
<td>7.2 x 10^-5</td>
<td>2.8 x 10^-4</td>
<td>1.8 x 10^-5</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Putative recipient</th>
<th>1030S</th>
<th>1030(C)S</th>
<th>1030(52)S</th>
<th>1030(79)S</th>
<th>1030(55)S</th>
<th>1030(71)S</th>
<th>1030(95)S</th>
</tr>
</thead>
<tbody>
<tr>
<td>1030tet-r (non-lysogenic)</td>
<td>1.6 x 10^-2</td>
<td>3.9 x 10^-3</td>
<td>4.0 x 10^-3</td>
<td>2.0 x 10^-5</td>
<td>2.5 x 10^-4</td>
<td>3.2 x 10^-3</td>
<td>4.5 x 10^-5</td>
</tr>
<tr>
<td>1030(C)tet-r</td>
<td>1.8 x 10^-3</td>
<td>3.5 x 10^-3</td>
<td>6.5 x 10^-3</td>
<td>1.2 x 10^-5</td>
<td>5.0 x 10^-4</td>
<td>1.5 x 10^-4</td>
<td>2.5 x 10^-3</td>
</tr>
<tr>
<td>1030(71)tet-r</td>
<td>8.8 x 10^-7</td>
<td>6.1 x 10^-5</td>
<td>5.5 x 10^-5</td>
<td>2.0 x 10^-5</td>
<td>3.0 x 10^-5</td>
<td>4.3 x 10^-5</td>
<td>4.3 x 10^-5</td>
</tr>
<tr>
<td>1030(55)tet-r</td>
<td>7.4 x 10^-5</td>
<td>1.1 x 10^-5</td>
<td>2.9 x 10^-5</td>
<td>6.0 x 10^-4</td>
<td>5.0 x 10^-5</td>
<td>1.1 x 10^-5</td>
<td>1.1 x 10^-5</td>
</tr>
<tr>
<td>1030(95)tet-r</td>
<td>6.3 x 10^-5</td>
<td>1.8 x 10^-5</td>
<td>2.1 x 10^-5</td>
<td>3.1 x 10^-4</td>
<td>8.9 x 10^-5</td>
<td>3.5 x 10^-5</td>
<td>3.5 x 10^-5</td>
</tr>
<tr>
<td>1030(42E)tet-r</td>
<td>2.9 x 10^-7</td>
<td>7.0 x 10^-7</td>
<td>5.0 x 10^-4</td>
<td>2.9 x 10^-5</td>
<td>5.3 x 10^-5</td>
<td>1.8 x 10^-6</td>
<td>1.8 x 10^-6</td>
</tr>
<tr>
<td>1030(54)tet-r</td>
<td>3.5 x 10^-2</td>
<td>5.7 x 10^-2</td>
<td>4.8 x 10^-2</td>
<td>7.6 x 10^-5</td>
<td>1.1 x 10^-5</td>
<td>2.9 x 10^-2</td>
<td>2.9 x 10^-2</td>
</tr>
<tr>
<td>1030(77)tet-r</td>
<td>1.5 x 10^-3</td>
<td>2.1 x 10^-3</td>
<td>9.5 x 10^-5</td>
<td>2.2 x 10^-4</td>
<td>1.1 x 10^-5</td>
<td>3.2 x 10^-4</td>
<td>3.2 x 10^-4</td>
</tr>
<tr>
<td>1030(88)tet-r</td>
<td>8.9 x 10^-7</td>
<td>6.1 x 10^-7</td>
<td>3.2 x 10^-6</td>
<td>1.3 x 10^-5</td>
<td>0.0</td>
<td>3.1 x 10^-4</td>
<td>2.7 x 10^-4</td>
</tr>
<tr>
<td>1030(96)tet-r</td>
<td>4.5 x 10^-3</td>
<td>5.3 x 10^-4</td>
<td>7.5 x 10^-3</td>
<td>3.8 x 10^-5</td>
<td>1.6 x 10^-5</td>
<td>1.6 x 10^-5</td>
<td>1.6 x 10^-5</td>
</tr>
</tbody>
</table>

