On the role of Cro in λ prophage induction

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The lysogenic state of bacteriophage λ is exceptionally stable yet the prophage is readily induced in response to DNA damage. This delicate epigenetic switch is believed to be regulated by two proteins; the lysogenic maintenance promoting protein CI and the early lytic protein Cro. First, we confirm, in the native configuration, the previous observation that the DNA loop mediated by oligomerization of CI bound to two distinct operator regions (O_L and O_R), increases repression of the early lytic promoters and is important for stable maintenance of lysogeny. Second, we show that the presence of the cro gene might be unimportant for the lysogenic to lytic switch during induction of the λ prophage. We revisit the idea that Cro’s primary role in induction is instead to mediate weak repression of the early lytic promoters.

CI protein | genetic switch | transcription

The λ prophage of Escherichia coli can escape lysogeny and enter lytic development by prophage induction (1). Induction is triggered by the host SOS response, which in turn is activated by damage to the host cell DNA. Thus, induction provides a way for the prophage to escape from a challenged or dying host. During lysogeny, the λ lytic genes are repressed by a phage-encoded repressor, the product of the cI gene (2). The lytic genes of λ are arranged in a sequential and temporal manner where the expression of one group of genes is required for expression of the next and henceforth. The CI repressor silences all of the lytic genes by preventing transcription from the two earliest lytic promoters. The regulatory region of promoter right (P_R) and promoter left (P_L) each contain an operator, operator right (O_R) and operator left (O_L), respectively, each consisting of three binding sites for CI (reviewed in ref. 3; see also Fig. 1B). In the lysogen, two CI dimers are bound cooperatively to O_R1/O_R2 and O_L1/O_L2 to prevent transcription from P_R and P_L. The CI dimer bound to O_R2 activates transcription from maintenance promoter (P_RM) by a direct protein–protein interaction with RNA polymerase (4–7). CI has a lower binding affinity for O_R3 relative to O_L3 and the other operators. Recent studies have shown that O_R3 and O_L3 are simultaneously occupied by CI in lysogens ∼60% of the time (8). CI binding to O_R3 excludes RNA polymerase from initiating transcription from P_RM, thus cI transcription is both positively and negatively autoregulated.

The repression of P_RM is enhanced by an O_L–CI–O_R complex, formed by octamerization of the CI dimers bound to O_R1/O_R2 and O_L1/O_L2 (refs. 9–11; see Fig. 1B); thus, the left operator region participates in the regulation of the right operator region and vice versa. The early lytic protein Cro, transcribed from P_R, antagonizes CI after prophage induction (12, 13). Cro binds to the same three sites at O_R and O_L as CI binds, but does so with the opposite affinities of CI. When Cro binds to O_R3, it prevents transcription of cI from P_RM, and only at higher concentrations does Cro bind O_R2/O_R1 and repress transcription of P_R. It has been proposed that Cro is also important for the regulation of the switch from lysogenic to lytic growth during λ prophage induction (12–14).

To induce the prophage, P_R and P_L must be derepressed to initiate transcription of the lytic genes. After initiation of the SOS response, this is accomplished through activation of the host RecA coprotease, which binds to CI and promotes autocleavage of CI (15). In a recA− host, SOS-mediated prophage induction is defective; the residual induction, caused by fluctuation of CI levels, occurs very rarely, less than once per million cells (16). This frequency is lower than that of mutational inactivation of the cI gene (17). Thus, the regulatory network maintaining lysogeny is extremely stable, more stable than the genes encoding it. The high stability is not obtained by precise control of the concentration of CI repressor because single-cell studies have shown that the concentration of repressor in stable lysogens vary greatly from cell to cell (11).

The switch between lysogenic and lytic growth of phage λ was the first genetic switch to be deciphered and the system has contributed immensely to our present understanding of developmental pathways (3). The recent observation that a DNA loop forms between O_L and O_R prompted us to investigate the role of the loop in the switching process of induction. We show here that the interaction between repressors bound at O_L and O_R increases the tolerance of the switch to fluctuations in CI concentration and argues against the role of Cro in the switch that leads to prophage induction.

