Transduction of Swarming in Proteus mirabilis

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SUMMARY

A phage able to transduce a streptomycin resistance marker in Proteus mirabilis can also separately transduce the swarming characteristic between variants of two strains of P. mirabilis. Motile non-swarming variants were made to swarm on agar by incorporation of a swarming gene via a phage lysate of a swarming strain. The production of swarms by motile non-swarming variants when treated with phage lysates of other phenotypically similar strains indicated that at least three non-homologous factors control swarming. An O variant could also act as a donor of the swarming gene. This is because the O strain possesses an intact swarming centre which is masked by the absence of active flagella. This O variant was transduced to swarming by phage lysates of motile non-swarming strains or of a swarming strain. The gene transduced here was concerned with the presence of flagella. Factors controlling two morphological varieties of swarming were separately transduced to suitable recipients and a locus able to modify wild-type swarms was identified.

INTRODUCTION

Many organisms may swarm under suitable conditions. Thus Salmonella (Stocker, Zinder & Lederberg, 1953), Escherichia coli (Furness, 1958) and members of the genus Bacillus (Brown, Cherry, Moody & Gordon, 1955) swarm on semi-solid agar. Proteus morganii swarms on 1% agar at 20–28° (Rauss, 1936) and members of the genus Clostridium may swarm, particularly on moist media (Fildes, 1925). The A and C phases (Belyavin, 1951) of P. hauseri are able to swarm vigorously on well-dried nutrient agar.

Many explanations for the swarming of Proteus hauseri have been given (Lominski & Lendrum, 1947), but the phenomenon is now generally attributed to negative chemotaxis exerted by toxins which form during the growth of the culture (Lominski & Lendrum, 1947; Hughes, 1957). These toxins have not been identified. As regards the mechanism of swarming it is agreed (Topley & Wilson, 1955) that non-motile organisms do not swarm; active flagella are a prerequisite. Coetzee & Sacks (1960a) reported that certain temperate Proteus phages could transduce a streptomycin resistance marker. Swarming and non-swarming variants of some of the strains used were available (Coetzee, 1959; Coetzee & Sacks, 1960b), and it was decided to apply transduction techniques to the study of the phenomenon of swarming in P. mirabilis.
METHODS

Media. The composition of the broth and the MacConkey-type agar employed have been described (Coetzee & Sacks, 1960b). In addition, a nutrient agar made and used routinely by the Department was used. In some experiments to detect abortive transductants the agar concentration in media was reduced to 0.6% (w/v). Plates were well dried before use. The incubation temperature was 37°C.

Organisms. Variants of *Proteus mirabilis* strains 13 and 193 were used (Coetzee, 1959; Coetzee & Sacks, 1960a, b). These were 13-Y, 13-W, 13-Z, 13-O, 13-Y *str-r*, 193-Y, 193-W_4_, 193-W_15_. After 12 hr. incubation organism 13-Y has swarmed vigorously across nutrient agar or MacConkey plates in two or three bounds (Pl. 1, fig. 1). Variant 13-Z swarms as vigorously but in one continuous sheet of growth (Pl. 1, fig. 2). The Y and Z variants correspond respectively to the A and C phases of Belyavin (1951). Organism 193-Y swarms in compact rings and only covers half the area of a plate in the same period of time (Pl. 1, fig. 3). It hardly swarms at all on MacConkey agar. Agglutinin absorption tests (Coetzee, unpublished) showed that the H antigens of organisms 13 and 193 were identical while the O antigens differed. This difference could possibly be reflected in the fact that strain 13 is lysed by Proteus phages 68, 78 (Coetzee, 1958), while strain 193 lacks these phage receptors. The W forms of these two strains are motile and non-swarming and correspond closely to the B phase described by Belyavin (1951). They agglutinate to titre with 13-Y or 193-Y H sera. O variants—i.e. non-motile strains which do not agglutinate with H sera (Friewer & Leifson, 1952) are rare in these strains and only one (13-O) has been isolated. The W variants mutate to corresponding Y forms at low rates (Coetzee & Sacks, 1960b) and the O variant does the same (Coetzee, unpublished). Strains were stored on nutrient agar slopes at 4°C.

Phage. Phage 34/13 (Coetzee & Sacks, 1960a) was used. Phage lysates were prepared, sterilized and titrated according to methods described in the same paper. Grown on organism 13 it has a relative efficiency of plating of unity on strain 193.

