The C-terminal domain of the Escherichia coli RNA polymerase α subunit plays a role in the CI-dependent activation of the bacteriophage λ pm promoter

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ABSTRACT

The bacteriophage λ pm promoter is required for maintenance of the λ prophage in Escherichia coli, as it facilitates transcription of the cl gene, encoding the λ repressor (CI). CI levels are maintained through a transcriptional feedback mechanism whereby CI can serve as an activator or a repressor of pm. CI activates pm through cooperative binding to the O1 and O2 sites within the O operator, with the O2-bound CI dimer making contact with domain 4 of the RNA polymerase α subunit (αCTD). Here we demonstrate that the 261 and 287 determinants of the C-terminal domain of the RNA polymerase α subunit (αCTD), as well as the DNA-binding determinant, are important for CI-dependent activation of pm. We also show that the location of αCTD at the pm promoter changes in the presence of CI. Thus, in the absence of CI, one αCTD is located on the DNA at position −44 relative to the transcription start site, whereas in the presence of CI, αCTD is located at position −54, between the CI-binding sites at O1 and O2. These results suggest that contacts between CI and both αCTD and σ are required for efficient CI-dependent activation of pm.

INTRODUCTION

Bacteriophage λ is a temperate phage which can enter one of two alternative developmental pathways, lytic or lysogenic, upon infection of its host, Escherichia coli (1,2). When the lysogenic pathway is chosen, phage DNA is incorporated into the E. coli genome, forming a prophage that can be maintained in this state for many cell generations. Stable maintenance of the prophage is achieved through the action of the phage-encoded repressor, the λ repressor (CI), which both represses the lytic promoters, pm and pr, and stimulates transcription of its own gene from the pm promoter (3). The pr and pm promoters are divergently arranged with their start sites separated by only 82 bp. Both promoters are regulated by the binding of CI dimers to three related 17-bp sequences, O1, O2 and O3, located at −74 to −58, −50 to −34 and −27 to −11, respectively, with respect to the transcription start site at pm. A CI dimer bound at the high-affinity operator, O1, acts as a repressor of the pm promoter but also stabilizes the binding of a second CI dimer to a lower-affinity operator, O2, and the second dimer, in turn, interacts with RNA polymerase (RNAP) to stimulate transcription from pm above basal levels (3,4). This stimulation occurs at the isomerization step (kD) in the transcription initiation pathway that leads to open complex formation (5,6). At higher concentrations, CI also binds to O3, thereby repressing pm (7).

Each CI monomer comprises an N-terminal DNA-binding domain (residues 1–92) and a C-terminal oligomerization domain (residues 132–236) connected by an interdomain linker known as the 'hinge' region (8). Detailed structural information is available for the isolated N-terminal and C-terminal domains (9–13). The oligomerization domain participates in dimerization of CI monomers and is also involved in weaker cooperative interactions between pairs of dimers bound to adjacent operator sites. The nature of both of these types of interaction have been elucidated by X-ray crystallography (12,13). It has also been shown that repressor tetramers (i.e. pairs of dimers) bound at O1–O2 and O1–O2 can interact through their oligomerization domains over a distance of ~3 kb, forming an octamer that enhances
repression of $p_R$ (13–15). The N-terminal domain of CI contains a DNA-binding helix-turn-helix motif which is responsible for operator recognition. In addition, residues exposed on the first helix (specifically E34 and D38) generate a negatively charged patch which, in the case of the downstream subunit of the CI dimer bound to $O_R2$, is involved in interactions with positively charged residues (R588, K593 and R596) on the surface of domain 4 of the RNAP $\sigma^70$ subunit ($\sigma_4$) during activation of $p_M$ (6,16–23). For this reason, CI is classified as a Class II activator, along with other activators which bind to sites overlapping the −35 region and, in most cases, activate transcription by contacting $\sigma_4$ (22,24,25).

