Formation of the Vegetative Pool by Induced Defective and Healthy Lysogenic Strains of Escherichia coli

BY J. F. WHITFIELD AND R. K. APPLEYARD

Biology Division, Atomic Energy of Canada Ltd., Chalk River, Ontario, Canada

SUMMARY: Comparison of two defective λ prophages carried by strains of Escherichia coli, with respect to their physiological properties following induction, has revealed that one defective mutant (λ i2) can multiply normally as vegetative phage but cannot mature into normal infectious phage, whereas the other defective phage (λ i1) can undergo only limited multiplication since it is unstable in the vegetative state. Consideration of the properties of λ i1 has led to an hypothesis that radiation induction causes the prophage to produce a vegetative replica which becomes autonomous rather than conversion of the prophage into vegetative phage.

Lysogenic strains of Escherichia coli containing defective forms of the bacteriophage λ have been investigated by Appleyard (1954, 1956) and by Jacob & Wollman (1956). They are distinguished by an almost total failure to liberate healthy phage following induction, and Appleyard (1956) showed that the defect may be transmitted by extracellular defective phage. The phage genes responsible for the defects resemble the recessive lethal genes of higher organisms; induction by ultraviolet irradiation of a doubly lysogenic strain, in which a normal and a defective prophage are present in the same cell, is followed by liberation at lysis of healthy phage particles, some of which have acquired genes from the defective prophage by recombination during the latent period.

As with recessive lethal genes, phage defects of independent origin would only rarely be expected to be genetically or physiologically identical. Appleyard & Whitfield (1956) demonstrated recombination between two such defects to give rise to normal progeny. Therefore, they are not allelic but are genetically distinct factors. In the present study the physiological effects of these defects have been compared.

METHODS

Bacteria and bacteriophages

We used as experimental material strains of Escherichia coli and bacteriophages described in earlier papers from this laboratory (Appleyard, 1956; Appleyard, McGregor & Baird, 1956). The two defects of independent origin chosen for investigation were those originally isolated in strains C33 (λ i2) and C60 (λ i1) (Appleyard, 1954). Both defects can be transmitted through extracellular phage to the same sensitive bacterium (Appleyard, 1956) with the advantage that observed physiological differences cannot then be due to strain differences between the bacterial hosts in which the defects were originally found. The cultures C112 (λ i1) and C112 (λ i2) as well as the healthy lysogen...
C112 (λ s) used in our experiments were prepared by artificial lysogenization of the same sensitive strain C112, a descendant of strain 12-112 of Wollman (1953). Bacteriophage λ-s was chosen as the healthy control phage, since both it and λ i1 and λ i2 were derived from the same ancestral strain of E. coli K12 (λ).

Production of healthy phage by ultraviolet-stimulated reverse mutation

If the defects arise as mutations of phage genes, many of them should be reversible through 'back mutation'. To study the frequency of u.v. stimulated reversion at various steps of phage development, a culture of defective lysogenic organisms was first induced at time zero by an exposure to a small dose of u.v. radiation. The induced organisms were allowed to develop and at various times during development several portions of the induced culture were exposed to different doses of u.v. radiation and after each dose the bacteria yielding healthy phage were counted. At any given time, the slope of the mutation/dose curve near the origin was taken as a measure of mutability. The detailed procedure was as follows.

Defective lysogens were prepared for induction by one of two methods:

1. Organisms were grown on the surface of nutrient agar plates and, while still in the logarithmic phase of growth, were washed off in ice-cold buffer and diluted to a turbidity value (nephelometrically determined) which corresponded to a concentration of $3 \times 10^8$ viable organisms/ml.

2. Organisms were grown in liquid Tryptone (Difco) broth with aeration to a concentration of $3 \times 10^8$ viable organisms/ml. They were then separated from the broth by centrifugation at $4^\circ$ for 3 min. at 8000 g and resuspended in an equal volume of ice-cold buffer. In either case the organisms were at once exposed to u.v. radiation (2537 Å) for 15 sec. (225 ergs/mm.2) to induce them. Tryptone was added to a concentration of 1% (w/v) (1 ml. of a 10% (w/v) solution of Tryptone to 9 ml. suspension of organisms) and the culture was aerated at $37^\circ$.

