Specialized Transduction of Kanamycin Resistance in a Providence Strain

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SUMMARY

Properties of a transducing system with a phage able to transduce a kanamycin-resistance marker of the T compatibility group plasmid R394 at a frequency of $2 \times 10^{-2}$/plaque-forming unit adsorbed are described. The phage was detected in Providence strain P29 transduced to kanamycin resistance by Providence phage PL25 grown on this strain harbouring the R factor. Four P29 transductants, specially selected at the lowest multiplicities of infection of the high frequency transducing (HFT) phage, were defective lysogens. They plated PL25 with an efficiency of 1 and only one liberated low-titre phage spontaneously or on u.v. induction. The defect in maturation function could be corrected by introduction of a wild PL25 prophage. The transducing phage was serologically identical to PL25. It could transduce in single infection, but transduction frequency was increased by the simultaneous presence of homologous non-transducing phage. Transductants did not transfer the kanamycin-resistance marker by conjugation, and produced kanamycin-sensitive segregants at a moderate rate. These segregants could be transduced to kanamycin resistance by the HFT phage. Irradiation of HFT lysates by u.v. produced an exponential fall in transduction frequency. It was concluded that the defective phage transduced by lysogenization. Kanamycin-resistant transductants could themselves be transduced to streptomycin resistance by PL25 reared on a streptomycin-resistant mutant. Lysogenic transductants produced by the HFT phage did not always liberate HFT phage on u.v. induction. Possible explanations are considered.

INTRODUCTION

Transduction of a Proteus rettgeri R factor R394 (Coetzee, Datta & Hedges, 1972) by vector phage 34 (Coetzee & Sacks, 1960) to Proteus mirabilis strain PM5006 yielded some transductants which only registered the kanamycin-resistance marker of the plasmid and could not transmit the marker by conjugation. The latter transductants were lysogenic and liberated phage able to transduce kanamycin resistance to PM5006 at a frequency of $3 \times 10^{-2}$/plaque-forming unit (p.f.u.) adsorbed (Coetzee, 1974b). Transfer of R394 to these transductants yielded some progeny which, on u.v. induction, produced lysates that could transduce markers of kanamycin and ampicillin resistance at frequencies of $4 \times 10^{-3}$/p.f.u. adsorbed (Coetzee, 1975). Apart from the additional marker of the latter phage, properties of the two high frequency transducing (HFT) phages were similar. They were serologically identical to phage 34 and could transduce at a multiplicity of infection (m.o.i.) of less than 0.01. Transduction frequencies were increased about tenfold by the presence of homologous non-transducing phage. HFT lysates could also transduce various chromosomal markers at low frequencies. Transductants produced kanamycin-sensitive lysogenic segregants at high rates. Strain PM5006 is cryptically lysogenic (Krizsanovich, 1973) for a phage
## Table 1. Bacteria, plasmids and phage

<table>
<thead>
<tr>
<th>Bacteria</th>
<th>Properties*</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Providence strain NCTC9211 biogroup 1.</td>
<td>Recipient for general transducing phage PL25.</td>
<td>Coetzee (1974a); Coetzee et al. (1966)</td>
</tr>
<tr>
<td>Providence strain NCTC9211 biogroup 1.</td>
<td>Resistant to A</td>
<td>Coetzee et al. (1966)</td>
</tr>
<tr>
<td>Strain P29 lysogenized with PL25</td>
<td>Resistant to 1 mg streptomycin sulphate/ml</td>
<td>Coetzee et al. (1966)</td>
</tr>
<tr>
<td>Spontaneous rough colonial variant</td>
<td></td>
<td>Coetzee (unpublished)</td>
</tr>
<tr>
<td>Auxotrophic mutants produced by NTG treatment</td>
<td></td>
<td>Coetzee (unpublished)</td>
</tr>
<tr>
<td>Nalidixic acid-resistant mutant of 353, an F- lac+, pro, met mutant of E. coli K12</td>
<td></td>
<td>Clowes &amp; Hayes (1968); Coetzee et al. (1972)</td>
</tr>
<tr>
<td>Strain P29 with R plasmid R394 introduced by conjugation. R394 is a T compatibility group plasmid with markers A, K</td>
<td></td>
<td>Coetzee (unpublished); Coetzee et al. (1972)</td>
</tr>
<tr>
<td>Original transductant clone produced at high m.o.i. Superior line indicates that R394 was introduced by transduction with vector phage PL25</td>
<td></td>
<td>Coetzee (unpublished)</td>
</tr>
<tr>
<td>General transducing phage for P29</td>
<td></td>
<td>Coetzee et al. (1966)</td>
</tr>
<tr>
<td>Produced by lytic infection of P29str-r with PL25</td>
<td></td>
<td>Coetzee et al. (1966)</td>
</tr>
<tr>
<td>Produced by lytic infection of P29(R394) with PL25</td>
<td></td>
<td>Coetzee et al. (1966)</td>
</tr>
</tbody>
</table>