Transduction techniques. The recipient organism was washed off a stock agar slope with broth and a suspension containing c. 1 x 10^8 organisms/ml made. This was done by visual comparison with a standard suspension. One ml. was then centrifuged, the supernatant discarded and the deposit mixed with 2 ml of phage lysate of the donor strain (plaque-forming titre c. 7 x 10^9/ml.). The mixture was incubated for 45 min. Loopfuls of the contents were then streaked on both types of solid media and the plates incubated for 15 hr. In some experiments the mixtures were diluted with broth before streaking, while in others the tubes were centrifuged and the deposits streaked out as before. Controls included a lysate sterility control and a recipient control in the form of a lysate prepared on the recipient organism and mixed with that organism as above. As phage lysates of donor organisms might perhaps contain swarm-inducing toxins for the recipient (Hughes, 1957) an additional control was used. This consisted of the supernatant obtained by centrifuging a 4 hr. growth of the donor organism in soft top-layer agar used in lysate preparation, sterilized with chloroform, as were phage lysates, and mixed with the recipient organism as described above. Lysates which caused swarming of recipient organisms were treated with deoxyribonuclease and anti-phage serum as previously described (Coetzee & Sacks, 1960a). Streptomycin resistance was transduced as described.
in the same paper. Methods used to measure phage adsorption and to detect lysogeny in transductants were also described in the paper. The character of swarms which appeared in these experiments was investigated by subculturing them on to fresh agar plates.

*Serological techniques.* These were as described for Proteus by Kauffmann (1951).

**RESULTS**

*Transduction studies.* Phage controls were always sterile. The two recipient controls usually showed no swarms after 15 hr. incubation (Pl. 2, fig. 4) but often produced a couple of swarms after a further 12-24 hr. No differences were noted between the two types of recipient controls. A summary of experiments performed and the nature of the swarms evoked from recipient organisms by phage lysates is presented in Table 1. Numerous swarms were evoked from variants 13-O1, 13-W1, and 13-W7, by treatment with phage lysates of 13-Y or 13-Y str-r (Pl. 2, figs. 5, 6). No quantitative work was done, but the number of swarms produced by the above systems appeared to be of a similar order—about 20/streak. All the swarms had the 13-Y (broad zone) mode of spread. The swarm-inducing abilities of these lysates and all other lysates used in this investigation were not affected by prior treatment with deoxyribonuclease but were completely abolished by an exposure to homologous phage antiserum which reduced the plaque-forming titre below $10^6$/ml. It is concluded that the phenomena presented are examples of phage-mediated transduction. These swarms and all other transductant swarms investigated were lysogenic for strain 13 and did not adsorb phage 34/13. This is attributed to lysogenic conversion (Coetzee, 1961). Phage lysates of 13-Y str-r were also capable of transducing resistance to 1000$\mu$g. streptomycin/ml. in these variants (Coetzee & Sacks, 1960a). These two markers are apparently not very closely linked; for no joint transductants were observed amongst 200 swarms and 300 streptomycin-resistant colonies examined. Swarms of the 13-Y (broad zone) variety were evoked from variants 13-W1, 13-W7 by a phage lysate of 13-O1. It would appear that 13-O1 cannot swarm because it has lost a gene determining the presence of flagella. It retains the gene for swarming and this gene can be transduced from it to variants 13-W1 and 13-W7, which are motile but do not swarm. Numerous swarms were also evoked when 13-O1 was treated with phage lysates of 13-W1 or 13-W7. It is probable