At each time to be investigated, the organisms in 5 ml. of the culture were spun down and resuspended in an equal volume of ice-cold buffer as before. Separate 1 ml. portions of this buffer suspension were exposed to a further dose of u.v. radiation usually for 20, 40 and 60 sec., and were at once assayed, with an unirradiated control, as plaque-formers before lysis by the agar layer technique.

Recombinational analysis of pool size

It is possible to superinfect an induced lysogenic organism with a phage which has been suitably marked genetically, and to relate the number of superinfecting normal phage/induced lysogenic organism and the proportion of recombinant phage released at lysis to the genetic or vegetative pool size (Visconti & Delbrück, 1953) in the induced lysogenic organism at the time of superinfection. This relation will be described in full in the experimental section of the present paper. To determine the proportion of recombinants produced by superinfection following induction, cultures of C112 (λ s), C112
(λ i1) and C112 (λ i2) were grown to $3 \times 10^8$ viable organisms/ml in Tryptone broth. Ten ml culture were spun down at 7000 g. The organisms were resuspended in 9 ml buffer and u.v.-irradiated (225 ergs/mm$^2$). One ml. of a 10% (w/v) solution of Tryptone in distilled water was added to the induced culture, which was then aerated at 37°. After various time intervals, 0.9 ml samples of the culture were removed and transferred to a tube containing 0.1 ml at various titres of phage $\lambda v1 h$ (clear-plaque former) a host-range mutant of the weak virulent $\lambda$ phage (Appleyard, 1954; Appleyard et al. 1956). Beginning at any time between 0 and 45 min. after induction of the lysogenic host organisms 99.9% of the phage $\lambda v1 h$ was adsorbed within 15 min. at 37°. This fraction remained the same in mixtures containing up to 60 phage particles/organism.

Each infected culture was aerated until 30 min. after the appearance of visible lysis. The lysates were then shaken with 0.1 ml chloroform/ml lysate, diluted 1/100 in broth and assayed by the double layer method on plates containing medium to a depth of not less than 1 cm. (Jacob & Wollman, 1954) to determine the numbers of turbid (recombinant) and clear plaques (normal parental phenotype).

In order to study changes in the normal healthy lysogenic strain of Escherichia coli C112 (λ s) following induction, a slightly different procedure was followed. The lysates were diluted and mixed with sufficient organisms from a freshly grown culture of the sensitive strain CR600 (Appleyard et al. 1956) to obtain a phage to bacterium ratio of at most 1:100. The infected culture was plated before lysis on the host-range indicator strain E. coli CR63. This method of pre-adsorption eliminates any distortion of the data due to phenotypic mixing (Appleyard et al. 1956) as well as improving the clarity and size of the plaques. A phage to bacterium ratio of 1:100 was used to avoid double infection, which would also distort the results.

RESULTS

Physiological comparisons between lysogens carrying defects i1 and i2

The defects show superficial similarities. Both are transferable by infection and can therefore be studied in bacteria having otherwise identical genotypes. Mass lysis of bacteria carrying either defective prophage can be induced by small doses of u.v. radiation. It occurs 85 min. after exposure to u.v. light with liberation of an average of one healthy phage/10$^5$ to 10$^6$ induced organisms. In both defective lysogens the production of infective phage, i.e. 'reverse mutants', can be stimulated by exposure to further doses of u.v. radiation during development, an average of 6 to 10 phage particles being released from each exposed organism.

