Fine structure of *E. coli* RNA polymerase-promoter interactions: \(\alpha\) subunit binding to the UP element minor groove

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The \(\alpha\) subunit of *E. coli* RNAP plays an important role in the recognition of many promoters by binding to the A+T-rich UP element, a DNA sequence located upstream of the recognition elements for the \(\sigma\) subunit, the \(-35\) and \(-10\) hexamers. We examined DNA–RNAP interactions using high resolution interference and protection footprinting methods and using the minor groove-binding drug distamycin. Our results suggest that \(\alpha\) interacts with bases in the DNA minor groove and with the DNA backbone along the minor groove, but that UP element major groove surfaces do not make a significant contribution to \(\alpha\) binding. On the basis of these and previous results, we propose a model in which \(\alpha\) contacts UP element DNA through amino acid residues located in a pair of helix–hairpin–helix motifs. Furthermore, our experiments extend existing information about recognition of the core promoter by \(\sigma^{70}\) by identifying functional groups in the major grooves of the \(-35\) and \(-10\) hexamers in which modifications interfere with RNAP binding. These studies greatly improve the resolution of our picture of the promoter–RNAP interaction.

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Promoters in bacteria contain recognition sequences for RNA polymerase RNAP, subunit composition \(\alpha, \beta', \beta'\) in up to three distinct regions: the \(-10\) and \(-35\) elements, both recognized by the \(\sigma^{70}\) subunit (for review, see Record et al. 1996; Gross et al. 1998), and the UP element, recognized by the \(\alpha\) subunit (for review, see Gourse et al. 2000). The UP element, located just upstream of the \(-35\) hexamer, was identified as a binding site for \(\alpha\) in the rRNA promoter *rrnB* P1 of *Escherichia coli*, where it stimulates promoter activity ~30-fold (Ross et al. 1993). UP elements have been identified in other promoters in *E. coli*, in other bacterial species, and in promoters transcribed by holoenzymes containing alternative \(\sigma\) factors (e.g., Moran et al. 1982; Newlands et al. 1992; Ross et al. 1993, 1998 Fredrick et al. 1995; Negre et al. 1997). The optimal (consensus) UP element sequence was identified by in vitro selection and contains alternating A and T tracts in two distinct subsites (proximal and distal; Estrem et al. 1998, 1999). Analysis of the *E. coli* genome sequence suggests that UP elements, consisting of either one or two subsites with no more than two mismatches to consensus, occur in more than one-third of stable RNA promoters, but are also found in about 4% of mRNA promoters [Estrem et al. 1999]. Because sequences with much lesser degrees of match to consensus can still significantly affect promoter function [Ross et al. 1998], the number of promoters in which UP elements play some role in transcription is likely to be much larger. DNA–\(\alpha\) interactions also occur adjacent to many activator protein-binding sites, although, in these cases, \(\alpha\) interactions are often DNA sequence independent, the binding site(s) do not resemble the UP element consensus, and \(\alpha\) binding does not occur in the absence of the bound activator (Hochschild and Dove 1998; Busby and Ebright 1999; Gourse et al. 2000).

The ~37 kD \(\alpha\) subunit contains two independently folded domains [Blatter et al. 1994]. The amino-terminal domain [residues 8–233] is responsible for dimerization and interactions with \(\beta\) and \(\beta'\) [Hayward et al. 1991], and the carboxy-terminal domain (\(\alpha\)CTD), residues 245–329) contains the DNA-binding determinants [Blatter et al. 1994]. Genetic analyses have identified amino acid residues in the \(\alpha\)CTD critical for DNA binding and UP element function (Gaal et al. 1996; Murakami et al. 1996). These residues are essential for cell viability [Gaal et al. 1996] and are highly conserved in prokaryotic \(\alpha\) se-
quences. Amino acid sequence comparisons in conjunction with the solution structure of the αCTD, determined by nuclear magnetic resonance spectroscopy (Jeon et al. 1995; Gaal et al. 1996), suggest that α is a member of the helix–hairpin–helix (HHH) domain family of DNA-binding proteins (Shao and Grishin 2000).