* No. of cells resistant to both tetracycline and streptomycin/final no. of cells resistant to tetracycline.
† < 10^-9; the spontaneous frequency of mutation to tetracycline or streptomycin resistance was always < 10^-9.
Table 7. Effect of carriage of phage 55 on the transfer of tetracycline resistance
between clones of 1030

<table>
<thead>
<tr>
<th>Donor</th>
<th>Recipient</th>
<th>Transfer frequency* (duplicate expts)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1030tet-r</td>
<td>1030SF</td>
<td>0</td>
</tr>
<tr>
<td>1030tet-r</td>
<td>1030(55)SF</td>
<td>3.0 × 10⁻¹</td>
</tr>
<tr>
<td>1030(55)tet-r</td>
<td>1030SF</td>
<td>2.5 × 10⁻⁶</td>
</tr>
<tr>
<td>1030(55)tet-r</td>
<td>1030(55)SF</td>
<td>6.0 × 10⁻⁵</td>
</tr>
</tbody>
</table>

* No. of resistant recipients/final no. of donors.

of plasmid transfer was from the non-lysogenic culture 1030tet-r to culture 1030(55)S, a mutant of the latter was isolated with an additional chromosomal marker (resistance to fusidic acid, F). The loci of these two markers were widely separated since there was no cotransduction of the resistances, using phage 88 as vector (data not shown). Cultures of 1030tet-r and 1030(55)SF were mixed in mid-exponential phase and, with controls, sampled at 5, 10, 15, 20 and 30 min and 1, 2, 4, 6 and 20 h by plating undiluted and diluted cultures on to nutrient agar containing 4.0 mg tetracycline l⁻¹, 100 mg streptomycin l⁻¹ and 0.02 M-sodium citrate. Transfer of resistance was first detected after 20 min co-incubation. Forty of these derivatives were examined in detail; all possessed the phenotype 1030(55)SFtet-r including the phage typing pattern of 1030(55) and were lysogenic for a phage resembling 55 in its spectrum of lysis. The rapid appearance of these cells argued that transfer was not mediated by phage 55 released from the recipient and propagated on the donor before conveyance of the tet-r plasmid back to the recipient. Filtrates of broth cultures of 1030(55)SF contained phages lytic for 1030tet-r although little spontaneous phage activity was seen after application of the mixed culture to the surface of nutrient agar containing 0.01 M-CaCl₂. Furthermore, filtrates of the mixed culture after 4 h incubation could not transfer tetracycline resistance to 1030 wild. It is inferred that transfer of tetracycline resistance is in the direction from 1030tet-r to 1030(55)SF. The frequency (expressed as resistant recipients/donors) increased from about 10⁻⁷ at 20 min to 3.0 × 10⁻³ at 4 h, 3.6 × 10⁻² at 6 h and to 1.2 × 10⁻¹ or 7.0 × 10⁻¹ in replicate experiments at 20 h. Transfer (approximate frequency 10⁻² at 20 h) also occurred on the surface of nutrient agar incubated anaerobically, with precautions taken to exclude transfer during processing (see Methods). The addition of mammalian or micrococcal nucleases had no effect on the transfer under aerobic conditions. The optimal molarity of Ca²⁺ and Mg²⁺ for transfer was between 0.001 and 0.025 M, and transfer was abolished by the addition of 0.01 M-citrate to nutrient broth (this contained 0.05 mM-CaCl₂). The effect of cell density on transfer was investigated by altering the oxygen tension during incubation (Lacey & Stokes, 1979). The transfer frequency varied from about 3 × 10⁻³ at a final cell density of 10⁹ colony-forming units (c.f.u.) ml⁻¹ to 6 × 10⁻³ at a density of 10⁷ c.f.u. ml⁻¹.