Differences in frequency of u.v.-induced reversions of i to i+

A marked difference between the two defects was observed with respect to frequency of reverse mutant phage detected following the double irradiation technique described above (Fig. 1). Following induction of organisms of
Escherichia coli C112 (λ i2) in the logarithmic phase of growth, the frequency of reversions was over 10 times as great when the second irradiation was applied at 30 min. as when the second irradiation was applied at 0 min. (Fig. 1A). After 30 min., the frequency dropped sharply. In log phase organisms of C112 (λ i1), no such increase was observed (Fig. 1A). Although the variation in time of the u.v. stimulation was somewhat different for older cultures (phase of retarded growth) the marked contrast in behaviour between i1 and i2 remained (Fig. 1B). These observations suggested to us that the number of λ i2 vegetative phage units increases with time following induction, whereas the number of λ i1 vegetative phage units does not. To investigate this possibility it was necessary to devise ways of measuring the size of the vegetative or genetic pool in the induced defective and normal lysogens.

Recombinational estimate of pool size

Principles of experiment. By the genetic or vegetative pool we mean the total number of actively-recombining replicas of the phage genotype inside the organism (Visconti & Delbrück, 1953). The simplest method of estimating the pool size is to measure the number of recombining units by some kind of superinfection with a genetically marked phage, and experiments on these lines have been carried out by a number of workers (Levinthal & Visconti, 1953; Visconti & Garen, 1953; Jacob & Wollman, 1956). It is clear from the

Fig. 1. The variation with time of the u.v. stimulated production of healthy phage following induction of Escherichia coli C112 (λ i1) (○—○) and E. coli C112 (λ i2) (●—●), (A) from cultures in the logarithmic phase of growth, (B) from cultures in the retarded phase of growth.
Defective lysogeny

work of Jacob (1954), as well as from our own experiments, that the recombinational method includes the prophages in its measure of the pool size. It therefore strictly measures the genetic pool size rather than the vegetative. It was, however, necessary to avoid possible complications connected with selection for or against recombinants, with variations of the recombination process as a function either of time of irradiation (Jacob & Wollman, 1955) or of the physiological state of the organisms in which defective development was proceeding. Moreover, we were unable to measure the total yield because it included defective phage. We therefore used a variation of the simple method based on the following argument.

Let $m$ be the number of defective, recombinable phage replicas/induced organism and let $n$ be the number of superinfecting healthy phage particles/organism. If the ratio of $m$ to $n$ be constant throughout the remainder of the latent period, then the number of recombinants, $N$, issuing from such an infection should be directly proportional to the fraction of matings in which recombination is possible. If matings are at random, this fraction is $2mn/(m+n)^2$.

Then,

$$N = \frac{2Kmn}{(m+n)^2}.$$  (1)

Since the numbers of reciprocal recombinants are statistically equal in infections with $\lambda$ phage (Jacob & Wollman, 1954; Wollman & Jacob, 1954), the fraction of viable recombinants in the yield will be some constant fraction of $N$

$$N' = \frac{K'mn}{(m+n)^2},$$  (2)

whereas the total fraction of viable phage particles will be $n/(m+n)$.

Therefore, the proportion, $p$, of recombinants among the total yield of active phage particles will be

$$p = \frac{K'm}{(m+n)}.$$  (3)

(3) may be rewritten as a linear relation between the observable quantities $(np)$ and $p$:

$$np = -mp + K'm.$$  (4)

By comparing different multiplicities of superinfection, $K'$ may be eliminated

$$\frac{p}{p'} = \frac{m+n'}{m+n}.$$  (5)

To determine the pool size in induced defective lysogens, we studied recombination between the clear-plaque marker, $v1$, and the defect $i1$ or $i2$ when the superinfecting phage was $v1\ 1^+\ i2^+$ and the carried phage $v1^+\ i1\ i2^+$ or $v1^+\ i1^+\ i2$; in such an experiment the only viable recombinant is $v1^+\ i1^+\ i2^+$. The complementary recombinant $v1\ i1$ or $v1\ i2$ is not scorable. To determine the pool size in induced healthy lysogens, we studied recombination between the markers $v1$ and $h$ when the superinfecting phage was $v1\ h$ and the carried
phage was $v1^+ h^+$. Scoring only one recombinant class $v1^+ h$ among the total $h$ phage in this case is analogous to the method of estimating the pool size in defective lysogens.