Defining the interactions between components of the transcription initiation apparatus remains a central problem in molecular biology. Recently, there have been dramatic advances in our understanding of the structure of bacterial core RNAP at the atomic level (Zhang et al. 1999), but the αCTD was not resolved in this structure, and no information at the atomic level exists for complexes containing DNA bound to α or RNAP holoenzyme.

Footprinting and cross-linking studies have provided some information about the α–UP element interaction and the promoter–RNAP interaction. Footprints of RNAP on rrnB P1 and other promoters extend about 60 bp upstream of the transcription start site, and protection upstream of ∼40 is attributable to interactions with α (Kolb et al. 1993; Ross et al. 1993). In the rrnB P1 UP element, DNA backbone regions centered at ∼42 and ∼52 are protected against hydroxyl radical cleavage by α or RNAP [Newlands et al. 1991; Estrem et al. 1998]. However, a full understanding of the mode of α binding has been hindered by complicating features of its interactions with different promoter upstream regions. Not only can α bind to DNA in either a sequence-specific or a sequence-nonspecific manner, but α-dependent protection signals have also been observed upstream of ∼60, and sometimes multiple regions of protection have been observed in the same upstream region [Newlands et al. 1992, Kolb et al. 1993; Fredrick et al. 1995; Belyaeva et al. 1998]. Comprehensive cross-linking studies further suggest that the two αCTDs may oscillate among different binding sites (like fingers on a keyboard, Naryshkin et al. 2000), or alternatively may bind to different sites in different molecules in the population. Variability in the position(s) of α binding sites is most likely attributable to the long, flexible tether linking the α amino- and carboxy-terminal domains [Blatter et al. 1994; Jeon et al. 1997], as well as to intrinsic DNA distortions and protein-induced DNA bending in initiation complexes containing activators [Gourse et al. 2000; Naryshkin et al. 2000].

A binding site for a single α subunit (the consensus proximal subsite, see Fig. 1A) was defined by in vitro selection and by the use of RNAPs heterodimeric for α, [i.e., containing one wild-type α subunit and one α lacking its DNA-binding domain (ΔαCTD, Estrem et al. 1999)]. Only one αCTD was required for function of an UP element with only a proximal subsite and for protection of this subsite in hydroxyl radical footprints. The protected positions (about 4 bp on each DNA strand) are offset on the two strands by ∼3 bp, and are thus located in close proximity across the DNA minor groove [Estrem et al. 1999]. Likewise, studies with a cross-linking agent tethered by a 10-Å linker indicated that αCTD interactions with a promoter upstream region lacking a func-

![Figure 1. Promoters and base analogs used in footprinting experiments. (A) RNAP binding to rrnB P1 with an UP element containing a single proximal subsite. The proximal subsite (black rectangle) is a binding site for one αCTD. The position of the other αCTD in this complex is not known. The σ subunit binds to the −10 and −35 hexamers (grey rectangles). The non-template strand sequences of the rrnB P1 −10 and −35 hexamers and the two UP elements used in this study [4547, 4549; Estrem et al. 1999] are shown. (B) Structures of an adenine:thymine (A:T) base pair and of the base analogs 7-deaza-7-nitro adenine (A*) and uracil (U). Surfaces facing the major and minor grooves are indicated for the A:T pair. Minor groove N3 position of A (bold) was probed in DMS footprints. Major groove positions N7 of A [modified in 7-deaza-7-nitro adenine (A*)] and uracil (U). Surfaces facing the major and minor grooves are indicated for the A:T pair. Minor groove N3 position of A (bold) was probed in DMS footprints. Major groove positions N7 of A (bold) and the C5 methyl group of T (in bold; not present in U) were probed in interference footprints using templates containing A* or U.](Image 334x306 to 565x733)
information to build a structural model of the sequence-specific α-CTD–UP element interaction; for example, proteins that bind to two successive major grooves are also centered on the minor groove. The previous studies do not distinguish whether the α-CTD makes base contacts within a DNA groove, or whether these contacts, if they occur, are with the major groove or minor groove surfaces of the bases.