At many bacterial promoters, the C-terminal domain of the RNAP $\alpha$ subunit ($\alpha$CTD) interacts with upstream promoter DNA, the RNAP $\sigma^70$ subunit and/or transcription activator proteins (24,26). These interactions are mediated by determinants on the surface of $\alpha$CTD and are facilitated by the presence of a flexible linker connecting $\alpha$CTD to the N-terminal domain (27–29). For example, residue 265, and neighbouring residues, contribute to the 265 determinant, which is responsible for interactions with DNA (30–33). Similarly, residue 261 and neighbouring residues contribute to the 261 determinant, that can contact $\sigma_4$ (34–36), whereas the side chains of valine 287 and neighbouring residues form a surface-exposed patch, the 287 determinant, which interacts with an activatory surface, AR1, on CRP (cyclic AMP receptor protein) and the 287 determinant, which interacts with an activatory and neighbouring residues form a surface-exposed patch, leading to substitution of glutamate for lysine at position 271 within $\alpha$CTD, decreases $\lambda$ prophage stability (39,40). This observation could be explained by a defective interaction between the mutant $\alpha$CTD and the CI repressor at $p_M$. Therefore, the aim of this work was to determine whether $\alpha$CTD plays a role in CI-dependent activation of $p_M$. Our results show that determinants on the surface of $\alpha$CTD are required for fully efficient activation of CI. In addition, we demonstrate that the location of $\alpha$CTD at $p_M$ is shifted further upstream in the presence of CI. These observations suggest that CI makes direct contact with $\alpha$CTD at $p_M$ and that this interaction is important for transcription activation by CI.

**MATERIALS AND METHODS**

**Bacterial strains**

The *E. coli* $rpoA^+$ strain, WAM106 [araD139, $\Delta$(argF-lac) U169, $\Delta$(his-gnd), thi, proL510, glgS, fliB5301, relA1, deoC1, rbsR], and its otherwise isogenic $rpoA341$ derivative (WAM105), bearing a chromosomal mutation that results in the K271E substitution in the RNAP $\alpha$ subunit (39), were used. Strains WAM140, WAM141 and WAM144, harbouring chromosomal $rpoA261$, $rpoA269$ and $rpoA287$ alleles, encoding $\alpha$ subunits with alanine substitutions at positions 261, 269 and 287, respectively, are otherwise isogenic with WAM106 and were isolated by a previously described procedure (34,41,42). Strain WAM142, bearing the chromosomal mutation $rpoA271$, which results in substitution K271A in $\alpha$ was isolated as a Cym$^+$ Mel$^+$ pseudorevertant of strain WAM105. The *E. coli* strain, TAP90 (supE44, supF58, hisD, pro, leuB, thi-1, rpsL, lacY, tonA1, recD1903::miniTet) was used to titrate bacteriophage containing the S7 amber allele (43).

**Bacteriophage, plasmids and gene fusions**

Bacteriophage $\lambda$exR57S7 (44), which is unable to lyse *E. coli* cells unless the supF suppressor allele is present, was used for measuring prophage stability. For the expression of mutant $rpoA$ alleles for the $\alpha$CTD alanine scan analysis, derivatives of plasmid pHf1 were used. For the expression of mutant $rpoA$ alleles, both of which are p15A derivatives, were used to express the phage $\lambda$ CI gene. pGW857 encodes the thermostabile $C_k357$ protein under control of the lac promoter (48) and thereby allows for complete inactivation of repressor function by growth at 42°C. Plasmid pACcI was used to overexpress the wild-type CI gene from the lacUV5 promoter (49). For measuring the activity of the $p_M$ promoter, two $p_F$–lacZ fusion plasmids were used: pHAI, a pBR322-based replicon, and pTJSpM, an RK2-based replicon. To construct pHAI, the wild-type $p_M$ promoter region (248 bp) was amplified by PCR using the $\lambda$ plasmid pKB2 (50) as a template, and the following primers: 5'-GCC GGA TCC CCA TCT TGT CTG C and 5'-TAT GCG TTG TTA GCT ATA GAC TCC TTA GTA C (35 cycles of the following program were performed: denaturation at 95°C for 30 s, annealing at 55.4°C for 30 s, extension at 72°C for 30 s). The product of the amplification was digested with BamHI and cloned with other activators (33,34,37,38).