<table>
<thead>
<tr>
<th>Donor</th>
<th>Recipient</th>
<th>Type of swarm evoked</th>
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<tbody>
<tr>
<td>13-Y</td>
<td>13-O1, 13-W1, 13-W7, 193-W4, 193-W15</td>
<td>13-Y</td>
</tr>
<tr>
<td>13-O1</td>
<td>13-W1, 13-W7, 193-W4, 193-W15</td>
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<td>13-W7</td>
<td>13-O1, 13-W1, 193-W4, 193-W15</td>
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<td>13-Z</td>
<td>13-O1, 13-W1, 193-W7, 193-W4, 193-W15</td>
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<td>13-Z</td>
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that in the first series as well as in the latter experiments a gene concerned with flagellation was transmitted from 13-Y, 13-W₁, 13-W₁⁻ to 13-0₁. Swarms were evoked by phage lysates of 13-Y, 13-O₁, 13-W₁, 13-W₁⁻ from variants 193-W₄ and 193-W₁₅. The swarms were all of the 13-Y (broad zone) type. Recipient controls showed a few swarms after 24 hr. incubation but these were all of the 193-Y (compact) type. In years of work with strain 193 it has never swarmed in any other way than the tight-ring form previously described (Pl. 1, fig. 3). The complexity of the swarming locus was further demonstrated by the finding that swarms of the 13-Y (broad zone) variety were elicited from variant 13-W₁ by treatment with phage lysate 13-W₁⁻ and vice versa. The reason why all the swarms evoked from 193-W recipients by phage lysates of strain 13 and its variants should be of the 13-Y (broad zone) and not the 193-Y (compact) variety may be due to the proximity of the 193 swarming-type locus and the locus responsible for loss of swarming in the two recipients, so that both loci are replaced by their alleles present on the transducing fragment. No more W variants of strain 193 were available to test this hypothesis and it was not possible (see Discussion) to test the interaction of variants 193-W₄ and 193-W₁₅. However, the relationship between the motile non-swarming variants means that the swarming locus in these strains is composed of at least three sites. Allogenic transformations in pneumococci are well known (Jackson, 1962) and transductions of a similar nature were first described amongst a group of galactose-non-fermenting mutants in Salmonella typhimurium (Zinder & Lederberg, 1952). Organisms 13-O₁, 13-W₁, 13-W₁⁻, 193-W₄, 193-W₁₅ were treated with phage lysates of organism 18-Z. Numerous swarms were produced but they were all of the 13-Y (broad zone) type. In case the transduction rate of the Z type of swarming was very low the transduction mixtures were centrifuged and the deposits streaked. Again only 13-Y type swarms were detected. The following hypothesis may explain these results. Swarming of Proteus hauseri is normally (Coetzee & Sacks, 1960b) of the discontinuous 13-Y (broad zone) or 193-Y (compact) type. This swarming is controlled by a swarming locus and the 13-Y and 193-Y types are controlled by alleles \( Y^+ \) and \( Y^{+13} \) (Clowes, 1960). The Z-type continuous swarming is produced when a modifier gene (the Z gene) mutates in a \( Y^f \) genome. This Z locus is not closely linked to the swarmer locus \( Y \) or to the locus (or loci) controlling the presence of flagella. When a 13-Z lysate is applied to 13-O₁ transductants acquiring the flagellar gene are selected and the newly flagellated organism then swarms because it possesses a latent swarming locus. When the 18-Z lysate is applied to any of the W variants transductants acquiring the swarming gene are selected, and the resultant swarms therefore manifest the \( Y \) character of the donor. Independent incorporation of the \( Z^+ \) factor remains undetected because the \( W \) recipients have defective swarming loci. This hypothesis was tested by making use of the very restricted swarming of 193-Y on MacConkey agar. A transduction experiment was done with 193-Y as recipient and a donor lysate from 13-Z (Table 1). Controls on MacConkey plates showed the restricted 193-Y swarming. The test plates had numerous large swarms after 12 hr. incubation. The periphery of 50 discrete swarms were subcultured on to nutrient agar and incubated overnight. Twenty-nine of the swarms were of the 18-Y (broad zone) type and 21 had the 13-Z (continuous sheet) character. In this experiment both the flagellation and swarming loci of the recipient organism were intact. If the recipient incorporated the exogenous swarming gene then 18-Y type swarms
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resulted; when the Z+ locus of the donor was incorporated, the swarming of 198-Y was modified to the Z pattern.

No trails of colonies (Stocker et al. 1953; Stocker, 1956) at a distance from the original streaks were detected even with the use of semi-solid media in the transduction of motility to variant 13-O1. Trails could be expected if abortive transductions to swarming occurred. Because the swarms may have obscured such trails and because abortive transductions are almost always more numerous than the complete form (Ozeki, 1956) transduction mixtures were diluted before streaking. This had the effect of reducing the number of swarms/streak, but no trails of colonies were seen.