In this work, we address the basis for the specific interaction of α with UP element DNA. An understanding of bacterial promoter recognition and the architecture of transcription complexes containing activators requires detailed information about the α-CTD–DNA interaction. Therefore, we carried out RNAP protection footprinting studies with dimethylsulfate (DMS) and interference footprinting studies with promoter fragments containing the thymine analog uracil or the adenine analog 7-deaza-7-nitro-adenine (A*; Fig. 1B). We also examined the effects of the minor groove-binding drug distamycin on α binding and on UP element–dependent transcription. These results and those from hydroxyl radical footprinting studies (Estrem et al. 1999) indicate that α binds to the DNA backbone across the minor groove and to bases within the minor groove. Together with genetic analyses of α-CTD (Gaal et al. 1996; Murakami et al. 1996) and the recent proposal that α contains two HhH DNA binding motifs [an [HhH]_2 domain; Shao and Grishin 2000], our data suggest a model for the interaction of α with UP element DNA. In addition, our experiments provide detailed information about the proximity of the RNAP σ subunit to the major groove surface of thymine and adenine bases in the −35 and −10 hexamers.

Results

We carried out footprinting studies of RNAP on templates containing the trnB P1 core promoter fused to UP elements consisting of only a single proximal subsite (UP elements 4547 or 4549; Estrem et al. 1999, see Fig. 1A). UP elements containing only a proximal subsite are the most frequent class of UP elements in the E. coli genome (Estrem et al. 1999). Most important, use of templates with only a proximal subsite simplifies interpretation to a single DNA–α interaction.

The UP elements used in these experiments are representative of the proximal subsite sequences identified by in vitro selection and can stimulate promoter activity >100-fold in vivo. Each contains an A tract extending from −46 to −41 but has a different sequence at positions −40 and −39 (Estrem et al. 1999). The region upstream of the proximal subsite in these promoters is not protected by RNAP and does not affect promoter function (Estrem et al. 1999).

Missing bases in the UP element interfere with RNAP binding

Limited information was available about the relative importance of individual positions in the UP elements, and no information was available about whether bases on the nontemplate strand, template strand, or both make contacts with α. As a first approach to understanding the role of specific bases in proximal subsite function, we carried out missing base interference footprinting (Brunelle and Schleif 1987). This method allows the importance of a single base at any promoter position to be determined by the selective depletion of templates lacking that base from the population of fragments able to form open complexes. Modified promoter fragment populations that contained on average one missing thymine per molecule were prepared by low frequency substitution of dUTP for TTP (Fig. 1B) followed by removal of the uracil base by use of uracil DNA glycosylase (UDG; Devchand et al. 1993). Only the base was removed, leaving the phosphodiester backbone intact.

Labeled promoter fragments containing missing bases were then incubated with RNAP. Because the α–UP element interaction increases the rate of association of the promoter with RNAP (Rao et al. 1994), the RNAP binding reaction was carried out at low RNAP concentration and in 20-fold molar excess of modified promoter fragment relative to RNAP, conditions in which complex formation was strongly dependent on the α–UP element interaction. The resulting promoter–RNAP complexes were separated from unbound DNA by electrophoresis on a nondenaturing gel, purified, subjected to strand cleavage at the abasic sites, and then fractionated on a high resolution denaturing gel. Missing bases that interfered with RNAP binding were under-represented in the strand cleavage profiles of bound DNA relative to their representation in the DNA population that was not incubated with RNAP.

A typical missing T interference experiment carried out on a promoter with a consensus proximal subsite is shown in Figure 2. Each of the thymine positions in the proximal subsite (eight of nine positions on the template strand) was severely depleted in the RNAP-bound sample, indicating that a missing T at each of eight positions in the UP element interfered with RNAP binding. The strongest interference signals occurred in the center of the T tract (−41 to −45), whereas missing Ts upstream of the UP element sequence did not interfere with RNAP binding (Fig. 2 and data not shown). Identical results were obtained with an RNAP preparation containing only one α-CTD (α/αΔ1 heterodimeric RNAP; data not shown), indicating that, as expected from previous work (Estrem et al. 1999), missing Ts interfering with wild-type RNAP binding to the proximal subsite are attributable to interactions involving only one α subunit. No interference signals in the UP element were observed with RNAP lacking both α-CTDs (data not shown).