Validity of the method. The linearity of the relation (4) between $(np)$ and $p$ has been confirmed over a wide range of times and multiplicities of superinfection. A typical set of data is given in Table 1 and plotted as a test of linearity in Fig. 2. The straight lines derived from a number of other superinfections are plotted in Figs. 3 and 4.

Table 1. Calculation of the values $(np)$ and $p$ of text relation (4) using experimental data from superinfection of Escherichia coli C112 ($\lambda i^2$) with phage $\lambda v1 i^2+ h$ 15 min. after induction with u.v. light

<table>
<thead>
<tr>
<th>$n^*$</th>
<th>$v1^+ i^2+$</th>
<th>$v1 i^2+$</th>
<th>$p$</th>
<th>$(np)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.6</td>
<td>224</td>
<td>2763</td>
<td>0.075</td>
<td>0.110</td>
</tr>
<tr>
<td>4.7</td>
<td>221</td>
<td>3738</td>
<td>0.055</td>
<td>0.258</td>
</tr>
<tr>
<td>10.0</td>
<td>115</td>
<td>3184</td>
<td>0.035</td>
<td>0.350</td>
</tr>
<tr>
<td>21.2</td>
<td>140</td>
<td>6136</td>
<td>0.021</td>
<td>0.445</td>
</tr>
</tbody>
</table>

* In order to allow for the Poisson distribution of the infecting particles, the values of $n$ are calculated from the observed average multiplicity of infection, $x$, by the formula $n = x/(1 - e^{-x})$.

![Fig. 2. Test of the linearity of relation (4) in the text between $(np)$ and $p$. Data obtained from superinfection of Escherichia coli C112 ($\lambda i^2$) with $v1 i^2+ h$ at 15 min. after u.v. induction. The data and calculations are recorded in Table 1. This line is also line B of Fig. 4.]

Such straight lines as that of Fig. 2 have confirmed the validity of the assumptions concerning random mating and absence of selection upon which our estimates of pool size are based. They show that very high multiplicities of superinfecting phage can enter the induced cell and take part in recombi-
Defective lysogeny, in contrast to the situation in multiple infections of sensitive cells (Wollman & Jacob, 1954). Further, any selection for or against the superinfecting phage during multiplication should be a function of the extent of that multiplication and so, inversely, of the multiplicity of superinfection: hence it would give rise to curvature of the lines which is not observed. Loss of a constant appreciable fraction of the superinfecting phage, for example during penetration of the induced cell, is not excluded by the linearity and could give rise to a constant factor of error in the estimates of pool size. However, this is unlikely, in view of the good agreement of the pool size estimated shortly after induction (see Fig. 2 and below) with the number of prophages per cell estimated by other methods.

Measurements of pool size as a function of time

We have used the recombinational analysis just described to follow the size of the genetic pools as a function of time in the defective lysogens *Escherichia coli* C112 (\(\lambda i\)), C112 (\(\lambda i 2\)), and the healthy lysogen *E. coli* C112 (\(\lambda s\)). The results are shown in Figs. 5 and 6. The three strains behave differently and will be discussed in succession.

Healthy lysogen (Fig. 5, A). During the first 15 min. after infection the number of recombinable particles remained nearly constant, but by 35 min. it had increased from 8 to 20. By 40 min. the pool size had increased to 95 particles and there it remained up to 45 min. This increase with time is reflected in the difference between the slope of line A and lines B and C of Fig. 3. The observations of Weigle & Delbrück (1951), that the first intracellular mature phage particles appeared between 40 and 45 min. after induction have been shown to apply to lysogenic derivatives of *Escherichia coli*
of C112 under the conditions of our experiments (Appleyard, unpublished). We therefore attribute the levelling off of the pool size to the onset of maturation.

