Unstable Generalized Transduction in Achromobacter

By D. R. WOODS and JENNIFER A. THOMSON*

Department of Microbiology, Rhodes University, Grahamstown, South Africa

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SUMMARY

Six auxotrophic markers of a halotolerant collagenolytic strain of Achromobacter were transduced by four $\alpha$ phages. Abortive transduction was also demonstrated. The generalized transduction system is unusual as the transductants were unstable, characteristic of transduction by lysogeny. The Achromobacter strain is a cryptic lysogen for $\alpha$ and purified transductants were either sensitive or resistant to $\alpha$. Purified clones from four resistant transductants and one sensitive transductant liberated phage spontaneously. The host ranges of these spontaneous phages differed from that of the $\alpha$ phage used for the transduction experiment. Some initially resistant transductants became semi-sensitive to $\alpha$ (efficiency of plating [e.o.p.] $10^{-1}$ to $10^{-3}$) after repeated cloning.

INTRODUCTION

Four closely related bacteriophages ($\alpha$ phages) specific for halotolerant, collagenolytic strains of Achromobacter were isolated (Thomson & Woods, 1974). Achromobacter strains become resistant to the $\alpha$ phages and do not adsorb phage when they are lysogenized by one of them. The strain, Achromobacter sp. 2, has been shown to be a cryptic lysogen for phage $\alpha$ and is not immune to superinfection by homologous phage. As the prophage can be induced with mutagens, it is proposed that the cryptic nature of the prophage is not due to an extensive phage deletion (Fischer-Fantuzzi & Calef, 1964) but due to the mode of prophage integration and a defective excision mechanism (Krizsanovich, 1973; Thomson & Woods, 1974). Since transduction is especially useful for the analysis of genetic fine structure and function (Hayes, 1968), the transducing abilities of the $\alpha$ phages were tested with a view to developing a genetic mapping system in Achromobacter and studying the genetics of collagenase production.

Generalized transduction of a number of markers usually yields stable transductants by integration (Zinder, 1953) and specialized transduction of markers adjacent to the prophage attachment site yields unstable heterogenotes by lysogenization (Morse, Lederberg & Lederberg, 1956). Specialized transducing particles can only be obtained by lysogen induction, while phage from lytic infection can perform generalized transduction (Zinder, 1955; Morse et al. 1956). Generalized transducing particles usually contain bacterial DNA synthesized before infection and little or no phage DNA (Ikeda & Tomizawa, 1965; Ebel-Tsipis, Botstein & Fox, 1972). The formation of P22 transducing particles occurs independently of the known bacterial or phage recombination systems (Ebel-Tsipis, Fox & Botstein, 1972).

Generalized transducing phages have occasionally, however, been reported to yield heterogenote transductants (Luria, Adams & Ting, 1960). Moreover, Chakrabarty &

* Present address: Department of Microbiology and Molecular Genetics, Harvard Medical School, Boston, Massachusetts 02115, U.S.A.
Gunsalus (1969) showed that transduction of mandelate genes in \textit{Pseudomonas putida} by phage pf16ha could yield stable or unstable transductants, depending on the degree of homology between incoming and recipient DNA as well as on the residual phage function in the transducing particles. Thus the distinction between generalized and specialized transduction may not always be clearly defined.

\section*{Methods}

\textbf{Media.} Nutrient broth (Difco) supplemented with 0:4 M-NaCl was used. For double-agar-layer phage assays, the media were as described previously (Thomson & Woods, 1973). Minimal medium was that of Grabow & Smit (1967) supplemented with 0:4 M-NaCl.

\textbf{Bacteriophages.} Four \(\alpha\) phages (Thomson & Woods, 1974) were used.

\textbf{Isolation of mutants.} Auxotrophic mutants of \textit{Achromobacter} sp. 2 (Thomson, Woods & Welton, 1972) were isolated by treatment of exponentially-growing broth cultures with 45 \(\mu\)g \(N\)-methyl-\(N'\)-nitro-\(N\)-nitrosoguanidine (NTG)/ml for 15 min at 37 \(^\circ\)C. Six auxotrophs were isolated, \textit{arg, cysA} (temperature sensitive), \textit{cysB} (non-temperature sensitive), \textit{met, pro} and \textit{trp}.

\textbf{Phage techniques.} Phage lysates were prepared according to the double-agar-layer method of Adams (1959), and the rapid loss of infectivity (100-fold) after one week (Thomson & Woods, 1974) was prevented by concentration and purification. Concentration was achieved by a two-phase system using Na-Dextran 500 (Pharmacia) and polyethylene glycol 6000 originally developed by Albertsson (1960). The concentrated phage was purified by caesium chloride density gradient centrifugation.