Missing Ts at some positions on each strand in the core promoter region also strongly affected RNAP binding (Fig. 2 and data not shown; summarized in Fig. 6, below). All missing Ts in the −35 hexamer and some in the spacer region between the −10 and −35 hexamers reduced RNAP binding. In contrast, most missing Ts in the −10 hexamer strongly stimulated RNAP binding (all but nontemplate strand position −9), most likely because

RNAP α subunit binding to the DNA minor groove
unpaired nucleotides facilitate strand opening (see Fig. 2 and Discussion).

Missing purine interference experiments (data not shown) with templates prepared by limited acid depurination [Maxam and Gilbert 1980], were used to examine the interaction of RNAP with the nontemplate strand. The conclusions from these experiments were similar to those from the missing T experiments: missing bases at the eight UP element positions that are purines, and at purine positions in the spacer, but not at control positions further upstream, resulted in reduced RNAP binding (for summary, see Fig. 6, below). Missing purines at positions −41 and −42 reduced RNAP binding the most. As reported previously [Li and McClure 1998], and as found in the missing T experiments, most missing purines in the −10 region [all but nontemplate strand −13] stimulated RNAP binding.

**Major groove modifications affect RNAP binding to the core promoter but not to the UP element**

The missing-base interference experiments indicated that each base in the UP element proximal subsite was important for α binding and suggested that bases in the center of the subsite (−41 to −45) are most critical. However, these results did not distinguish which base surfaces (major groove, minor groove, or both) are contacted by α, or alternatively, whether the interference signals might derive solely from indirect effects of missing bases on DNA backbone structure.

To determine whether α makes base contacts in the major groove, interference footprinting was carried out with promoter fragments containing adenine or thymine bases modified on their major groove surfaces. Modifications were introduced by limited incorporation of the nucleotide analogs dUTP or dA*TP [7-deaza-7-nitro dATP; Fig. 1B]. U is identical to T except that it lacks the prominent major groove Cs methyl group. A* [Min et al. 1996] disrupts contacts to adenine at position 7 through steric exclusion and at position 6-NH through intramolecular hydrogen bonding. Both U and A* could potentially disrupt protein binding directly through loss of functional groups that contact amino acids, but, in addition, the adenine modification could potentially disrupt protein binding by steric hindrance. dUTP or dA*TP have not to our knowledge been used for footprints of RNAP, although they have been used in footprinting studies of other proteins [Pu and Struhl 1992; Devchand et al. 1993; Min et al. 1996].

To carry out dUTP interference footprints, RNAP–promoter complexes were formed with templates containing, on average, one U substitution per molecule (Fig. 3A,B). Complexes were then gel isolated, and the DNA was purified, subjected to strand cleavage at modified bases, and fractionated on a denaturing gel. Uracil substitution at any of the eight thymine positions in the UP element had very little effect on RNAP binding (template strand, Fig. 3B), suggesting that α does not make critical contacts with major groove thymine methyl groups in the UP element. In contrast, U substitutions at any of three core promoter positions (−36, −35, and −14) strongly reduced or eliminated RNAP binding, and U substitutions at either of two other core promoter positions (−31 and −9) also reduced RNAP binding (Fig. 3A,B, for summary, see Fig. 6, below).

Similar experiments were carried out with promoter fragments containing A* substitutions (Fig. 3C,D). As in the U substitution experiments, A* at UP element positions had little or no effect on RNAP binding (nontemplate strand, Fig. 3C). However, A* substitutions had large effects on RNAP interactions with the core promoter. A* substitutions reduced binding at five of the six A residues in the two strands of the −10 hexamer (Fig. 3C,D, for summary, see Fig. 6, below); A* at nontemplate strand position −10 virtually eliminated binding of RNAP, and at template strand position −33 (in the −35 hexamer) strongly enhanced RNAP binding (see Discussion).