Serology. Organisms 198-W4 and 193-W15 transduced to the 13-Y type of swarming by respectively phage lysates 13-Y and 13-O1, and organism 13-O1 transduced to swarming by a lysate from 13-Y were examined serologically. O and H sera were prepared and agglutinin absorption tests done to detect antigenic differences between the transductants or between the latter and their parent organisms, 193-Y and 13-Y. All the H antigens proved identical. The O antigens of the 13-Y type transductants of 193-W4 and 193-W15 were identical but the O sera against these strains contained antibodies which were not absorbed by strains 193 or 13. Similarly, an O serum against the swarming transductant derived from 13-O1, possessed antibodies which were not absorbed by strains 193 or 13. The differences between the transductants and their parent organisms were thought to result from lysogenic conversion (Coetzee, 1961).

DISCUSSION

Stocker et al. (1953) have shown that a number of distinct genes control the presence, antigenic structure and functions of flagella in Salmonella. Through lack of non-motile flagellated (paralysed) variants (Friewer & Leifson, 1952) it was not possible to identify separate loci controlling the presence of flagella and their motility-conferring properties in this investigation. Having only one O variant also made it impossible to examine the possible complexity of the flagellation locus. A nice distinction was however made between the latter locus (or loci) and the swarming locus, which was found to consist of at least three sites. It was not possible to determine whether the swarming and flagella loci were linked. The Y (ring swarming) variant is the wild type of Proteus hauseri (Coetzee & Sacks, 1960b), and this type of swarming is controlled by the swarming locus. Because phage lysates of 13-Z (continuous swarming) transduce either the 13-Y or the 13-Z type swarming to suitable recipients, the 13-Z phenotype is thought to result from the action of a modifier gene (Z+) on the swarming locus. These two factors are not linked closely enough to be transduced together. Bryan (1961) described a modifier gene in Pneumococci whose only detectable expression was the enhancement of activity of specific streptomycin-resistance mutations.

No experiments are reported with the use of phage lysates of variants of organism 193, because phage 34/13 grown on 198 has an e.o.p. of 10^-3 on organism 13. The plaques are clear. Clear plaques are also formed on all variants of 198 and no transductions have been observed. This change appears to result from a phenotypic modification of the phage (Luria, 1958) but has not been extensively studied. Thorne (1962) described a Bacillus subtilis transducing system in which the transducing phage could not be propagated on the recipient organism.
No evidence for the occurrence of abortive transductions of swarming was encountered. This is not an exceptional finding as this type of transduction has only been reported in Salmonella (see Hartman, Hartman & Serman, 1960) and Escherichia coli systems (Gross & Englesberg, 1959) and not in Pseudomonas aeruginosa (Holloway & Monk, 1959), Staphylococcus aureus (Edgar & Stocker, 1961) or Bacillus subtilis (Thorne, 1962) transductions.

Because the flagellar antigens of organisms 193 and 13 are identical it has not been possible to analyse the genes which control H antigenic specificity. The system employing phage 12/57 and Proteus mirabilis strains 57 and 118 (Coetzee & Sacks, 1960a) would possibly have been more rewarding as these strains differed in somatic and flagellar antigens (Coetzee, unpublished) but strain 118 has been lost. Initially phage 12/57 was used. It too is capable of transducing the swarming character between Y and W variants of strain 57 but, as no other variants of this strain could be selected, the present system was chosen. Indeed the paucity of variants and the very restricted host-ranges of the transducing phages are serious obstacles in the transduction studies with P. hauseri.

This work has shown that phage 84/13 can independently transduce a streptomycin-resistance marker, a flagellation marker, the swarming-type and Z markers. It may now be classed as a phage capable of general transduction.

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REFERENCES


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EXPLANATION OF PLATES

(All figures life size.)

PLATE 1

Fig. 1. Organism 13-Y inoculated at centre of nutrient agar plate and incubated for 15 hr. at 37°.

Fig. 2. Organism 13-Z inoculated at centre of nutrient agar plate and incubated for 15 hr. at 37°.

Fig. 3. Organism 193-Y inoculated at centre of nutrient agar plate and incubated for 15 hr. at 37°.

PLATE 2

Fig. 4. A 13-W1 recipient control on nutrient agar. No swarms present after 15 hr. at 37°.

Fig. 5. Organism 13-O1 treated with a 13-Y phage lysate. Many small discrete swarms and a few large swarms are present on MacConkey agar after 15 hr. at 37°.

Fig. 6. Organism 13-W1 treated with a phage lysate of 13-Y. Swarms have merged but their points of origin can be seen. Nutrient agar after 12 hr. at 37°.