\textbf{Transduction.} Overnight broth cultures of auxotrophs were diluted 1:10 and aerated at 30 \(^\circ\)C for 5 h (1 \times 10^9 cells/ml). Two ml of culture were resuspended in 4 ml phage lysate prepared on the wild-type strain at a multiplicity of infection (m.o.i.) of 1 to 5, (where the m.o.i. is defined as per plaque-forming unit, (p.f.u.) adsorbed. The transduction mixture was incubated for 1 h for phage adsorption. Cells were harvested by centrifugation, resuspended in 0:4 M-NaCl and washed. The cells were added to minimal agar plates either by spreading or by means of a soft minimal agar overlay. Plates were scored for prototrophs after incubation at 30 \(^\circ\)C for 2 to 4 days. Controls consisted of cells incubated with 4 ml 0:4 M-NaCl and 4 ml phage propagated on the homologous strain. Phage sterility was tested by spreading 0:1 ml on to salt and minimal agar. Control transduction experiments were also carried out using phage pre-incubated with antiphage serum (\(k = 10\)) for 15 min.

\textbf{Small transductant colonies.} The small transductant colonies were streaked on to minimal agar and the isolated colonies assayed for cell counts on salt and minimal agar. The number of cells in these small colonies was compared with that of wild-type colonies on minimal agar.

\textbf{Transductant stability.} Transductants were streaked on to minimal agar before suspension in nutrient broth (5 \times 10^8 cells/ml) and incubated for 18 h. Dilutions were plated on to salt and minimal agar and the rate of segregation per cell per generation calculated after incubation for 48 h. The loss of \textit{trp} by transductants was investigated by growth of the transductants in nutrient broth and nutrient broth containing 10 \(\mu\)g acriflavine/ml. Dilutions were plated on to salt agar and 150 colonies from both acriflavine-treated and control cultures were tested for growth on minimal agar.

\textbf{Phage characteristics of transductants.} Twenty \textit{Trp}\textsuperscript{+} transductants obtained with an \(\alpha\text{3a}\) transducing lysate at a m.o.i. of 1 were purified by streaking on minimal agar (1st cloning). Duplicate samples of three clones from each of these 20 streaks were restreaked on minimal agar (2nd cloning). The 120 clones were again purified by streaking twice more on minimal
Table I. Transduction by Achromobacter phages*

Auxotrophic mutants of Achromobacter sp. 2 were used as the recipient strains at a m.o.i. of 1 to 5.

<table>
<thead>
<tr>
<th>Phage</th>
<th>Marker</th>
<th>$10^{-7} \times$ Transduction rate/ p.f.u. adsorbed</th>
<th>Phage</th>
<th>Marker</th>
<th>$10^{-7} \times$ Transduction rate/ p.f.u. adsorbed</th>
</tr>
</thead>
<tbody>
<tr>
<td>α1</td>
<td>arg</td>
<td>0:13</td>
<td>α3a</td>
<td>arg</td>
<td>3:3</td>
</tr>
<tr>
<td></td>
<td>cysA</td>
<td>0:9</td>
<td></td>
<td>cysA</td>
<td>3:2</td>
</tr>
<tr>
<td></td>
<td>cysB</td>
<td>1:5</td>
<td></td>
<td>cysB</td>
<td>6:5</td>
</tr>
<tr>
<td></td>
<td>met</td>
<td>1:77</td>
<td></td>
<td>met</td>
<td>4:3</td>
</tr>
<tr>
<td></td>
<td>pro</td>
<td>1:6</td>
<td></td>
<td>pro</td>
<td>4:8</td>
</tr>
<tr>
<td></td>
<td>trp</td>
<td>1:7</td>
<td></td>
<td>trp</td>
<td>7:3</td>
</tr>
<tr>
<td>α2</td>
<td>arg</td>
<td>0:0</td>
<td>α3b</td>
<td>arg</td>
<td>3:5</td>
</tr>
<tr>
<td></td>
<td>cysA</td>
<td>3:1</td>
<td></td>
<td>cysA</td>
<td>4:6</td>
</tr>
<tr>
<td></td>
<td>cysB</td>
<td>0:0</td>
<td></td>
<td>cysB</td>
<td>6:0</td>
</tr>
<tr>
<td></td>
<td>met</td>
<td>1:85</td>
<td></td>
<td>met</td>
<td>4:0</td>
</tr>
<tr>
<td></td>
<td>pro</td>
<td>0:74</td>
<td></td>
<td>pro</td>
<td>5:0</td>
</tr>
<tr>
<td></td>
<td>trp</td>
<td>2:0</td>
<td></td>
<td>trp</td>
<td>7:6</td>
</tr>
</tbody>
</table>

* Transductants isolated by spreading cells on to minimal agar.