In summary, the U and A* interference experiments indicate that major groove base surfaces in the UP element play little if any direct role in RNAP binding, although they do play a major role in interactions between RNAP and the −10 and −35 elements (presumably by
contacting the σ subunit; see Fig. 6, below and Discussion).

Distamycin inhibits σ binding and UP element function

Hydroxyl radical footprints with RNAP and with purified σ previously suggested that σ contacts the DNA backbone along the minor groove in the UP element proximal subsite (Newlands et al. 1991; Estrem et al. 1999). To investigate the role of the minor groove in σ binding, we used reagents that bind in the minor groove (distamycin) or that modify the N3 of adenine, which is located in the minor groove (dimethylsulfate, DMS). Distamycin, a small, well-characterized oligopeptide that binds within the minor groove of A+T-rich DNA (Coll et al. 1987; Churchill et al. 1990), was used to determine whether insertion of a small molecule in the minor groove of the UP element could compete with and prevent σ binding to the promoter. Binding sites for distamycin in the promoter fragment were first characterized by hydroxyl radical footprinting to determine whether, as expected, the drug would bind preferentially to the UP element (Fig. 4A). Specific, but partial, protection in the UP element (positions −42 to −46) was observed at 125 nM distamycin. Protection of this site was more complete at 250 nM, but, at 500 nM, protection by distamycin also occurred at three other sites: in the −35 hexamer, in an A+T-rich region of the spacer (−20 to −24), and in the −10 hexamer (−10 to −14). The length of the protected regions, 4–5 bp, is consistent with the length reported previously in hydroxyl radical footprints with distamycin (Churchill et al. 1990).

Because σ and distamycin protect the same UP element positions against hydroxyl radical cleavage, the identity of the molecule binding to the UP element cannot be distinguished by footprinting when both the protein and the drug are present. However, purified σ–UP element complexes migrate differently on gels from free or distamycin-bound DNA fragments. Therefore, we

Figure 3. Interference footprints with the base analogs uracil (U; A, B) or 7-deaza-7-nitro adenine (A*; C, D). Promoter fragments [rrnB P1 with UP element 4549], containing, on average, one modified base per molecule, were labeled on the nontemplate strand (A, C) or the template strand (B, D). RNAP–promoter complexes were isolated, and samples were examined by gel electrophoresis after strand cleavage at modified bases. Superimposed profiles from scans of normalized lanes are aligned with gels; RNAP-bound samples (+) are indicated by solid blue lines [uracil, (A, B)] or green lines [A*; C, D], and control samples without RNAP (−) are black dotted lines. UP element, −35 and −10 hexamer sequences are shown, with all T positions in blue (A, B), and all A positions in green (C, D). Modified bases that affected RNAP binding are indicated: [blue or green circles] interference with binding, [blue or green arrows] stimulation of binding. Symbol sizes correlate with the approximate degree of interference or stimulation observed.
used electrophoretic mobility-shift assays to determine the effect of distamycin on binding of purified \( \alpha \) to a promoter fragment.

The mobility shift assay was carried out with a concentration of \( \alpha \) (2 μM dimer) where \( \alpha \) bound specifically to the UP element (Ross et al. 1993). Formation of the \( \alpha \)-DNA complex was inhibited by distamycin (Fig. 4B). Half-maximal inhibition occurred at \( \sim 200 \) nM distamycin, a concentration where the drug bound specifically to the UP element and not to other regions of the promoter fragment (Fig. 4A). We conclude that distamycin competes with \( \alpha \) for binding to the UP element, consistent with the model that \( \alpha \) binds to the UP element minor groove. (At higher \( \alpha \) concentrations, where \( \alpha \) also binds to other DNA sites, higher concentrations of distamycin were required to inhibit \( \alpha \) binding in mobility shift assays; data not shown).