Previously, we have shown that the $rpoA341$ mutation, leading to substitution of glutamate for lysine at position 271 within $\alpha$CTD, decreases $\lambda$ prophage stability (39,40). This observation could be explained by a defective interaction between the mutant $\alpha$CTD and the CI repressor at $p_M$. Therefore, the aim of this work was to determine whether $\alpha$CTD plays a role in CI-dependent activation of $p_M$. Our results show that determinants on the surface of $\alpha$CTD are required for fully efficient activation of CI. In addition, we demonstrate that the location of $\alpha$CTD at $p_M$ is shifted further upstream in the presence of CI. These observations suggest that CI makes direct contact with $\alpha$CTD at $p_M$ and that this interaction is important for transcription activation by CI.
Measurement of the effect of mutant rpoA alleles on CI-dependent activation in vivo

For the alanine scanning experiment (merodiploid), expression of wild-type cl from pACcl, and mutated rpoA alleles from pHTT1z and pREIIz derivatives, was simultaneously induced by addition of IPTG (0.1 mM final concentration) to cultures of WAM106 harbouring pJMH1 and pTJSpM growing at 37°C. β-galactosidase activity was measured 1 h later. To assess the effect of haploid rpoA alleles on CI-dependent activation of ρM, strains harbouring chromosomal mutant rpoA alleles were transformed with pGW857 and pAHAl, and cultures were grown at 43°C to OD578 = 0.2 [the cI57(is) gene product is inactive under these conditions and β-galactosidase activity is very similar to that measured in cells devoid of pGW857; data not shown] whereupon IPTG was added (0.05 mM final concentration) and the culture was immediately shifted to 30°C. Following incubation at this temperature for 1 h the β-galactosidase activity was measured. This induction regime minimizes problems due to CI occupancy of O53 present on pAHAl (data not shown).

Measurement of β-galactosidase activity

The activity of β-galactosidase in bacterial cells was measured according to Miller (56). Since we used a multicopy lacZ fusion, the β-galactosidase activities were calculated per plasmid copy number, estimated as described previously (57), to compensate for any possible copy number variation between strains. For the alanine scanning experiment, bacteria were grown at 37°C to OD578 = 0.2, induced with 0.1 mM IPTG and, following further incubation for 1 h, β-galactosidase assays were performed. Results presented are averages of at least three independent experiments and are shown with standard deviations.

Measurement of the efficiency of prophage maintenance

λ prophage maintenance in lysogenic E.coli strains was estimated by measuring the efficiency of spontaneous induction of a λcI857S7 prophage as described previously (40). Briefly, samples (5 ml) of exponential phase cultures (OD578 0.2–0.5) of bacteria lysogenic for bacteriophage λcI857S7, growing at 30°C, were withdrawn and shaken vigorously with chloroform (0.5 ml) for 1 min to release progeny phage. Following centrifugation, liberated phages were titrated on the suppressor strain, TAP90, at 37°C. Other samples, withdrawn at the same time as those for phage titration, were centrifuged. Cell pellets were resuspended in 0.9% NaCl and used for titration of bacteria on LB agar at 30°C. Finally, the number of phages yielded per bacterial cell was calculated.

Protein purification and reconstitution of RNA polymerase

Plasmid pT7cl1Sa109Hi6 (21) was used for overproduction of C-terminally His-tagged CI protein, which was purified as described previously (21). For the reconstitution of RNAP, inclusion bodies of RNAP β, β' and σ70 subunits from strains XL1-Blue (MKSe2), BL21(DE3)(pT7β'), and BL21(DE3)(pLHN12σ), respectively, were prepared as described previously (58). His-tagged RNAP α subunits were prepared using plasmid pHTT71NHα (58). Derivatives of pHTT71NHα carrying mutant rpoA alleles were constructed by replacing the HindIII–BamHI fragment, which encodes αCTD and the interdomain linker, with the corresponding fragments from derivatives of pHTT1z and pREIIz encoding the appropriate alanine-substituted α mutants (see above) or from pLAW2phs (encoding α containing the K271E substitution) (39). Overexpression of the α subunits in strain BL21(DE3), purification of α by Ni2+-affinity chromatography and reconstitution into RNAP were performed essentially as described previously (30,58). Purification of α subunits with single cysteine residues, conjugation with Fe.BABE, and reconstitution into RNAP was performed as described by Lee et al. (59).