**Defective lysogen** *Escherichia coli* C112 (λ i2) (Fig. 5, B). The number of recombinable particles in induced *E. coli* C112 (λ i2) at first remained constant and then rose steeply as in *E. coli* C112 (λ s). However, instead of reaching a plateau it continued to rise to about 201 particles, twice the value in the healthy strain, and we were not able to detect any levelling off at the latest times at which we were able to make measurements. The rate of multiplication of phage λ i2 appeared to be about the same as that of phage λ s (compare Fig. 5, A B with 5, C). These results suggest that the maturation of λ i2 is impaired whereas its vegetative multiplication is normal.

![Fig. 5](image1.png)

**Fig. 5.** Variation of the pool size with time following induction. A: *Escherichia coli* C112 (λ s). B: *E. coli* C112 (λ i2). C: *E. coli* C112 (λ i1).

![Fig. 6](image2.png)

**Fig. 6.** Variation of the pool size with time following induction of *Escherichia coli* C112 (λ i1).

**Defective lysogen** *Escherichia coli* C112 (λ i1) (Figs. 5, C and 6 and Table 2). There were marked differences between the events in *E. coli* C112 (λ i1) and those in *E. coli* C112 (λ i2). The first multiplication of phage λ i1 took place between 25 and 30 min. after induction but was followed by a decline in numbers between 30 and 40 min. Two other experiments were performed to confirm this picture. In the second experiment the value of *m* was 14·6 at 30 min. and 16·3 at 35 min.; by 40 min. this number had declined to 8·8 and at 45 min. *m* had increased to 21·1. The results of these two experiments have been combined and presented in Fig. 5, C and Table 2.

Further analysis of the events described in the preceding paragraph was carried out. Samples of an induced culture of *Escherichia coli* C112 (λ i1)
Defective lysogeny

were superinfected at 2 min. intervals from 34 to 46 min. From the data presented in Fig. 6 it may be seen that as in the first two experiments the value of \( m \) between 34 and 36 min. was constant, or nearly so, at 10. However, by 38 min. \( m \) had decreased to 4.5, being followed by the usual rise, which in this case began between 40 and 42 min.

Table 2. *Calculation of the pool size (\( m \)) in cells of Escherichia coli C112 (\( \lambda i1 \)) superinfected with \( \lambda v1 i1^+ h \) at various times post-induction*

<table>
<thead>
<tr>
<th>Time post-induction (min.)</th>
<th>( n )</th>
<th>( v1^+ i1^+ )</th>
<th>( v1 i1^+ )</th>
<th>( p )</th>
<th>( m )</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>1.6</td>
<td>143</td>
<td>1547</td>
<td>0.085</td>
<td>5.5</td>
</tr>
<tr>
<td>10</td>
<td>1.6</td>
<td>342</td>
<td>4341</td>
<td>0.073</td>
<td>4.3</td>
</tr>
<tr>
<td>20</td>
<td>1.6</td>
<td>110</td>
<td>1218</td>
<td>0.083</td>
<td>4.8</td>
</tr>
<tr>
<td>25</td>
<td>1.6</td>
<td>156</td>
<td>1802</td>
<td>0.079</td>
<td>3.8</td>
</tr>
<tr>
<td>30</td>
<td>1.6</td>
<td>203</td>
<td>3146</td>
<td>0.061</td>
<td>12.0</td>
</tr>
<tr>
<td>10</td>
<td>206</td>
<td>4968</td>
<td></td>
<td>0.038</td>
<td></td>
</tr>
<tr>
<td>1.6</td>
<td>157</td>
<td>1662</td>
<td></td>
<td>0.086</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>92</td>
<td>1502</td>
<td></td>
<td>0.057</td>
<td></td>
</tr>
<tr>
<td>1.6</td>
<td>84</td>
<td>1062</td>
<td></td>
<td>0.073</td>
<td></td>
</tr>
<tr>
<td>5.2</td>
<td>115</td>
<td>1791</td>
<td></td>
<td>0.060</td>
<td></td>
</tr>
<tr>
<td>35</td>
<td>1.6</td>
<td>145</td>
<td>1737</td>
<td>0.077</td>
<td></td>
</tr>
<tr>
<td>7.0</td>
<td>302</td>
<td>4683</td>
<td></td>
<td>0.060</td>
<td></td>
</tr>
<tr>
<td>40</td>
<td>1.6</td>
<td>228</td>
<td>2605</td>
<td>0.080</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>46</td>
<td>1195</td>
<td></td>
<td>0.037</td>
<td></td>
</tr>
<tr>
<td>1.6</td>
<td>166</td>
<td>1746</td>
<td></td>
<td>0.086</td>
<td></td>
</tr>
<tr>
<td>7.0</td>
<td>123</td>
<td>2090</td>
<td></td>
<td>0.055</td>
<td></td>
</tr>
<tr>
<td>43</td>
<td>1.4</td>
<td>91</td>
<td>1520</td>
<td>0.056</td>
<td></td>
</tr>
<tr>
<td>5.0</td>
<td>131</td>
<td>3270</td>
<td></td>
<td>0.044</td>
<td></td>
</tr>
<tr>
<td>7.6</td>
<td>61</td>
<td>1503</td>
<td></td>
<td>0.038</td>
<td></td>
</tr>
<tr>
<td>18.5</td>
<td>66</td>
<td>2586</td>
<td></td>
<td>0.026</td>
<td></td>
</tr>
<tr>
<td>45</td>
<td>1.6</td>
<td>425</td>
<td>6734</td>
<td>0.059</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>168</td>
<td>3462</td>
<td></td>
<td>0.046</td>
<td></td>
</tr>
<tr>
<td>1.6</td>
<td>208</td>
<td>3335</td>
<td></td>
<td>0.058</td>
<td></td>
</tr>
<tr>
<td>8.0</td>
<td>50</td>
<td>1057</td>
<td></td>
<td>0.045</td>
<td></td>
</tr>
</tbody>
</table>