* Symbols designate resistance to: A, ampicillin; K, kanamycin.

morphologically and serologically identical to phage 34, and many features of the transduction systems were explicable in terms of interaction between resident cryptic prophage and the HFT phage.

Organisms of the Providence group may be readily distinguished from *P. mirabilis* by simple biochemical tests but they have properties in common, and the tendency has been to arrange Providence in the Proteus-Providence group (see Coetzee, 1972; Prozesky et al. 1973). Generalized transducing phages for certain strains of Providence have been described (Coetzee, Smit & Prozesky, 1966). During an investigation (Coetzee, unpublished) of R factor transduction by one of these phages (PL25) to strain NCTC9211 (re-named P29) it was found that transductants which had received the kanamycin-resistance marker of R394 were unable to transmit their antibiotic resistance by conjugation. It meant that the transmissibility system of the R factor was defective in Providence P29 or that the markers in question had integrated in another replicon which was not capable of conjugal transfer. This was reminiscent of the *P. mirabilis* systems just described. Non-homologous genes were being transduced and chances of recombination with the recipient chromosome were therefore reduced. This report deals with an investigation of the possibility that transductants were produced by phages carrying resistance genes and still able to lysogenize.

## METHODS

### Bacteria, plasmids and phages

These are listed in Table 1.

### Media

Media were as described previously (Coetzee et al. 1973) except that the minimal medium was that of Lederberg (1950). Selection for drug resistance was on MacConkey agar containing one or more of the following: ampicillin, kanamycin, nalidixic acid and
Lysate | Titre (p.f.u./ml) | Multiplicity of infection | Recipient | Kanamycin resistant | Streptomycin resistant |
--- | --- | --- | --- | --- | --- |
U.v.-induced | 6 x 10^7 | 10^-4 | p29 | 4 x 10^-2 | — |
\(\text{p}29\text{(R}394\text{j-o)}\) | 1 x 10^9 | 10^-3 | \(\text{p}29\text{gou-1})\) | 5 x 10^-4 | — |
\(\text{PL}25\text{-p}29\text{str-r)}\) | 1 x 10^9 | 10^-1 | p29 | 5 x 10^-7 | — |
| | | 10^-9 | \(\text{p}29\text{gou-1})\) | < 2 x 10^-7 | — |
| | | 10^-1 | \(\text{p}29\text{(394)}\text{o})\) | 1 x 10^-7 | — |

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Table 2. Transductions by u.v.-induced lysate of \(\text{p}29\text{(R}394\text{j-o)}\) and by phage \(\text{PL}25\text{-p}29\text{str-r)}\)

strepotomycin. Ampicillin, kanamycin and nalidixic acid were used at 50 µg/ml unless otherwise stated. Streptomycin was used at 1 mg/ml. Incubation temperature was 30 °C.

Conjugal transfer of plasmids and preparation of lytic phage lysates. Methods were as described by Coetzee et al. (1973).