Materials and Methods

Strains. All strains are derivatives of E. coli K12 and are listed in Table 4, which is published as supporting information on the PNAS web site. The parent strain of all λ fusions, NC398, is a bidirectional reporter that carries the luciferase (luc) reporter gene on one direction and the β-galactosidase (lacZ) reporter gene on the other in the chromosome at the lac locus. Any promoter region of interest can be inserted into NC398. All constructs were made by using recombineering (18).

β-Galactosidase Assays. To determine the kinetics of P_L activity upon temperature induction of cI857 lysogens, cultures were grown in LB (19) at 30°C overnight, diluted 1:200 the next morning, and induced by rapid transfer of the culture flasks to a 42°C water bath when OD600 reached 0.3–0.4. At each time point, 1 ml of culture was quickly transferred to an Eppendorf tube on wet ice. After all time points had been collected and cooled, OD600 was recorded and a 0.5-ml sample was immediately mixed with 0.5 ml of Z-buffer (20) containing 25 μl of chloroform and 25 μl of 0.1% SDS. β-Galactosidase activity in the sample was then measured as described by Miller (20).

To determine the relationship between growth temperature and activity of P_R in the cI857 lysogens, cultures were grown at 22°C overnight, diluted 1:200 the following morning, and incubated at the indicated temperatures to reach OD600 = 0.3–0.4. Samples were assayed for β-galactosidase activities.

Luciferase Assays. Luciferase activity was measured by using the reagents of the Promega Luciferase Assay System (catalog no. E1500) according to the company’s directions (21). Cells were grown and treated as described for the β-galactosidase assays up

Abbreviations: O_L, operator right; O_R, operator left; P_R, promoter right; P_L, promoter left; P_RM, maintenance promoter.

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to the point of addition of lysis buffer. Culture aliquots (0.1 ml) were centrifuged for 2 min at 21,000 \( \times \) g, and pellets were resuspended in 0.4 ml of Cell Culture Lysis Reagent with 2.5 mg/ml BSA and 1.25 mg/ml lysozyme. Twenty microliters of cell lysate was mixed with 0.1 ml of luciferase substrate (Promega Luciferase Assay Reagent, catalog no. E151A), incubated for 120 s and read in EG&G Berthold Lumat LB 9507 single sample luminometer for 10 s. The relative light unit (RLU) was normalized to \( \lambda_{600} \).

**Results**

The formation of an \( \lambda_{L}–\text{CI–}O_{R} \) regulatory complex containing a DNA loop has been demonstrated to increase repression of \( P_{RM} \) and \( P_{R} \) (8, 10, 11). This additional level of regulation is required for the prophage’s compensatory response to low doses of DNA damage, which sets the threshold level of DNA damage to which the prophage responds by induction (11). Repressor bound to \( \text{O}_{L} \) can also increase repression of the \( P_{R} \) promoter when artificially placed 3,600 bp downstream of the \( P_{R} \) transcription start site, suggesting that the formation of an \( \lambda_{L}–\text{CI–}O_{R} \) complex might also affect transcription from the early lytic promoters (9). To study the effects of the \( \lambda_{L}–\text{CI–}O_{R} \) DNA loop on regulation of the early lytic promoters, we constructed \( P_{K}–\text{lacZ} \) fusions by inserting part of the immunity region of \( \lambda \) between two reporter genes at the \( \text{lac} \) locus of \( E. \text{coli} \) as described in Supporting Text and Table 4, which are published as supporting information on the PNAS web site, and shown in Fig. 1C and D. The constructs fuse the \( cII \) gene with \( \text{lacZ} \), rendering the expression of \( \beta \)-galactosidase under \( P_{R} \) control. \( P_{R} \) is one of the two early lytic promoters that are repressed in the lysogen but activated upon induction; thus, production of \( \beta \)-galactosidase in the fusion strains reflects induction of the lytic pathway.