The difference between the pool sizes at 43–45 min. after induction of *Escherichia coli* C112 (\( \lambda i1 \)) and *E. coli* C112 (\( \lambda i2 \)) is reflected in the difference between the slope of line A and those of lines C and D of Fig. 4. It appears that phage \( \lambda i1 \) can undergo only limited multiplication in the vegetative state.

**DISCUSSION**

Measurements of pool size and u.v.-stimulated liberation of healthy phage show that defects \( i1 \) and \( i2 \) bring about different physiological changes following induction. In particular, estimates of genetic pool size show that
phage $\lambda i1$, unlike phage $\lambda i2$, experiences some difficulty in vegetative multiplication as was suggested by the curves of u.v.-stimulated reversions (Fig. 1, A). Comparison of the pool formed by each defective phage with that formed by healthy phage (Fig. 5) permits a more detailed interpretation of the nature of both defects.

The nature of defect $i2$. The characteristic difference between phage $\lambda i2$ and phage $\lambda s$ is the failure of the number of recombinable particles in the case of the induced $i2$ lysogen to stop rising and reach a plateau. Although the time at which the rapid increase in recombinable particles took place in *Escherichia coli* C112 ($\lambda i2$) was 5 min. earlier than that in *E. coli C112* ($\lambda s$), this is more likely to be correlated with later lysis of induced *E. coli C112* ($\lambda s$), than to be a characteristic of phage $\lambda i2$.

Weigle & Delbrück (1951) observed that the first mature infectious phages appeared in cells of induced healthy *Escherichia coli K12* ($\lambda$) between 40 and 45 min. after induction. This observation has been confirmed for our strains (Appleyard, unpublished). If the attainment of a constant pool size in the induced normal lysogenic organism, as in coliphage T2- or T4-infected organisms, is due to the intervention of maturation, then we can describe the $i2$ defect as a failure to undergo the first step of maturation, i.e. removal from the vegetative pool.