Phage neutralization. This was done according to the method of Adams (1956). The antiserum used was raised in a rabbit against phage PL25. It was stored freeze-dried, and had a K value of 120/min (Coetzee et al. 1966).

Determination of the stability of the kanamycin-resistance marker in \(\text{p}29\text{(R}394)\text{)}\). This was done as described by Coetzee (1974b) with the exception that PL25 antiserum (K = 60) was substituted.

Ultraviolet irradiation of phage and phage induction. This was done as described (Coetzee, 1974b).

Concentration of phage lysates. The method described for P22 by Yamamoto & Alberts (1970) was used.

Replica screening technique for detection of HFT phage and standard transduction experiments. These were done exactly as described by Coetzee (1974b). Briefly, adsorption mixtures were filtered and membranes with impinged bacteria were incubated on nutrient agar for 1 h before transfer to MacConkey agar containing 50 µg kanamycin/ml. For streptomycin resistance, membranes were maintained on agar for 4 h before transfer to MacConkey agar containing 1 mg streptomycin/ml.

RESULTS

Providence strain \(\text{p}29\text{)}\) is naturally resistant to ampicillin and sensitive variants have not been obtained (Coetzee, unpublished). Consequently only the kanamycin-resistance marker of R factor R394 could be selected. Sterile lysates prepared from strain \(\text{p}29\text{(R}394\text{)}\) with phage PL25 had titres of \(5 x 10^9\) p.f.u./ml and the lysates, designated PL25-p29(R394), transduced the kanamycin-resistance marker to \(\text{p}29\) at a frequency of \(5 x 10^-6\) p.f.u. adsorbed. The m.o.i. of the phage was about 2. Eight of 10 transductant clones failed to transfer kanamycin (or ampicillin) resistance to \textit{Escherichia coli} \(\text{J}53\text{-1)}\) by conjugation. One of the eight clones was chosen for further study and designated \(\text{p}29\text{(R}394\text{j-o)}\).

Properties of transductant \(\text{p}29\text{(R}394\text{j-o)}\)

The conjugation with \(\text{J}53\text{-1)}\) as recipient and \(\text{p}29\text{(R}394\text{j-o)}\) as donor was repeated with selection for progeny on nalidixic acid plates (used for counterselection) and containing kanamycin or ampicillin. The crosses were sterile. Phage PL25 did not form plaques on \(\text{p}29\text{(R}394\text{j-o)}\), and overnight broth cultures of the strain contained, per ml, about \(1 x 10^6\)
p.f.u. of a phage serologically identical to PL25 when assayed on P29. These titration plates were replicated to kanamycin agar which was then incubated overnight. The replicas, especially of the lower dilution plates, showed practically confluent growth. This has been described as a screening method for phage capable of high frequency transduction of antibiotic resistance (Coetzee, 1974b). Induction of the strain by u.v. irradiation yielded a lysate with a titre of about $6 \times 10^7$ p.f.u./ml. The lysate was capable of HFT of the kanamycin marker at a m.o.i. of $10^{-6}$ (Table 2). Minute colonies suggestive of abortive transduction were not observed. The number of transductant clones which appeared on membranes transferred to agar containing 50 $\mu$g kanamycin/ml immediately after filtration at the end of the adsorption period, could not be increased by prior incubation on non-selective media.

It was concluded that P29 expressed resistance to this concentration of kanamycin immediately after the termination of phage adsorption. Membranes had to be incubated for about 1 h on non-selective agar for transductants to achieve full resistance to 500 $\mu$g kanamycin/ml. A similar situation was encountered with the P. mirabilis HFT phage for kanamycin resistance (Coetzee, 1974b). Strain P29(R394)-0 could be transduced to streptomycin resistance by PL25·P29str-r (Table 2). This is in accord with findings (Coetzee et al. 1966) that auxotrophs of P29 could be transduced to prototrophy and the prototrophs could then be transduced to streptomycin resistance by the above phage. A rough mutant of P29 which could not be transduced to streptomycin resistance by the general transducing phage (Table 2) could be transduced to kanamycin resistance by the HFT lysate at a lower frequency than usual. This could be due to the rough mutant adsorbing phage so poorly that only HFT phage could register its marker. Four kanamycin-resistant clones produced by the infection of P29 at a m.o.i. of $10^{-6}$ in the presence of PL25 antiserum and designated $P29(R394)_{1-4}$ were chosen for further study.