Results

Transduction. All four α phages were able to transduce the auxotrophs to prototrophy at a m.o.i. of 1 to 5 (Table I). Only those experiments where controls showed no growth were recorded. No transductants were formed when the transducing lysate was pre-treated with antiphage serum or when phage lysates propagated on the homologous strain were used.

When α3a was used to transduce the trp marker and the transductants were selected by spreading on to minimal agar, a transduction frequency of $7.3 \times 10^{-7}$/p.f.u. adsorbed (Table 1) was obtained. This low transduction frequency was increased to $1.3 \times 10^{-6}$/p.f.u. adsorbed by adding the cells to minimal agar by means of a soft minimal agar overlay.

Small transductant colonies. Apart from wild-type transductant colonies two types of small transductant colonies were observed. The first type were approximately a quarter of
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Table 2. *Phage characteristics of transductants obtained with an a3a transducing lysate at a m.o.i. of 1*

Twenty transductants (Nos. 1 to 20) were streaked on to minimal agar and duplicate samples of three clones from each transductant were cloned a further three times on to minimal agar. Phage characteristics were determined after each cycle of cloning.

<table>
<thead>
<tr>
<th>Type</th>
<th>All 3 clones ex transductions initially resistant but at least one clone becoming semi-sensitive after 3rd or 4th cloning</th>
<th>Spontaneous and erratic phage liberation by one or more of the clones ex transductants*</th>
<th>U.V. induction of phage</th>
<th>Total no. of transductants and their identification numbers</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>4 (1-4)</td>
</tr>
<tr>
<td>2</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>8 (5-12)</td>
</tr>
<tr>
<td>3</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>1 (13)</td>
</tr>
<tr>
<td>4</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>2 (14, 15)</td>
</tr>
<tr>
<td>5</td>
<td>-</td>
<td>+</td>
<td>-</td>
<td>1 (16)</td>
</tr>
<tr>
<td>6</td>
<td>-</td>
<td>-</td>
<td>+†</td>
<td>1 (17)</td>
</tr>
<tr>
<td>7</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>2 (18, 19)</td>
</tr>
<tr>
<td>8</td>
<td>+</td>
<td>-</td>
<td>+</td>
<td>1 (20)</td>
</tr>
</tbody>
</table>

* Transductants went through a minimum of one cycle of cloning and growth in broth without phage release prior to spontaneous phage liberation.
† Phage liberation by resistant clone.

The size of the wild-type colonies. These small transductant colonies were obtained by both the spreading and overlay techniques and occurred most frequently when the transducing lysate used was at a m.o.i. of 50 to 100. They were not abortive transductants as every cell in the colony could grow on both minimal and salt agar. The number of cells in a small colony on minimal agar was 2.4 × 10⁵ compared with 6.0 × 10⁶ in wild-type colonies on minimal agar. The colonies derived from the cells of the original small colony were likewise of the same small size.

The second type of small transductant colonies were abortive transductants. Abortive transduction was only obtained when the overlay plating technique was used.

Transductant stability. The average rate of segregation of *trp*⁻ in Trp⁺ transductants when a3a was used as the transducing lysate was 3.6 × 10⁻³/cell/generation. The other markers showed similar rates of segregation. The Trp⁺ transductants were not cured by treatment with acriflavine. The segregation of *trp*⁻ was not correlated with a change in phage characteristics.

Phage characteristics of transductants. The phage characteristics of twenty Trp⁺ transductants (obtained with an a3a lysate at a m.o.i. of 1) which were purified by four cycles of cloning on minimal agar, could be divided into eight types (Table 2). Twelve of the transductants were either sensitive (type 1) or resistant (type 2) to a3a and did not liberate phage spontaneously or after u.v. induction. Two of the transductant colonies gave rise to both sensitive and resistant strains (types 3 and 6). An interesting observation was the isolation of resistant and sensitive strains which showed spontaneous phage liberation (types 5, 6, 7 and 8). The transductant colonies (types 5, 6, 7 and 8) which showed resistance to phage a3a gave rise to clones which released phage erratically but more frequently than the sensitive cryptic...
lysogen (Thomson & Woods, 1974) which was used as the recipient. The sensitive transducer (type 8) continually released phage and was easily inducible by ageing and u.v. irradiation, in contrast to the parent cryptic lysogen which is only inducible by NTG (Thomson & Woods, 1974). None of the resistant transductant clones were u.v. inducible.

Although the spontaneously liberated phage lysates and the induced lysate were tested for high frequency transduction (h.f.t.) of trp+, no h.f.t. preparations were detected.