Effect of prophage 55 in strain 1030tet-r on transfer frequency

Culture 1030(55)tet-r was incubated with 1030SF or 1030(55)SF. Transfer was detected after 30 min, and in each mixture the frequency increased until 20 h, although the final frequencies were less than those obtained for 1030tet-r and 1030(55)SF (Table 7). Examination of 20 derivatives from each mixture showed drug sensitivity and lysogeny consistent with a direction of transfer from the tetracycline-resistant culture to the putative recipient. Thus, the presence of phage 55 in the donor can itself promote transfer although at rather low frequency, but its presence in the donor inhibits transfer to a recipient that also harbours this phage.
Transfer of plasmids from the non-lysogenic donor 1030 to recipients derived from 1030

Non-lysogenic donors 1030neo-r, 1030chl-r, 1030cpns, 1030pen-r, 1030cad-r and 1030cp-1 could not transfer resistance (frequency < 10^-9) to the non-lysogenic recipient 1030s in mixed culture. Transfer to the other lysogenic derivatives of 1030 occurred at variable frequency; the plasmids from 1030cpns, 1030cad-r and 1030pen-r transferred at frequencies from 10^-4 to 10^-8, the plasmids from 1030neo-r and 1030chl-r at frequencies from 1.5 x 10^-2 to 5 x 10^-7 and the plasmid from 1030cp-1 at frequencies from 3 x 10^-1 to 1 x 10^-6. It is assumed from the study of the effects of phage 55 in 1030 that the direction of transfer is from the putative donor to the recipient. The highest frequencies were found when the recipient was lysogenic for phage 55, 88, 71 or 79 and the lowest when lysogenic for phage 52 and 42E where the transfer frequencies were often only fivefold greater than the frequency of spontaneous mutation to streptomycin resistance in the donor.

The chromosomal genes sn, derived from the element cpns (Lacey, 1979), were transduced to 1030 wild using a mitomycin C-induced lysate of strain 6936cpns as donor. The effect of ultraviolet irradiation on a transducing lysate indicated a probable chromosomal locus for sn in 1030 (data not shown). Incubation of 1030sn with 1030s and each of the lysogenic recipients did not result in transfer in any mixture [selection was made with 4.0 pg neomycin ml^-1 plus 400 pg streptomycin ml^-1 since sn confers low level (minimum inhibitory concentration 40 µg ml^-1) resistance to streptomycin]. This finding is consistent with the observations that plasmid genes are readily transferred whilst chromosomal genes are not.

Transfer of cp-2 and cp-3 in mixed cultures of derivatives of strain 1030

When cultures of strains 1030cp-2 and 1030cp-3 were incubated with each lysogenic recipient, transfer occurred at frequencies similar to or slightly greater than that from 1030cp-1. However, both 1030cp-2 and 1030cp-3 transferred their elements at high frequency (2 x 10^-1 and 4 x 10^-1, respectively) to the non-lysogenic strain 1030s. Cultures of 1030cp-2 and 1030cp-3 were inducible with mitomycin C and ultraviolet light; hence, they probably harbour prophages. These filtrates contained about 10^-4 transducing particles per original donor cell, although no plaque-forming activity has been found after the application of mitomycin C-induced lysates to several indicator cultures including 1030 wild. This prophage might become easily associated with, or even be covalently linked to, the resistance determinants of plasmids cp-2 and cp-3. If so, it should be possible to transfer this postulated element through several clones of non-lysogenic 1030. Elements cp-2 and cp-3 each in 10 clones of 1030 were successively transferred (at frequencies of 10^-1 to 10^-3) from 1030s to 1030F and back to 1030s. Ten derivatives of each of the final resistant recipients (1030scp-2, 1030scp-3) were tested for inducibility to mitomycin C and ultraviolet light. All were inducible with either agent. Thus, it is believed that linked transfer of plasmid functions and prophage genome has occurred, because the possibility of simultaneous transfer of plasmid and prophage at the high frequencies obtained is extremely unlikely.

The presence of plasmid cp-1, cp-2 or cp-3 in recipient 1030s did not alter the frequency of transfer of plasmids from either 1030(55)tet-r, 1030(55)neo-r or 1030(55)chl-r in mixed cultures (data not shown).