**Properties of $P29(R394)_{1-4}$**

The four clones resembled P29(R394)-0 in antibiotic resistance and did not transmit resistance by conjugation to E. coli 153-1. They differed from the original transductant in that only one (No. 4) liberated phage spontaneously or on u.v. induction and then only in low titre (approx. 50 p.f.u./ml), and all four plated PL25 with an efficiency of 1. Unlike P29(R394)-0 these transductants were selected at low m.o.i. and arose in the presence of phage antiserum. They probably resulted from single particle infection, and the above-mentioned properties suggested defective immunity and maturation functions of the transducing phage. The four transductants segregated kanamycin-sensitive clones (designated $P29(R394)_{kS}$) at about $1 \times 10^{-4}$/bacterium/generation. Segregants were fully sensitive to PL25 and did not liberate phage. The rate of segregation of the marker is evidence that transduction proceeded by lysogenization (Luria, Adams & Ting, 1960). These four transductants, like P29(R394)-0, could be transduced to streptomycin resistance by phage PL25·P29str-r (not shown). Turbid centres of the PL25 plaques on the four kanamycin-resistant transductants $P29(R394)_{1-4}$ were streaked on kanamycin agar, and single clones corresponding to each transductant, lysogenic for PL25, were obtained. These were designated $P29(R394PL25)_{1-4}$.

**Properties of $P29(R394PL25)_{1-4}$**

These strains differed from $P29(R394)_{1-4}$ in that they did not plate PL25 and liberated phage spontaneously which was serologically identical to PL25. They could however be transduced to streptomycin resistance by phage PL25·P29str-r, and gave rise to kanamycin-sensitive lysogenic segregants (designated $P29(R394PL25)_{kS}$) at a frequency of about
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Table 3. Transduction with u.v.-induced lysates of P29(R394PL25)1 to 4

<table>
<thead>
<tr>
<th>Strain from which lyse was induced</th>
<th>Titre (p.f.u./ml)</th>
<th>Multiplicity of infection</th>
<th>Recipient</th>
<th>Kanamycin-resistant (or prototrophic) transductants*/p.f.u. adsorbed</th>
</tr>
</thead>
<tbody>
<tr>
<td>P29(R394PL25)-1</td>
<td>6 × 10^6</td>
<td>10^-6</td>
<td>P29</td>
<td>2 × 10^-2</td>
</tr>
<tr>
<td></td>
<td>8 × 10^7</td>
<td>10^-6</td>
<td>P29</td>
<td>1 × 10^-2</td>
</tr>
<tr>
<td></td>
<td>1 × 10^7</td>
<td>10^-4</td>
<td>P29</td>
<td>1 × 10^-4</td>
</tr>
<tr>
<td></td>
<td>3 × 10^7</td>
<td>10^-3</td>
<td>P29</td>
<td>8 × 10^-6</td>
</tr>
<tr>
<td>P29(R394PL25)-2</td>
<td>6 × 10^6</td>
<td>10^-6</td>
<td>P29(PL25)</td>
<td>1 × 10^-3</td>
</tr>
<tr>
<td></td>
<td>1 × 10^9</td>
<td>10^-1</td>
<td>P29arg-1</td>
<td>&lt; 5 × 10^-9</td>
</tr>
<tr>
<td></td>
<td>1 × 10^9</td>
<td>10^-1</td>
<td>P29met-1</td>
<td>&lt; 5 × 10^-9</td>
</tr>
<tr>
<td></td>
<td>1 × 10^9</td>
<td>10^-1</td>
<td>P29ura-1</td>
<td>&lt; 5 × 10^-9</td>
</tr>
<tr>
<td>P29(R394PL25)-1</td>
<td>6 × 10^6</td>
<td>10^-6</td>
<td>P29(R394)k^8</td>
<td>1 × 10^-2</td>
</tr>
<tr>
<td></td>
<td>6 × 10^6</td>
<td>10^-6</td>
<td>P29(R394PL25)k^8</td>
<td>5 × 10^-4</td>
</tr>
</tbody>
</table>