How much of the machinery of maturation can be set up by phage $\lambda i2$ remains unknown. However, Dr E. Kellenberger (personal communication) observed that empty phage heads and tails both appeared in lysates after induction of the strain C83 ($\lambda i2$). Whitfield, Appleyard & Baird (unpublished) prepared doubly lysogenic strains of *Escherichia coli C112* which carried both phage $\lambda i2$ and a host range ($h$) mutant of normal phage $\lambda$. Among the phage in lysates from such u.v.-induced strains, they observed healthy phage which were genotypically $h$ but which possessed $h^+$ coats derived from the defective phage. Possession of an $h$ coat permits the individual phage to absorb on to the host range indicator strain *E. coli CR63* while a phage particle with an $h^+$ coat cannot absorb on to this strain (Appleyard *et al.* 1956). Therefore, it seems that phage $\lambda i2$ can initiate construction of and at least in part control the specificity of the morphological components required for maturation.

Another defective $\lambda$ phage was studied by Jacob & Wollman (1956). They concluded that their defect prevents maturation of vegetative $\lambda$ phage. Since these authors were unable to demonstrate the presence of mature defective phage even after superinfection with healthy phage, they suggested that each defective vegetative unit interferes with its own maturation. However, in the presence of healthy phage, $\lambda i2$ can be matured (Appleyard, 1956). We therefore conclude that it belongs to a different class of maturation defects.

The nature of $i1$. In the prophage state, phage $\lambda i1$ is stable, for it can be transmitted through many cell generations without loss. In induced and spontaneous lysates from the double lysogenic strain *Escherichia coli C112* ($\lambda i1$, $\lambda Clh$), the major part of the progeny population is genotypically $h$ but phenotypically $h^+$. It would appear from this that phage $\lambda i1$, as is the case
with phage \( \lambda i2 \), can direct the synthesis of those elements which confer upon the mature virus its host range properties (Whitfield, Appleyard & Baird, unpublished).

In the vegetative state, phage \( \lambda i1 \) can multiply, since two successive bursts of multiplication are observed to follow induction. However, these two events are separated by an interval of decrease in the number of vegetative particles. Lambda-\( i1 \) phage is therefore unstable in the vegetative state. The present experiments do not enable us to distinguish clearly between a genuine physico-chemical instability of vegetative phage \( \lambda i1 \) and a cessation of multiplication coupled with some hypothetical normal process of removal from the vegetative pool. However, removal could not in this case be through a normal maturation process, as it begins too soon.

Some support for believing vegetative phage \( \lambda i1 \) to be unstable is provided by the observations of Appleyard & Whitfield (1956). They observed that superinfection immediately following u.v. irradiation of cultures of Escherichia coli C112 (\( \lambda i2 \)) with phage \( \lambda i1 \) resulted in the appearance of no healthy re-combinant phage-yielding organisms. Superinfection at later times produced increasing numbers of normal phage yielders. In the opposite situation, with phage \( \lambda i2 \) and E. coli C112 (\( \lambda i1 \)), superinfection immediately following u.v. irradiation resulted in the appearance of nearly the maximum number of normal phage-yielders possible under the conditions of the experiment. These observations suggest that phage \( \lambda i1 \), unable to maintain its integrity in the host cytoplasm, would have to enter the host at a later time in order to be present in an intact state to participate in recombination with phage \( \lambda i2 \) when that phage begins its vegetative multiplication.

Analyses of the deoxyribonucleic acid content of Escherichia coli C112 (\( \lambda i1 \)) cultures at various times after u.v.-induction indicate that the DNA content remains constant during the first 30 min. after induction but shortly thereafter increases linearly at the same rate as that in induced normal lysogenic cultures of E. coli C112 (\( \lambda s \)) (Whitfield, unpublished). It should be pointed out that the time of resumption of DNA synthesis following induction of these strains correlates well with the time at which the values of \( m \) begin to rise and also with the time at which the frequency of u.v.-stimulated reversions sharply increases in E. coli C112 (\( \lambda i2 \)). These biochemical studies, to be published later, were carried out under the same conditions as the pool-size estimations.

If the instability of phage \( \lambda i1 \) be due to extensive fragmentation resulting from the operation of some factor such as deoxyribonuclease, then large amounts of DNA fragments would appear in the acid-soluble nucleotide fractions from 30 min. to lysis of the culture. This was not observed. However, limited fragmentation of the vegetative particles remains a possibility.