In vitro transcription

Single round in vitro transcription reactions were performed in a total volume of 20 μl in buffer containing 50 mM KCl, 40 mM Tris-HCl (pH 8.0), 10 mM MgCl2, 1 mM DTT, 100 μg/ml BSA and 30 ng linear template DNA. Template DNA containing the ρM promoter was prepared by isolating the 1200-bp Ndel–EcoRI fragment from plasmid pRLGpMmut. The 1313-bp Ndel–PstI fragment from the same plasmid, containing the RNA I gene, served as the internal control. The binding reaction of CI (80 ng) to the DNA (30 ng) was carried out at 37°C for 10 min, after which time in vitro reconstituted RNAP was added and the incubation continued for a further 10 min (this concentration of CI gave rise to ~4-fold activation of ρM in the presence of wild-type reconstituted RNAP (results not shown)). After the addition of nucleotides (CTP, GTP and ATP, each to a final concentration of 150 μM, UTP to 15 μM and 0.6 μCi [α32P]-UTP per reaction) and heparin to 50 μg/ml, the samples were incubated at 37°C for 15 min and the reactions were stopped by the addition of an equal volume of 95% formamide containing 20 mM EDTA, 0.05% bromophenol blue and 0.05% xylene cyanol. The samples were separated by electrophoresis in 6% polyacrylamide gels containing 46% urea in TBE buffer. The gel was dried, and RNA bands were visualized and quantified, following background subtraction, using a PhosphorImager (Bio-Rad). Concentrations of RNAP, calibrated to give the same amount of transcription from the activator-independent RNA-I promoter, were: 46 nM wild-type RNAP, 34 nM RNAP αK271E, 54 nM RNAP αK271A, 13 nM RNAP D258A, 28 nM RNAP αE261A, 35 nM RNAP αR265A, 34 nM RNAP αV287A.

Fe-BABE-mediated hydroxyl radical footprinting

A 150-bp DNA fragment containing the ρM promoter was amplified from bacteriophage λ DNA by PCR using primers 5'-GCT TTA AGC TTA CTT GCG TCC TCA AGC TGC-3' and 5'-CCT GAA TTC ATG CAA CCA TTA CCA CCG-3', cleaved with HindIII and EcoRI and cloned into the vector pSR (60). A 220-bp AatII–HindIII fragment was purified from the resultant plasmid
important role in CI-dependent activation of \( p_M \). RNAP was also reconstituted with the R265A \( \alpha \) subunit. In addition, due to our previous observation that the K271E substitution in \( \alpha \) causes decreased prophage stability, we included RNAP reconstituted with the 271E and 271A \( \alpha \) subunits in the analysis.

Our results are in general agreement with the in vivo results, i.e. the abundance of \( p_M \)-derived transcripts was significantly decreased when RNAP was reconstituted with \( \alpha \) containing the 258A, 261A and 287A substitutions, whereas the efficiency of transcription obtained using RNAP reconstituted with \( \alpha \) harbouring the 265A substitution was comparable to that of wild-type RNAP (Figure 2). Consistent with its effect on prophage stability, RNAP reconstituted with 271E \( \alpha \) was significantly less active at the \( p_M \) promoter in vitro. This was also the case with 271A \( \alpha \), although alanine substitution at this position does not exert a negative effect at \( p_M \) in vivo (Figure 1A).