Figure 4C shows the effects of a range of distamycin concentrations on UP element-dependent transcription in vitro. We examined expression from three different promoters, the \( rrnB \) P1 core promoter with a consensus proximal subsite, the \( rrnB \) P1 core promoter without an UP element, and the RNA I promoter (which also lacks an UP element) encoded by the plasmid vector. Transcription from the UP element-containing promoter was inhibited at concentrations of distamycin from 62.5 to 250 nM, whereas transcription from the promoters lacking UP elements was not affected, correlating well with the concentrations at which distamycin bound specifically to the UP element (Fig. 4A) and at which distamycin specifically inhibited \( \alpha \) binding (Fig. 4B). We conclude that distamycin binding in the minor groove blocks UP element function in transcription. At higher distamycin concentrations (\( \geq 500 \) nM), transcription was inhibited from all three promoters (Fig. 4C), consistent with the observed binding of distamycin to core promoter sites. At these core promoter sites, high concentrations of distamycin may compete with RNAP for binding to DNA (e.g., at positions in the –10 region or in the spacer region that are protected by both distamycin and by RNAP in hydroxyl radical footprint; Fig. 4A and Newlands et al. 1991).

The UP element minor groove is protected by \( \alpha \) CTD from modification by DMS

The distamycin experiments (Fig. 4) suggest that the DNA minor groove plays an important role in the binding of \( \alpha \) to the UP element. To determine whether \( \alpha \) contacts (or is in very close proximity to) specific bases within the minor groove, we asked whether prebound RNAP or purified \( \alpha \) subunit protected the N3 position of adenines in the UP element minor groove (see Fig. 1B) against methylation by DMS. N3A methylation, as well as N7G methylation in the major groove, is detectable by elimination of the modified base at high temperature, followed by strand cleavage at the abasic site (Maxam and Gilbert 1980, see Materials and Methods).

DMS protection footprints were carried out with a promoter in which the UP element contained adenines at

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**Figure 4.** Distamycin experiments. (A) Scans of hydroxyl radical footprints of distamycin bound to a fragment containing \( rrnB \) P1 with UP element 4547. Profiles of distamycin-bound fragment [red, 125 nM distamycin; green, 250 nM; blue, 500 nM] are superimposed on profiles from the control fragment [no distamycin, grey lines]. RNAP recognition elements (UP element, –35 and –10 hexamers), indicated by rectangles, are aligned with the scans. (B) Effect of pre-bound distamycin on the binding of purified \( \alpha \) to a fragment containing \( rrnB \) P1 with UP element 4547. \( \alpha \) binding was determined by a gel mobility shift assay, the percent of input DNA bound by \( \alpha \) (averages with standard deviation from three separate experiments) is shown as a function of distamycin concentration. (C) Effect of pre-bound distamycin on transcription in vitro. Transcription from \( rrnB \) P1 with UP element 4547 [filled circles], \( rrnB \) P1 with no UP element [open circles], and from the RNA I promoter [filled triangles] was quantified from gel scans, and is expressed as a percentage of transcription from each promoter in the absence of distamycin. Results are the average [with range] from two independent experiments.
seven of nine positions in the nontemplate strand. Profiles from footprints with purified α, with wild-type RNAP, and with RNAP lacking the αCTD (αΔ235) are illustrated in Figure 5. In each case, the profile is compared with a control profile lacking protein. Methylation of several N3A positions was significantly reduced in the presence of purified α [Fig. 5A] or wild-type RNAP [Fig. 5B]. Protection of the UP element from modification by DMS was not observed when RNAP lacked the αCTD (αΔ235 RNAP; Fig. 5C). Adenine residues are less reactive to DMS than guanine residues, and, therefore, the peak heights at A residues are lower than those for G even in the absence of bound protein, which tends to de-emphasize protection signals [Maxam and Gilbert 1980]. However, the N3A protections by RNAP and α were observed reproducibly in many experiments with both native and reconstituted RNAPs [for summary, see Fig. 5D, Fig. 6]. We determined the fraction of the N3A signals protected by RNAP [Fig. 5E]. RNAP reduced the methylation of N3s at positions −41 and −42 by −40%, a value that is consistent with partial protection signals observed in other DMS footprints [e.g., Johnsrud et al. 1978; Craig and Nash 1984; Newlands et al. 1991]. Protection of N3A by RNAP was also observed in UP elements from the native rrrB P1 and rrrD P1 promoters [data not shown].