Simultaneous transfer of plasmids

Cultures were constructed that harboured four plasmids (Table 8) and resistance transfer to recipient 1030s was studied. Co-transfer of either neomycin or tetracycline resistance with resistance to Cd^{2+} occurred from the donor to the recipient as long as the elements cp-2 or cp-3 were in the donor (Table 8). However, examination of five of each of the doubly-resistant recipients revealed that the two genes were subsequently transferred in mixed culture to 1030F separately. Thus, it was inferred that coincidental transfer of the two elements had occurred rather than the transfer of single structures. Mitomycin C-induced
Plasmid transfer in *S. aureus*

**Table 8. Transfer of plasmids in mixed cultures of derivatives of strain 1030**

<table>
<thead>
<tr>
<th>Donor</th>
<th>Recipient</th>
<th>Cd(^{2+})</th>
<th>Tetracycline</th>
<th>Neomycin</th>
<th>Cd(^{2+}) and tetracycline</th>
<th>Cd(^{2+}) and neomycin</th>
</tr>
</thead>
<tbody>
<tr>
<td>1030cp-1 tet-r neo-r chl-r</td>
<td>1030S</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>1030cp-2 tet-r neo-r chl-r</td>
<td>1030S</td>
<td>4.7 × 10^-1</td>
<td>2.0 × 10^-3</td>
<td>2.9 × 10^-4</td>
<td>9.1 × 10^-6</td>
<td>3.0 × 10^-7</td>
</tr>
<tr>
<td>1030cp-3 tet-r neo-r chl-r</td>
<td>1030S</td>
<td>1.9 × 10^-1</td>
<td>6.2 × 10^-3</td>
<td>6.8 × 10^-4</td>
<td>2.2 × 10^-4</td>
<td>6.3 × 10^-7</td>
</tr>
</tbody>
</table>

* No. of resistant colonies/final no. of donors.
† Selection for resistance to stated antibacterial agent(s).

Lysates of culture 1030cp-2 neo-r tet-r chl-r were used to transduce these resistances to 1030 wild. Transduction of *cp*-2 (at a frequency of about 10\(^{-4}\) per original donor cell) was observed, with no evidence of simultaneous transduction of other resistances. Similarly, after phage 88 was propagated on the culture 1030cp-2 neo-r tet-r chl-r each resistance was transduced separately (frequency about 10\(^{-4}\)), but no cotransduction was detected.

**Transduction of tetracycline resistance from lysogenic derivatives of strain 1030**

Mitomycin C-induced lysates of nine tetracycline-resistant lysogenic derivatives of strain 1030 and also non-lysogenic 1030tet-r were examined for their ability to transduce tetracycline resistance to 1030 wild. The cultures lysogenic for phages C, 54, 96, 71, 42E, 88, 55 and 95 transduced tetracycline resistance at frequencies from 6.0 × 10\(^{-4}\) for phage 96 to 3.8 × 10\(^{-7}\) for phage 88 and 1.0 × 10\(^{-8}\) for phage 42E. Lysates of cultures 1030(77)tet-r did not transduce (<10\(^{-9}\)). Culture 1030tet-r was non-inducible and its filtrate after mitomycin C treatment could not transduce tetracycline resistance. Thus, several phages in addition to those traditionally considered to be transducing phages can mediate this type of resistance transfer.

**Discussion**

In addition to the well-known involvement of bacteriophages in transformation and in transduction using cell-free preparations, the presence of bacteriophages in *S. aureus* can affect the occurrence and frequency of transfer of plasmids in mixed cultures in at least five ways. Normal lysogeny in either the donor or the recipient, or the presence of bacteriophage genes as part of a plasmid (*cp*-2, *cp*-3) in the donor, can increase the frequency of spontaneous transfer in mixed cultures, and the presence of some bacteriophages in either donor or recipient can inhibit transfer. Evidence is presented here that spontaneous transfer in mixed cultures has characteristics distinct from that mediated by conventional transduction, specifically: (i) transfer of resistance can occur from a non-lysogenic donor to a recipient; (ii) a culture lysogenic for one phage [1030(52)] cannot acquire resistance by transduction with phage C but can acquire resistance following incubation in mixed culture with 1030(C) derivatives; (iii) the frequency of transfer can be as high as 7.0 × 10\(^{-1}\) (resistant recipients/donor) following mixed culture incubation; (iv) little spontaneous phage activity is seen following subculture of mixed broth cultures to solid agar although a high frequency of transfer may have occurred; (v) filtrates of mixed cultures may not contain agents able to promote transfer, although transfer occurs between cells at high frequency.