Effect of substitutions in \( \alpha \)CTD determinants important for CI function in vitro in the absence of wild-type \( \alpha \)

In vivo transcription assays. To investigate the full effect of amino acid substitution within \( \alpha \)CTD on CI-dependent activation of the \( p_M \) promoter in vivo, we constructed \( E. coli \) mutant strains harbouring mutations within the chromosomal \( rpoA \) gene that result in alanine codon substitutions at positions 261, 269, 271 and 287 (\( rpoA261, rpoA269, rpoA271 \) and \( rpoA287 \), respectively) [it was not possible to transfer to the \( E. coli \) chromosome alleles encoding substitutions at positions 265, 268 or 299 within the DNA-binding determinant (34; M.S.T., unpublished data)]. The mutant strains were transformed with a plasmid containing inducible CI function and a plasmid harbouring a \( p_M \)–\( lacZ \) fusion, and the effect of induction of \( cl \) expression on \( p_M \) activity was measured.

Under these conditions we observed ~5-fold activation of transcription from \( p_M \) in the \( rpoA^+ \) host (Table 1), which compares favourably with previously reported induction ratios (19,20). However, in strains harbouring the mutant \( rpoA \) alleles, CI-dependent activation of \( p_M \) was only 45–60% as efficient as in the wild-type strain, with the C269A substitution causing the most profound effect (Table 1). By way of comparison, the \( p_M \) activity in the strain harbouring the \( rpoA341 \) allele, encoding the K271E substitution in \( \alpha \) (39,40), was ~55% as efficient as in the wild-type strain (Table 1). These results confirm the important roles played by the 261 and 287 determinants and the DNA-binding region of \( \alpha \)CTD in CI-dependent activation at \( p_M \).

**\( \lambda \). Prophage stability.** As maintenance of a \( \lambda \) prophage only requires CI function, we investigated whether substitutions within \( \alpha \)CTD which impair CI-dependent activation of the \( p_M \) promoter also impair \( \lambda \) prophage maintenance. To do this, we compared the efficiency of spontaneous induction of a \( \lambda cI857S7 \) prophage in hosts harbouring wild-type or mutant \( rpoA \) alleles on the chromosome. As expected, we found that alanine substitution at positions 261, 269, 271 and 287 in \( \alpha \) resulted in
a higher frequency of spontaneous induction of the \( \lambda \) prophage relative to the wild-type host (3–8-fold increase, depending on the position of the substitution) (Table 1). Consistent with the \( p_M \) promoter activity measurements, the prophage was most unstable in the host carrying the \( rpoA269 \) allele. As shown previously, we measured a 5-fold increase in spontaneous induction of \( \lambda \) prophages in the \( rpoA341 \) mutant relative to the wild-type (Table 1; 40). In support of the hypothesis that decreased prophage stability was due to decreased CI levels, overexpression of the \( cI \) gene from plasmid pAC\( cI \) resulted in equally efficient maintenance of the prophage in the wild-type and in all tested mutant strains (data not shown).

**Figure 1.** Identification of \( \alpha \)CTD residues important for CI-dependent activation of \( p_M \) in vivo. (A) Strain WAM106, containing plasmids pTJSpM, pJMHI and pAC\( cI \), was transformed with each of a set of plasmids encoding the RNAP \( \alpha \) subunit in which each residue of \( \alpha \)CTD was changed individually to alanine. Cultures were grown at 37°C to OD\( O_600 \) = 0.2 in LB medium containing appropriate antibiotics, at which time IPTG was added to a final concentration of 0.1 mM. After 60 min induction of \( \alpha \) and CI synthesis, the \( \beta \)-galactosidase activities were determined. The activities are presented relative to the activity of the strain harbouring plasmid pLAW2 encoding wild-type \( \alpha \) (100% = 2300 Miller units) and are averages of at least three independent experiments. Grey bars indicate positions when alanine occurs naturally. Black bars correspond to the residues in which alanine substitution causes a decrease in activity of \( \geq 20\% \) compared to wild-type \( \alpha \). (B) Structure of \( \alpha \)CTD, showing in black the residues that are important for CI-dependent activation of \( p_M \). Residue K271 is highlighted in grey for reference.