Strains NC416 and NC417 contain the part of the \( \lambda \) immunity region ranging from the AUG initiation codon of gene \( N \) gene on the left to the 30th codon of \( cII \) on the right. These strains retain both the left and the right operator regions (Fig. 1D). In strains NC414 and NC415, only the right operator region is retained; the strains carry the \( \lambda \) immunity region from the AUG initiation codon of the \( \text{rexA} \) gene on the left to the 30th codon of \( cII \) on the right (Fig. 1C). The AUG codon of \( N \) or \( \text{rexA} \) is fused to the 2nd codon of the \( \text{lac} \) reporter gene, while the 30th codon of \( cII \) is fused to the 7th codon of \( \text{lacZ} \). In strains NC414 and NC415, \( \text{lacZ} \) is expressed from \( P_{RM} \), whereas in NC416 and NC417, the reporter is under the control of the lytic \( P_{L} \) promoter. The fusions also vary with respect to the \( \text{cro} \) gene. NC414 and NC415 carry the wild-type \( \text{cro} \) gene, whereas NC415 and NC417 carry a nonfunctional missense \( \text{cro27} \) allele (12). Together, the four strains represent the four possible combinations of the \( \text{O}_{L} \) or \( \Delta \text{OL} \) and \( \text{cro}^{+} \) or \( \text{cro}^{-} \)genotypes.

**Table 1. Activity of \( \beta \)-galactosidase in repressed lysogens**

<table>
<thead>
<tr>
<th>Strain (relevant genotype)</th>
<th>( \beta )-Galactosidase activity, Miller units</th>
</tr>
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<tbody>
<tr>
<td>NC414 ( (\Delta \text{O}_L, \text{cro}^{+}) )</td>
<td>21 (3.1)</td>
</tr>
<tr>
<td>NC415 ( (\Delta \text{O}_L, \text{cro}^{-}) )</td>
<td>15 (1.1)</td>
</tr>
<tr>
<td>NC416 ( (\text{O}_L^{+}, \text{cro}^{+}) )</td>
<td>4 (1.2)</td>
</tr>
<tr>
<td>NC417 ( (\text{O}_L^{+}, \text{cro}^{-}) )</td>
<td>4 (1.3)</td>
</tr>
</tbody>
</table>

The \( \lambda_{L}–\text{CI–}O_{R} \) complex increases repression of \( P_{R} \) 4-fold. Samples were grown at 30°C and assayed for \( \beta \)-galactosidase activity in early log phase as described in Materials and Methods. The averages of four independent experiments are shown. Standard deviations are given in parentheses.
The OL–CI–OR Complex Increases Repression of Pr in the Native Prophage Configuration. The expression of β-galactosidase in log phase NC414, NC415, NC416, and NC417 cultures growing at 30°C under repressed conditions is shown in Table 1. It is evident that the presence of the OL region increases repression of Pr when the immunity region is located in the chromosome in its native configuration confirming the results obtained in multicopy systems (8–10). Note that our OL-deletion also removes the rex genes between OL and cl. We addressed the possibility that the decreased repression in the OL-deletion is not because of the absence of rex and demonstrated that the presence or absence of rex genes in the OL background does not measurably affect repression of Pr at temperatures ranging from 30°C to 37°C (data not shown). In the absence of functional Cro, the β-galactosidase activity is ~4-fold higher in the absence of the left operator region than in its presence, reflecting that OL increases the repression of Pr 4-fold. The 4-fold increase in Pr activity with OL deleted is essentially the same as when Cl is supplied at a constant level in trans (8–10), showing that OL is increasing the efficiency of Pr repression at the normal lysogenic concentration of Cl by >4-fold. We noted that in the absence of the left operator region, Pr activity is decreased ~30% in the cro– mutant (NC415) as compared to cro+ (NC414). This observation could either reflect a uniformly elevated expression from Pr in the cro– cells in comparison with the cro+ cells, or it could reflect that a fraction of the cro+ population has switched to the “lytic” state where Pr is not repressed by CI. The latter possibility is supported by the observation that, when the four strains are grown on MacConkey lactose agar plates, red colonies appear more frequently among NC414 (OL– cro+) cells than among NC415, NC416, and NC417 cells (Fig. 2). The red color indicates that these cells are producing higher levels of β-galactosidase, which metabolizes lactose. However, when both OL and OR regions are present, we were unable to detect any difference between cro+ and cro– in Pr expression (Table 1, compare NC414 and NC417). This observation could indicate that the interaction between OL and OR prevents Cro from switching the prophage from the lysogenic to the lytic state. To examine this hypothesis further, we studied the activity of Pr at low repressor concentrations. The cl allele present in our reporter strains is the cI857 allele, which encodes a temperature-sensitive repressor protein (22). The cI857 allele behaves as wild-type cl at 30°C, whereas there is essentially no repressor activity at temperatures above 40°C. Thus, a range of repressor activities can be obtained by growing the strains at a range of temperatures between the permissive and restrictive temperatures.