During the decline the number of recombinable particles decreases not to zero, but to approximately the number preceding the first burst of multiplication; thus there seems to be a small persisting number of stable particles. We suggest that this persistent fraction represents the original prophages of the cell which by continued association with the bacterial nuclear material
retain their stability. This hypothesis implies that in the process of induction, prophage is not itself physically converted to vegetative phage, for all the intracellular particles would then be unstable, but makes at least one vegetative replica which initiates vegetative multiplication. In the case of the unstable phage λ i1, because the picture is not obliterated by large numbers of vegetative phage units, it can be seen to initiate two such attempts at vegetative multiplication, separated from each other and from the moment of induction by very approximately the division time of the un-irradiated lysogenic organisms.

The nature of u.v. radiation induction. We therefore propose the following hypothesis concerning the mechanism of u.v. radiation induction. The effect of the radiation is to modify the cell's nuclear apparatus in such a way that replicas formed by the prophage at its subsequent divisions become vegetative phage. It is of interest to relate this hypothesis in detail to a number of established facts. The work of Lwoff and his colleagues (Lwoff, 1953b) suggests the intervention of physiological processes in the induction process. Direct absorption of radiation energy by prophage is not needed to accomplish induction. We suggest that the effect of the radiation upon the nuclear apparatus is somewhat analogous to the mitotic delay well known as an effect of radiation upon the cells of higher organisms. DNA synthesis in both lysogenic and non-lysogenic microbial cells is inhibited by small doses of u.v. radiation (Siminovitch & Rapkine, 1951; Kelner, 1953; Barner & Cohen, 1956; Whitfield, unpublished), although various intermediates of DNA synthesis including deoxyribonucleotides appear as accumulated products in the medium (Kanazir, 1954; Kanazir & Errera, 1954). Such changes would prevent the multiplication of the bacterial nuclear apparatus. If the prophage DNA were not similarly affected, the fresh replica would find itself in a predominantly cytoplasmic milieu. The availability of accumulated DNA products would permit it to multiply as an autonomous unit. The unbalanced condition of the cell would thus greatly favour vegetative multiplication. Similar hypotheses have been proposed by Jacob & Stent (cited by Lwoff, 1953a).

Morse, Lederberg & Lederberg (1956) observed that in Escherichia coli transduction of gal markers by λ phage was effective only when the phage was obtained by induction of a lysogenic strain, and is not observed when the phage issued from a lytic infection. Their findings are most naturally explained if vegetative λ phage does not become attached to galactose loci but after induction the linkage of some units to gal markers which they initially had as prophages persists, as does the associated stability in our experiments.

The number of prophages/organism. Bertani (1953) and Jacob (1954) were able to obtain approximate calculations of the number of prophages/lysogenic bacterium. Both workers (Bertani with lysogenic Shigella dysenteriae and Jacob with lysogenic Pseudomonas aeruginosa) concluded that the number of prophages/bacterium was close to the number of nuclei. The present studies lead to the same conclusion. The number of prophages can be inferred from the number of recombinable particles at zero minutes after induction. This
Defective lysogeny

number ranges from 4.6 to 6.0 with an average of about 5 and so closely approximates the average number of nucleci/cell in the logarithmic phase of growth (Whitfield, unpublished).

The authors would like to express their gratitude to Mr K. M. Baird for his assistance in carrying out some of the experiments recorded in this paper.

REFERENCES

JACOB, F. & WOLLMAN, E. L. (1956). Recherches sur les bactériès lysogènes défec-
KELNER, A. (1953). Growth, respiration, and nucleic acid synthesis in ultraviolet-
irradiated and in photoreactivated Escherichia coli. J. Bact. 65, 252.
MORSE, M. L., LEDERBERG, E. M. & LEDERBERG, J. (1956). Transduction in Escher-
ichia coli K12. Genetics, 41, 142.


(Received 8 April 1957)