**Location of the \( \alpha \)CTD–DNA interactions at the \( p_M \) promoter**

To determine the location of \( \alpha \)CTD at the \( p_M \) promoter we exploited the DNA cleavage reagent, iron [\( S \)]-[\( p \)-bromoacetamidobenzyl] ethylenediaminetetraacetate (Fe-BABE), that can be attached to cysteine residues introduced at specific locations within \( \alpha \)CTD (59,64,65). Thus, we derivatized \( \alpha \)CTD with Fe-BABE by employing a functional \( \alpha \) subunit in which cysteine was introduced at position 273, and used the derivatized product to reconstitute RNAP (53,59).

Analysis of DNA scission products following formation of the RNAP–Fe-BABE–\( p_M \) complex revealed that,
of CI in vitro. (A) The efficiency of transcription from pM in the presence of reconstituted mutant RNAPs is shown in a typical transcription gel. Single-round in vitro transcription experiments were performed using linear template DNA containing pM or specifying RNA-I, together with CI and RNAP reconstituted with hexahistidine-tagged α derivatives containing alanine substitutions at the positions indicated. The activities of purified RNAPs were normalized at the transcription from RNA-I, together with CI and RNAP reconstituted with hexahistidine-tagged α (to 0.05 mM) and simultaneous shift to 30°C yield obtained with wild-type RNAP. (B) The efficiency of CI-dependent transcription from pM in the presence of each reconstituted mutant RNAP. The results are from three independent experiments. Values (with standard deviation) are expressed as percentages of the transcript yield obtained with wild-type RNAP.

**Table 1.** Effect of different chromosomal rpoA alleles on CI-dependent activation of pM and on prophage stability

<table>
<thead>
<tr>
<th>Chromosomal rpoA allele (α subunit)</th>
<th>Activation of pM by CI</th>
<th>Relative frequency of prophage induction</th>
</tr>
</thead>
<tbody>
<tr>
<td>rpoA4 (α wild-type)</td>
<td>4.9</td>
<td>1</td>
</tr>
<tr>
<td>rpoA341 (α K271E)</td>
<td>2.7</td>
<td>4.9</td>
</tr>
<tr>
<td>rpoA271 (α K271A)</td>
<td>2.7</td>
<td>4.7</td>
</tr>
<tr>
<td>rpoA261 (α E261A)</td>
<td>2.9</td>
<td>2.7</td>
</tr>
<tr>
<td>rpoA269 (α C269A)</td>
<td>2.2</td>
<td>7.9</td>
</tr>
<tr>
<td>rpoA287 (α V287A)</td>
<td>3.0</td>
<td>5.5</td>
</tr>
</tbody>
</table>

β-galactosidase activities were measured in cells harbouring pAHA1 and pGW857 at 43°C (basal pM activity) and 1 h after IPTG addition (to 0.05 mM) and simultaneous shift to 30°C (CI-stimulated pM activity), and calculated per single copy of pAHA1 per cell. The values are presented in the table represent the induction ratios and were calculated by dividing the value for the stimulated pM activity by the value for the basal activity. The efficiency of transcription from pM in the presence of each reconstituted mutant RNAP. Values (with standard deviation) are expressed as percentages of the transcript yield obtained with wild-type RNAP.

**DISCUSSION**

The location of the stimulatory CI-binding site (O_R2) at pM (see Figure 3B) suggests that CI activates this promoter by a Class II-type mechanism (22,24,69). Consistent with this, we have shown that a negatively charged patch on the surface of the CI DNA-binding domain, located in helix 1 of the HTH motif, stimulates transcription from pM through making contact with a positively charged patch on σ_70 (23). In this report, we have demonstrated that determinants on αCTD also contribute to CI-dependent activation of pM. Alanine scanning analysis indicated that some of the surface-exposed residues on αCTD which are required for efficient CI-dependent activation are located within or near the previously identified 261 determinant (i.e. R255, P256, D258, E261 and K271) and the 287 determinant (V287). These determinants are located on opposite sides of αCTD and have been shown to play roles in activator-dependent transcription at other promoters. It is intriguing that the 261 determinant is implicated in CI-dependent activation, as it has previously been shown to play a role only at Class I CRP-dependent promoters and at some UP element-dependent promoters, where it interacts with σ_70 (34–36). At other Class II promoters, where αCTD is not in a position to interact with σ_70, the 261 determinant does not play a role in transcription activation (37). Our results with Fe-BABE-derivatized RNAP show that, in the presence of CI, αCTD is located close to position −54 at pM, i.e. between O_R1 and O_R2, and therefore is also not in a position to contact σ. Therefore, the simplest explanation for our observations is that the 261 determinant is involved in contacts with CI.