* Kanamycin-resistant transductions were as described in Methods. The phages used for prototrophic transductions were concentrated and membranes with impinged cells were placed on minimal medium immediately after filtration.

1 × 10^-4/bacterium/generation. Induction by u.v. irradiation yielded lysates with titres of 6 × 10^6 to 8 × 10^7 p.f.u./ml. Lysates invariably contained an unusually high proportion (up to 10%) of particles which produced clear plaques on P29. Some of these clear and turbid plaques were picked off and used to prepare separate lysates of P29(R394)-1, the first of the four transductant clones produced at low m.o.i. The lysates transduced P29 to kanamycin resistance at a frequency of about 2 × 10^-9/p.f.u. adsorbed (not shown). It is probable therefore that kanamycin-resistance transduction by HFT phage was effected by non-plaque forming particles (see Smith-Keary, 1966). Lysates induced by u.v. irradiation of strains P29(R394PL25)1 to 4 differed in their transduction frequencies for kanamycin resistance (Table 3). The reason is not understood. The producer strains were similar in all other respects and lysates had comparable plaque-forming titres. This variation in transduction frequency was also a variable property: further lysates prepared from strains which yielded low transducing frequency phage sometimes yielded phage with much higher transducing ability. Lysate P29(R394PL25)-2 (Table 3, line 2) had many more clear-plaque forming particles than the other lysates and so it was unlikely that results could be explained in terms of lysis of transductants by virulent phage.

Transduction of ability to yield HFT lysates

Seventeen transductant clones from experiments done simultaneously with those reported in Table 3 but produced at a m.o.i. of about 4, and therefore lysogenic, were examined for ability to yield HFT phage on u.v. induction. Titres of the lysates varied from 5 × 10^6 to 7 × 10^7 p.f.u./ml. Only ten of the lysates transduced at a frequency of about 2 × 10^-2/p.f.u. adsorbed. The transduction frequencies of the remaining seven lysates varied between 4 × 10^-4 and 3 × 10^-4/p.f.u. adsorbed.

These seven transductants segregated kanamycin-sensitive clones at the same rate as the other ten. Many other lysogenic transductants phenotypically identical to those mentioned above were tested during the course of this investigation. Failure to yield lysates with transduction frequencies above 1 × 10^-2/p.f.u. adsorbed was quite a common, but variable, feature.
The HFT phage was concentrated to $1 \times 10^8$ p.f.u./ml and used in transduction experiments with various auxotrophs of P29 as recipients. Unlike the HFT phages described by Kondo & Mitsuhashi (1964), Jessop (1972) and Coetzee (1974a, 1975), lysates could not transduce other markers (Table 3). Results presented in Table 3 also indicate that P29(PL25) is transduced at a frequency about 20-fold lower than P29. If HFT transduction proceeds by lysogenization, then this could be explained on the grounds that, as the principal attachment site was occupied, transducing phage had to contend with less efficient secondary sites. Weigle (1957) and Hoppe & Roth (1974) encountered similar phenomena with λigg and a P22 HFT lysate respectively. Transduction of phage-sensitive, as well as lysogenic kanamycin-sensitive, segregants with HFT phage lysates revealed that the former were transduced at a frequency similar to wild-type P29 while with the latter the frequency was about 20-fold lower (Table 3). In contrast, generalized transduction (Table 2) was not affected to the same degree by recipients being lysogenic and this fact supported the thesis that HFT proceeded by lysogenization.