Some of the transductant clones were initially resistant to a3a but after three or four clonings became semi-sensitive (types 4 and 7). The e.o.p. of a3a by these semi-sensitive colonies varied between $10^{-1}$ and $10^{-2}$. The decrease in e.o.p. was investigated to determine whether the phenomenon was due to restriction and modification. Plaques on a lawn of semi-sensitive cells were stabbed and the phage titre estimated on both the semi-sensitive strain and the wild-type bacterium. The semi-sensitive strain still showed the same reduction in e.o.p. as was obtained with the original a3a suspension, indicating that a3a had not been modified by growth on the semi-sensitive strain.

Host range of spontaneously liberated phage. The host range of each of the five spontaneously liberated phages was greater and differed from that of a3a. The spontaneously liberated phages were able to form plaques on the strains which were sensitive or semi-sensitive to a3a (Table 2) as well as on some strains which were resistant to a3a. Each of the phages liberated after transduction was able to infect the respective transductants from which they originated. All the liberated phages infected the clones from transductants 17 and 14 which were resistant to a3a. Furthermore, phage from transductants 16, 17 and 18 formed plaques on all the clones from transductants 10, 15 and 16. The phage from transductant 20 infected the clones from transductant 9.

Isolation of double lysogens in complete and minimal medium. The ten sensitive Achromobacter sp. 2 colonies, which were inoculated into complete broth containing phage a3a, resulted in cultures which were resistant to a3a. Clones isolated from these cultures were both resistant to a3a and u.v. inducible, and were therefore double lysogens (Thomson & Woods, 1974). These ten colonies were also inoculated into minimal broth containing a3a but formed cultures which were still sensitive to a3a. No resistant colonies or u.v.-inducible double lysogens were isolated from the minimal-medium cultures.

Thomson & Woods (1974) reported the isolation of double lysogens from colonies from the centre of turbid plaques on complete-medium double-agar-layer plates. In contrast, the plaques formed on Achromobacter sp. 2 by a3a on predominantly minimal-medium double-agar-layer plates (35 ml minimal agar bottom layer with a 2.5 ml soft complete-medium overlay) were completely clear, indicating that no resistant double lysogens had formed.

DISCUSSION

As the a phages all transduced a number of auxotrophic markers they were capable of generalized transduction. Generalized transductants are usually stable (Zinder, 1953) but Achromobacter transductants segregated auxotrophs. The transductants were not cured by treatment with acriflavine, suggesting that the transducing DNA did not remain as a plasmid in the recipient.

The range of phage characteristics and the spontaneous liberation of phage by purified transductant colonies is interesting, and was not due to either contamination by a3a or incomplete purification of the transductants. Four of the phages were liberated by strains resistant to a3a. All the released phages showed a different host range from a3a and were able to infect some strains which were resistant to a3a. The a3a-sensitive strain which re-
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leased phage was inducible by u.v. and ageing. The transduction system involved a cryptic lysogen (Thomson & Woods, 1974) and a hypothesis is suggested to explain the range of phage characteristics of the transductants. It is envisaged that the generalized transducing particles contain bacterial DNA as well as different amounts of phage DNA. The cryptic lysogen is characterized by defective immunity and excision systems, and complementation between the resident cryptic prophage and phage genes on the transducing DNA could account for the range of phage characteristics of the transductants and the release of recombinant phage types. Luria et al. (1960) and Chakrabarty & Gunsalus (1969) isolated transducing P1 and pf16h2 particles containing varying amounts of phage genes.

None of the transductants showed the characteristics of normal double lysogens which are resistant to α and u.v. inducible (Thomson & Woods, 1974). This suggests that only segments of the phage genome were available for complementation with the cryptic prophage and argues against the phage characteristics of the transductants being due to double lysogeny by contaminating non-transducing particles during transduction. Furthermore, double lysogens have only been isolated in complete medium, and attempts to demonstrate double lysogens in both liquid and agar minimal media were unsuccessful. This suggests that double lysogens cannot form in minimal media. The transductants were selected, purified and maintained on minimal medium.

As yet no h.f.t. lysates have been isolated. Watanabe, Ogata, Chan & Botstein (1972) obtained only three h.f.t. lysates by u.v. induction of 26 independently isolated transductants. Therefore the screening of more transductants might result in the isolation of an h.f.t. lysate.

There have been reports of generalized transducing phages causing transduction either by integration or lysogenization. The choice depends on the degree of homology between the bacterial DNA in the transducing particles and the recipient strain, as well as on the amount of residual phage DNA in the transducing particles (Luria et al. 1960; Chakrabarty & Gunsalus, 1969). However, although the phages worked with, P1 and pf16h2, are capable of generalized transduction, DNA incorporation by lysogenization was in both cases shown only for a specific set of genes. With P1, lac and with pf16h2, mandelate genes were transduced. Thus the α system is the first to show generalized transduction by a method other than the normal integration for a number of different markers.

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