The 287 determinant has been shown to interact with CRP at Class I and Class II CRP-dependent promoters and there is evidence that it interacts with MelR at the pmelAB promoter (34,37,38). Our results suggest that CI is another activator that utilizes this determinant. The involvement of residues on opposite sides of αCTD in CI-dependent activation could occur if αCTD is sandwiched between the two CI dimers, as demonstrated by the Fe-BABE analysis, with each determinant

in the absence of CI, cleavages occur in clusters separated by 10–11 bp, with the strongest signals occurring near position −44 relative to the transcription start site (Figure 3). This is consistent with the fact that pM serves as a weak promoter in the absence of CI (66). The pattern of cleavages is similar to that found at other promoters that are active in the absence of transcription activators, such as rrnB P1 or CC(−61.5)-p127 (59), and suggests that one of the two α subunits binds to the first available minor groove upstream of the −35 region while the second αCTD binds to successive minor grooves (i.e. −54, −65 and −54 being the most favoured position) (Figure 3). This is in accordance with previously published results, which suggested that the α subunit contacts sequences upstream of pM in a sequence non-specific manner (67). In the presence of CI, the strongest signals were observed near position −54, which is located in the minor groove between two CI dimers bound to major grooves within O_R2 (−34 to −50) and O_R1 (−58 to −74) (68) (Figure 3). Therefore, binding of CI results in re-positioning of αCTD at the pM promoter.
interacting with a different CI dimer. This is analogous to the situation at the artificial Class II promoter, \(ML(\text{p}L_{74.5})\), which contains tandem CRP sites centred at \(\text{p}L_{41.5}\) and \(\text{p}L_{74.5}\). At \(ML(\text{p}L_{74.5})\), one \(\alpha\)-CTD is recruited to the DNA between the two CRP-binding sites, whereas the other \(\alpha\)-CTD binds to DNA upstream of the CRP dimer bound at \(\text{p}L_{74.5}\) (70). Furthermore, the determinants of the \(\alpha\)-CTD sandwiched between the CRP dimers are likely to be aligned along the axis of the DNA, with the \(\alpha\)-CTD interacting with AR1 of the promoter-proximal CRP, as shown for the simple Class II CRP-dependent promoter \(CC(\text{p}L_{41.5})\) (37,59).

Although the location of the second \(\alpha\)-CTD at \(\text{p}M\) was not addressed in this investigation, one intriguing possibility is that, in a situation where \(OL\) (the CI operator overlapping the \(\text{p}L\) promoter) is also present, the second \(\alpha\)-CTD binds \(OL\) between the pair of CI dimers bound to the \(OL_{1}\) and \(OL_{2}\) sites.

Our results also revealed that alanine substitution of amino acids S266, N268, C269, L270 and S299 impaired CI-dependent activation. These residues are located within or near the DNA-binding surface of \(\alpha\)-CTD (33,71) (although L270 does not participate directly in DNA binding, the side chain is buried within the structure of \(\alpha\)-CTD and therefore substitution by alanine may cause a conformational change in the DNA-binding region). The DNA-binding determinant plays a role in UP element-dependent transcription initiation and at many activator-dependent promoters (24,30,34,37,53). Its involvement in CI-dependent transcription activation suggests that an interaction between \(\alpha\)-CTD and the promoter is important for CI-dependent activation. The results of the Fe.BABE analysis suggest that the important \(\alpha\)-CTD–DNA interaction is likely to be due to the \(\alpha\)-CTD positioned near \(\text{p}L_{54}\). It is noteworthy that the side chain of R265, which plays an important role in DNA binding at many promoters, does not appear to be required for efficient CI-dependent activation. However, it has been shown previously that the contribution of this residue to DNA binding at some activator-dependent promoters is minimal (34). On the other hand, the broader Fe.BABE cleavage pattern that occurs at \(\text{p}L_{54}\) in the presence of bound CI, in comparison to the more focussed cleavage at \(-44\) in the absence of CI, may indicate that \(\alpha\)-CTD is not in intimate contact with the DNA when CI is present (i.e. \(\alpha\)-CTD may be interacting with CI ‘off the DNA’) or that the interaction of the DNA-binding determinant with the promoter is different to that which occurs at many other promoters. One possible reason for this is that, for steric reasons, \(\alpha\)-CTD may not be able to readily access the \(-54\) region on the same side of the DNA as CI (Figure 4).