Relatedness of PL25 and transducing particles of HFT lysates

Samples of a HFT lysate derived from P29(R394PL25)-1 (Table 3) were treated with dilutions of PL25 antiserum and plaque-forming and transducing activities of the lysate determined as a function of amount of antiserum added. Transducing activity was assayed in the presence of helper phage to ensure that diminished transducing activity was not due to loss of helper. The two parameters diminished at the same rate (Fig. 1). This is evidence for the serological identity of the particles concerned and eliminates the possibility that the vector was a serologically different prophage present in P29.

Defectiveness of HFT phage

It was demonstrated (Tables 2 and 3) that the HFT phage could transduce at very low m.o.i. and it was probable that a single particle could transduce. Results presented in Fig. 2 again emphasize this point. The rapid fall of the curve for HFT alone (solid circles) between m.o.i. of 1 and $10^{-2}$ (where chances of simultaneous infection by transducing and normal particles present in the lysate are slight) is not indicative of mutual aid between particles (Chan et al. 1972; see also Rae & Stodolsky, 1974). The presence of helper phage, at a m.o.i. of 3, increased the transduction frequency about fivefold.

Effect of U.V. irradiation of HFT lysates on transduction

A HFT lysate was irradiated for different periods of time. Samples were titrated and quantitative transduction experiments performed with each sample. The curves (Fig. 3) relating time of irradiation to transduction frequency and phage survival, showed exponential declines with the slope of the former about half that of the latter. This implies that some phage functions are not required for transduction. The slope of the transduction curve differs from that obtained when the frequency of streptomycin-resistance transduction was measured as a function of time of irradiation of PL25 (Coetzee et al. 1966) where an increase in transductants, at low doses, was followed by slow inactivation. The initial increase was attributed to greater frequency of recombination between chromosome and the exogenote (Arber, 1958; Luria et al. 1960). Arber (1960) demonstrated an exponential fall in numbers of transductants of various episomes with increasing doses of irradiation applied to transducing lysates, and Luria et al. (1960) ascribed the simple decline in numbers of lac+ Shigella transductants produced by irradiation of Pf lysates of E. coli to failure of the
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Fig. 1. Effect of phage PL25 antiserum on transduction frequency of a HFT lysate derived from p29(R394PL25)-1. One ml of serial twofold dilutions of phage PL25 antiserum in 0.85% (w/v) NaCl were mixed with an equal volume of lysate and incubated for 5 min. The p.f.u. titre of the samples was determined and then 0.5 ml of phage PL25-p29(titre 6 x 10^10 p.f.u./ml) was added to the samples and quantitative kanamycin-resistance transductions performed. ●, Plaque-forming titre/ml; ○, no. kanamycin-resistant transductants/ml.

Fig. 2. Effect of helper phage on efficiency of transduction. Transductions were with HFT lysate p29(R394PL25)-1 at different m.o.i. Helper phage, PL25-p29Str-r, was used at m.o.i. of 2-9. ●, HFT phage; ○, HFT phage and helper phage.

Fig. 3. The effect of u.v. irradiation on a HFT lysate. Quantitative transduction experiments were done with the irradiated lysates at a m.o.i. of 1 p.f.u. of the unirradiated phage. ○, Transducing activity; ●, plaque formation.
particles to persist as prophages. A similar interpretation may apply here. This supports the contention, derived from the rate at which kanamycin-sensitive segregants arose and also from the disparity in the rates of HFT to lysogenic and non-lysogenic recipients, that the R factor genes formed part of the phage which transduced by lysogenization (Coetzee, 1974b, 1975).