Role of Cro in the Switch Process? The expression of lacZ as a function of growth temperature for NC414, NC415, NC416 and NC417 is shown in Fig. 3. If cro is critical for the switch (14), we expect that derepression of Pr will occur at lower temperatures in the cro+ than in the cro– cells, suggesting that Cro can sway the switch from lysogenic to lytic growth at intermediate concentrations of CI. We obtained that result for the strains that do not carry the left operator region: NC414(cro+) and NC415(cro–) (Fig. 3). However, the expected pattern was not seen in the strains that carry both the left and the right operator regions. Although the role of rex genes, if any, in the switch process remains to be investigated in strains containing both operator regions, no difference in β-galactosidase levels were observed between NC416(cro+) and NC417(cro–) cells. In fact, the derepression of Pr as a function of growth temperature appears identical for NC415, NC416, and NC417, suggesting that the DNA loop between the left and right operator regions renders the switch insensitive to the presence of the cro+ gene.

The Effect of Cro Mediated Repression of Pr Transcription in the Presence and Absence of OL. We investigated whether the left operator region had any influence on the regulation of Pr in the absence of functional Cl. We inactivated Cl by shifting growing cultures from 30°C to 42°C and followed the kinetics of Pr derepression by monitoring β-galactosidase activity at various times.
after the temperature switch. The activities of β-galactosidase as a function of time after induction for NC414, NC415, NC416, and NC417 are shown in Fig. 4. For NC414 and NC416, which carry cro<sup>-</sup>, it can be seen that ~10 min after induction, Cro presumably has accumulated to a level where it binds O<sub>R</sub> and/or O<sub>L</sub>2 and represses P<sub>R</sub>, causing the cro<sup>-</sup> curves to diverge from the cro<sup>+</sup> curves (NC415 and NC417). It took equally long for Cro to accumulate to a concentration where it negatively autoregulates P<sub>R</sub> in NC414 (ΔO<sub>L</sub> and NC416 (O<sub>L</sub>), and the degree of Cro-mediated repression of P<sub>R</sub> was similar in the two strains (Table 2). Therefore, we saw no indication that the left operator region participated in regulation of P<sub>R</sub> by Cro when CI was absent as expected (8), consistent with the hypothesis that O<sub>L</sub> participates in regulation at O<sub>R</sub> through the CI-mediated DNA loop (10).

Repression by Cro. It was suggested earlier on that the primary role of Cro in prophage induction is to repress the early lytic promoters, P<sub>R</sub> and P<sub>L</sub>, as well as the maintenance promoter P<sub>RM</sub>. The relationship between CI and Cro is that CI is a strong repressor specialized for complete turnoff of lytic functions, whereas Cro is a weak repressor functioning in a partial turn down of the early lytic promoters to allow progression into the late lytic phase (23, 24). Our unique divergent reporter constructs allowed us to measure the degree of repression of P<sub>R</sub>, P<sub>L</sub>, and P<sub>RM</sub> by CI and Cro in this study.