However, the transfer of resistance from culture 13136 to 1030 in mixed cultures could well be by spontaneous transduction, i.e. the plasmids are enclosed in phage protein, although it is impossible to establish this with absolute certainty. Similarities between transfer in mixed culture and transfer by transduction are also seen in the requirement for low concentrations (about 0.001 to 0.01 M) of Ca\(^{2+}\) or Mg\(^{2+}\) and the higher frequencies of transfer of plasmid genes compared with transfer of chromosomal genes.

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The elements \(cp-2\) and \(cp-3\) can effect their own transfer at high frequency, probably by becoming part of a phage genome. These elements arose spontaneously when cells of strain M7 were incubated with strain 6936 (Lacey, 1979). Strain M7 probably contains defective bacteriophage(s) (Lacey & Stokes, 1979) and the element \(cpns\) may well have recombined with this genome to produce the elements \(cp-2\) and \(cp-3\). These elements are not identical since the amount and control of beta-lactamase determined by them differs (V. T. Rosdahl, personal communication). Although their origin cannot be identified with certainty, the presence of elements determining production of beta-lactamase that can effect their own transfer among strains of \(S.\) aureus is a considerable threat to antimicrobial therapy. It is notable that the element \(cpns\) passes spontaneously between clones of culture 6936 at a higher frequency than do \(cp-I,\) \(cp-2\) or \(cp-3\) (Lacey, 1979, and unpublished observations). Possibly, phages in this culture inhibit the transfer of the latter elements to a greater extent than that of \(cpns\).

It is suggested that the plasmid transfer in mixed cultures from a non-lysogenic donor be described as "phage-mediated conjugation" since direct cell-to-cell contact seems necessary, although the mechanism is obscure. The staphylococcal cell surface requires further study before detailed understanding of this mechanism is obtained. However, the occurrence of two types of gene transfer in this species, in addition to transformation, could account for the discrepancies mentioned in the Introduction. Because of the requirement for a high molarity of \(\text{Ca}^{2+} (0.1 \text{ M})\) and the absence of DNAase for optimal frequencies, gene transfer by transformation in staphylococci is probably not common in nature. Spontaneous transfer by generalized transduction \textit{in vivo} is also of limited potential because of the probable death of the donor cell and some recipients. In contrast, resistance transfer by phage-mediated conjugation would be expected to occur without a requirement for the death of the donor cell. Most staphylococcal plasmids exist in multiple copies in the cell (Chopra et al., 1973). It is therefore possible for a donor cell to transfer several of its copies of a particular plasmid to different recipient cells with a total increase in the numbers of resistant cells in the culture.

Resistance to neomycin and tetracycline can be co-transferred with the penicillinase plasmids \(cp-2\) or \(cp-3\), although each element is subsequently transferred individually from the recipient.

The presence of phage genome could cause cell-to-cell adhesion, presumably by altering surface proteins, and occasionally two plasmids pass to the recipient coincidentally. Thus, the elements \(cp-2\) and \(cp-3\) not only effect their own transfer, but can mediate transfer of plasmids coding for neomycin or tetracycline resistance. The high frequencies of transfer of plasmids obtained when non-lysogenic derivatives of 1030 are used as donors are probably atypical since most staphylococci are lysogenic.

An incidental finding during these studies has been that lysogenic derivatives of strain 1030 and, to a lesser extent, plasmid-containing cells are consistently more sensitive to both sulphadiazine and trimethoprim than the corresponding wild strain. The presence of the additional cellular DNA may cause an increase in sensitivity to these agents although the mechanism is not established. Lysogeny might increase the efficiency of thymineless death provoked by these agents. These findings could be of value for the isolation of non-lysogenic bacteria or in the detection of plasmid elimination, and might be relevant to the use of these agents therapeutically.

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**REFERENCES**