**Discussion**

Luria *et al.* (1960) were the first to employ transduction of non-homologous genes to select HFT phage capable of conveying the genes while retaining ability to lysogenize. This principle has been brilliantly exploited by Hoppe & Roth (1974); see also Rae & Stodolsky (1974). Specialized transducing phages vary in the complement of phage functions retained. There are the tet<sup>+</sup> P22 (Chan *et al.* 1972) and P<sub>1dl</sub> (Rae & Stodolsky, 1974) phages which cannot multiply or lysogenize on single infection, the variants of phage epsilon (Kameda *et al.* 1965) and P22 (Hoppe & Roth, 1974) which can lysogenize but exhibit no immunity or maturation functions, and the semi-defective particles of <i>ldg</i> and P22 which can lysogenize and retain immunity properties but possess no (Campbell, 1957; Smith-Keary, 1966) or virtually no (Dubnau & Stocker, 1964) maturation function. Finally, there are converting phages P1CM (Kondo & Mitsuhashi, 1964) and φ Amp<sup>+</sup> (Williams Smith, 1972) which retain all phage functions. The phage described here has many of the properties of the phage epsilon tetracycline-resistance HFT lysate described by Kameda *et al.* (1965). The latter particles could also transduce on single infection and were defective in immunity and maturation functions. They differed from the Providence phage in that helper phage did not affect transduction frequency. The helper effect encountered here may be explained as by Coetzee (1974b, 1975), that non-transducing phage supplied some function which increased the probability of successful lysogenization by transducing phage. It was shown that the HFT Providence phage was defective (amongst other functions) in maturation and this could be corrected by a normal prophage. A disturbing feature was the fact that though phage titres were comparable, the transduction frequencies of different lysates varied considerably. The defaulting lysates were unlikely to have arisen by double cross-over events with the chromosome as Matsushiro (1963) sometimes encountered with φ <i>8dlt</i>, as such an explanation is incompatible with the frequency of segregation of kanamycin-sensitive clones from all transductants tested (see Bernstein, Rolfe & Onodera, 1973). In addition, the Providence HFT phage was not capable of general transduction. A possible explanation is that a permutatation of the excision event (Luria *et al.* 1960; Matsushiro, 1963; Rae & Stodolsky, 1974; see also Weigle, Meselson & Paigen, 1959; Adler & Templeton, 1963) of the prophage occurred in individual bacteria of a lysate and this resulted in individual vegetative phage genomes containing complete or defective portions of the kanamycin-resistance gene. Watanabe *et al.* (1972) found that nine out of ten transductants obtained from a P22 HFT lysate for tetracycline resistance produced HFT lysates on u.v. induction. Schmieger (1972), Gratia (1973), Wall & Harriman (1974) and Backhaus & Schmieger (1974) have demonstrated that phage does not necessarily play the passive ‘phenotypic mixing’ role in general transduction previously (see Ozeki & Ikeda, 1968) assigned to it and the possibility exists that PL25 acquired the kanamycin (and possibly ampicillin) genes from the R plasmid by integration of the R factor into the phage on the host chromosome.

Many PL25 lysates of P29 strains harbouring various R factors have been screened (Coetzee, unpublished) and only lysates of P29 carrying R394 yielded HFT phage for antibiotic resistance. The antibiotic-resistance markers of R394 have also combined with the <i>P. mirabilis</i> transducing phage 5006M (Coetzee, 1974b, 1975). It may be that this proclivity of
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R394 for recombination with phage genomes results from its resistance markers being contained in transposons (Hedges & Jacob, 1974). Transposition on to vegetative, or prophage, genes could account for the missing phage functions, but the increased molecular weight of the genome could result in encapsidation problems. Physical characteristics of the genome of PL25 have not yet been determined but particles of the HFT phage can transduce on single infection. They thus differ from the particles of P22 tet\(^4\) and P1\(^{dl}\) phages described by Chan et al. (1972) and Rae & Stodolsky (1974), respectively, which must be complemented by another particle to achieve expression.

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REFERENCES

JESSOP, A. P. (1972). A specialized transducing phage of P22 for which the ability to form plaques is associated with transduction of the proAB region. Molecular and General Genetics 114, 214–222.