Firstly, the diameter of \(\alpha\)-CTD (measured from the 261

**Figure 3.** Location of \(\alpha\)-CTD–DNA interactions at the \(\text{p}M\) promoter. (A) An autoradiogram of a polyacrylamide sequencing gel showing DNA cleavage resulting from attack triggered by RNAP reconstituted with Fe-BABE-derivatized \(\alpha\) subunits. \(\text{p}M\) promoter DNA was end-labelled on the template strand and incubated with or without CI and RNAP reconstituted with \(\alpha\) derivatized with Fe-BABE at position 273. (B) DNA sequence of the divergently arranged \(\text{p}M\) and \(\text{p}R\) promoters, showing Fe-BABE-induced cleavage positions in the presence or absence of CI protein indicated by black or grey stars, respectively. Transcription startpoints (+1 position) are indicated by bent arrows. The \(-55\) and \(-10\) hexamer sequences of the \(\text{p}M\) and \(\text{p}R\) promoters are shown in boxes. Base pair co-ordinates in A and B are numbered with respect to the \(\text{p}M\) transcription startpoint.
determinant to the 287 determinant) is ~25 Å. Although the distance between the two operators, \( O_{R1} \) and \( O_{R2} \), is ~24 Å (based on a rise of 3.4 Å per bp), the separation between the two CI dimers is likely to be less than this. This is due to the fact that the adenine tract between the two operator sites contains a static bend of the order of 18°, which becomes further bent by 15–18° upon binding CI, in a large part due to untwisting of the DNA (13,72–74). Access to the DNA between \( O_{R1} \) and \( O_{R2} \) may be further restricted by the cooperative interactions which occur between the C-terminal oligomerization domains of CI (12,13,73).

The other important observation from this investigation is that the location of \( \alpha \)CTD at \( p_M \) is different in the presence and absence of CI. In the absence of CI, one \( \alpha \)CTD is located adjacent to \( \sigma_70 \) at a site that overlaps \( O_{R2} \). In the presence of CI, \( O_{R2} \) is occupied by CI and \( \alpha \)CTD is relocated to a DNA site located between \( O_{R1} \) and \( O_{R2} \) (Figure 4). This observation, together with the analysis of \( \alpha \) mutants, is consistent with a model in which activation of RNAP at \( p_M \) is mainly the result of the interaction between CI bound at \( O_{R2} \) and \( \sigma_70 \), as previously proposed (19–22). The role of the \( \alpha \)CTD–CI interaction may be to stabilize the interaction of \( \alpha \)CTD with DNA upstream of \( O_{R2} \), facilitating CI-dependent stimulation of the \( k_f \) step.

CI is not the only Class II transcription activator to make contact with \( \alpha \)CTD in addition to \( \sigma_4 \). Both MelR and CRP (at the \( galP1 \) promoter) also make a specific contact with \( \alpha \)CTD, and this interaction contributes to the overall stimulatory activity of the regulatory protein (24,38,75,76). Other examples of so-called ‘ambidextrous’ activators include LuxR and the phage Mu Mor protein (76–79). In such cases, \( \alpha \)CTD binds to the first available minor groove upstream of the activator binding site, with a preference for binding to the same face of the DNA as RNAP (38,53). In the case of \( p_M \), the first available minor groove is located between the two CI dimers bound at \( O_{R1} \) and \( O_{R2} \).

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