To measure the degree of repression of P<sub>R</sub>, P<sub>L</sub>, or P<sub>RM</sub> by Cro, we compared the luciferase activity of NC416 (N::luc, cro<sup>-</sup>) and NC417 (N::luc, cro<sup>-</sup>) at 42°C, where CI857 is inactive (Table 3). Likewise, to measure the degree of repression of P<sub>RM</sub> by Cro, we compared luciferase in NC414 (rexA::luc, cro<sup>-</sup>) and NC415 (rexA::luc, cro<sup>-</sup>) at 42°C (Table 3). The degree of repression of P<sub>L</sub> and P<sub>R</sub> by CI in the presence of both the left and right operator regions was estimated by comparing the activity of the respective reporter genes in NC417 (P<sub>L</sub>::luc<sub>Z</sub>, P<sub>L</sub>::luc, cro<sup>-</sup>) grown at 30°C, where CI857 is active, with NC417 grown at 42°C, where CI857 is inactive (Table 3). The luciferase protein is inherently heat sensitive and forms inactive aggregates after temperature shifts, but enzyme activity is restored with the aid of host chaperones after a few minutes at the high temperature (25).

We observed full restoration of luciferase activity ~15 min after shifting from 30°C to 42°C in a reporter containing wild-type CI (not heat-sensitive; strain WAS) (data not shown). Luciferase activities reported here were measured 60 min after the temperature shift. We observed that, in the absence of CI, Cro represses the P<sub>R</sub> promoter ~1.7-fold (Table 3), although a stronger repression was reported when a fusion in which transcription terminators were present between P<sub>R</sub> and a reporter gene (galK) was used (26). In the current bidirectional reporter setup, Cro repressed P<sub>RM</sub> promoter ~2.5-fold. (Table 3).

Discussion

Repression of P<sub>R</sub> by the O<sub>R</sub>–CI–O<sub>L</sub> Complex in its Native Configuration. Tight repression of P<sub>R</sub> is believed to be vital for the maintenance of lysogeny, because otherwise Cro would be produced, bind to O<sub>R</sub>3, and repress P<sub>RM</sub>, turning off cl transcription (27). Mathematical modeling studies of the lysis–lysogeny switch have concluded that CI bound at O<sub>R</sub> cannot repress Cro production sufficiently to stably maintain the lysogen unless there are additional levels of repression of P<sub>R</sub> (16, 28). We have confirmed that an interaction between the left and right operator regions increases CI-mediated repression of P<sub>R</sub> ~4-fold. Thus, our data are consistent with that of Revet et al. (9) and Dodd et al. (8) in suggesting that the DNA loop between O<sub>L</sub> and O<sub>R</sub> constitutes the missing level of regulation. Our data also support the model of Dodd et al. (8) to explain that the means by which the presence of O<sub>L</sub> allows tighter repression of P<sub>R</sub> at physiological CI concentrations. The extra level of cooperativity added by the formation of a CI octamer increases the binding affinity of CI to the involved operator sites, thus lowering the critical concentration of CI needed to maintain lysogeny.

O<sub>L</sub> Contributes to Repression of Cro Synthesis. According to the classic model, Cro's major role during induction is to bind O<sub>R</sub>3 and thereby inhibit cl expression from P<sub>RM</sub> (14, 27). If the reduction in cl expression were important for effective induction of the early lytic promoters, we would expect a cl857 cro<sup>-</sup> prophage grown at intermediate temperatures to contain more repressor than the corresponding cro<sup>+</sup> prophage, and thus require inactivation of a larger fraction of the repressor proteins to effectively derepress P<sub>R</sub>. Instead, we observe the same amount of derepression of P<sub>R</sub> at the same growth temperature in cro<sup>+</sup> (NC416) and cro<sup>-</sup> (NC417),
indicating that the same fraction of CI must be denatured to de-repress \( P_R \) in the two strains (Fig. 3). It is possible that differences in de-repression temperature exist, which we cannot detect with our assay. The fraction of CI inactivated is not a linear function of the growth temperature, and it is likely that the concentration of active CI varies dramatically within a limited range of growth temperatures (22).

When \( O_1 \cdot CI - O_R \) complex formation is prevented by the absence of \( O_1 \), \( P_R \) is less repressed, more Cro is produced, \( P_{RM} \) is more repressed, and less CI is made, so it is easier to switch to the lytic mode. In contrast, in the presence of \( O_1 \), \( P_R \) is tightly repressed, and so Cro production is blocked, thus maintaining the lysogenic mode. These results suggest that Cro may not play any role in the switch from lysogenic to lytic state of \( A \) and are in agreement with the recent findings of Dodd and colleagues (8, 10), who showed that Cro may have a lesser role in prophage induction than previously perceived.

**Cro as a Weak Repressor of Early Lytic Functions.** The primary role of Cro in prophage induction may be to repress \( P_R \) and \( P_L \) after inactivation of CI (23, 24). In support of this hypothesis, a \( \lambda EO17 \) cro\(^{-} \) phage is capable of forming plaques when grown at 37°C, although no plaques are observed at 30°C or 42°C (24). Apparently, efficient plaque formation requires partial repression of the early lytic promoters and the repression can be exerted either by Cro or can, in the absence of Cro, be substituted by partially active CI. The Gibbs free energies of Cro and CI dimers binding to the operator regions (29, 30) support a stronger repression of \( P_R \) and \( P_L \) by CI than by Cro, and this has also been demonstrated for \( P_R \) in vivo (31). Our double reporter constructs allowed us to measure the degree of repression of \( P_R \) and \( P_L \) by Cro and CI all in the same genetic background, allowing a direct comparison of the obtained values. As expected, we observe a much higher degree of repression of both lytic promoters by CI (~400-fold) in the absence of Cro than by Cro (~1.7-fold) in the absence of CI, consistent with the idea that CI acts as a strong repressor of lytic functions and Cro acts as a weak repressor of the early lytic promoters. Because Cro plays little, if any, role in effecting the lysogenic to lytic epigenetic switch, as discussed above, and because single-copy cro partly represses \( P_{RM} \) and the lytic promoters, we conclude that this ability of Cro to partly repress \( P_{RM} \) and/or the lytic promoters is not needed in inducing the “thermal labile lysogen.” Formally, it remains possible that Cro repression of one or more of these promoters is needed for induction of the complete prophage.

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### Table 3. Repression of \( P_R \), \( P_L \), and \( P_{RM} \) by Cro and CI

<table>
<thead>
<tr>
<th>Repression</th>
<th>( P_R ) ( \beta )-galactosidase, Miller units</th>
<th>( P_L ) luciferase, ( O_L ) luciferase units</th>
<th>( P_{RM} ) luciferase, ( O_{RM} ) luciferase units</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cro</td>
<td>( O_L ) \ 1.516 \ 1.555 \ ( 5.9 \times 10^6 ) \ 6.7 \times 10^5</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>( \Delta O_L ) \ 1.7 \ 1.6 \ 1.7 \ 2.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CI</td>
<td>( O_L ) \ 1.516 \ ( 5.9 \times 10^6 )</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>( \Delta O_L ) \ 4 \ 1.4 \times 10^4</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*For repression by Cro in the absence of CI, cultures of NC414, NC415, NC416, and NC417 were grown at 30°C overnight, then diluted and shifted to 42°C to reach OD\(_{600}\) = 0.3–0.4. \( \beta \)-galactosidase and luciferase activities were measured as described in Materials and Methods. For repression by CI in the absence of Cro, cultures of NC417 were grown at 30°C overnight, then split, diluted 1:200, and incubated at 30°C and 42°C, respectively, to reach OD\(_{600}\) = 0.3–0.4. \( \beta \)-Galactosidase and luciferase activities were measured as described.*