The major groove N7G at −40 was partially protected in RNAP complexes [Fig. 5]. This protection was also observed [although to a lesser extent] with purified α and with RNAP lacking the αCTD (Fig. 5A,C). Thus, it is not clear whether the −40 N7G protection observed with wild-type RNAP is attributable to α or to σ interactions with the DNA or if it results from interactions between the two subunits in the vicinity of −40 (see Discussion).

Methylation of several N3A and N7G positions in the core promoter region was affected by the presence of bound RNAP [Figs. 5, 6; data not shown]. The observed protections or enhancements are consistent with those reported previously for DMS footprints of this and other promoters [Johnsrud 1978; Siebenlist et al. 1980; Duval-Valentin and Ehrlich 1986; Newlands et al. 1991]. The results from the DMS protection footprints, from the U and A* interference footprints, and from previously reported hydroxyl radical footprints on this promoter are summarized in Figure 6.

**Discussion**

**DNA features critical for α binding**

Binding of one αCTD to the UP element proximal subsite produces a very small footprint whose features suggest interactions with the phosphodiester backbone and with bases in the minor groove [Fig. 7A]. The DMS footprinting data indicate that α is in very close proximity to position N3 of adenine bases in the UP element minor groove. Protection of N3A positions upstream of the −35 hexamer was reported in early DMS footprints of RNAP on other promoters [e.g., Siebenlist et al. 1980; Duval-Valentin and Ehrlich 1986], although the role of α in
Promoter recognition was not known at the time of those experiments. X-ray structures of minor groove-binding proteins in complex with DNA have confirmed that protection of N3A positions in DMS footprints correlates with minor groove contacts [e.g., Hin recombinase (Glasgow et al. 1989; Feng et al. 1994) and IHF (Craig and Nash 1984; Rice et al. 1996)].

Our experiments using distamycin, which binds snugly in the minor groove of A+T-rich DNA (Coll et al. 1987), support the role of minor groove interactions in α binding. Distamycin protected essentially the same set of UP element residues as α in our hydroxyl radical footprinting experiments (Estrem et al. 1999; Figs. 4,6) and inhibited α binding to UP element DNA. Distamycin has been shown previously to interfere with minor groove binding of other proteins [e.g., HMG Y/I; Reeves and Nissen 1990].

Several backbone positions on each strand in the UP element proximal subsite are protected by RNAP from hydroxyl radical cleavage (Estrem et al. 1999; Figs. 4,6). Hydroxyl radicals react primarily with the hydroxyls on positions C4 and C5 on the minor groove side of deoxyribose (Balasubramanian et al. 1998). Therefore, the UP element base and backbone positions protected by α are located in very close proximity (Fig. 7A). Although not tested in our experiments, the O2 positions of thymine bases in the minor groove would also be candidates for direct hydrogen bond contacts with α.

Our results suggest that α does not make significant major groove contacts. Substitution of U or A* for T or A, respectively, in the UP element had little if any effect on RNAP binding to the promoter (Fig. 3). Although the partial protection of major groove position N7G at −40 against DMS methylation (Fig. 5) may indicate close proximity of RNAP to this position, other considerations suggest that the identity of this residue is not critical to UP element function. G is not strictly conserved at −40 in UP element proximal subsites (Estrem et al. 1999), and no steric hindrance to RNAP binding was observed when A* was substituted for A at this same position in the UP element (Fig. 3). In addition, although some degree of protection of N7G at −40 was observed with purified α, protection was also observed with RNAP lacking the αCTD. Thus, it is not clear whether it is α or σ that is in close proximity to the N7G at −40 in the context of RNAP holoenzyme. Furthermore, the reduced methylation of the N7G at −40 could be attributable to altered DNA structure resulting from RNAP binding rather than from direct protein contact. The latter interpretation is supported by the strong enhancement of DNase I cleavage at position −38 observed in footprints of RNAP on rRNA P1 and other promoters (Fig. 7A; Gourse 1988; Ross et al. 1993; Craig et al. 1995).

Although our footprints indicated that α contacts only a small subset of base surfaces, a missing base at any position in the UP element interfered very strongly with RNAP binding (Fig. 2). The strongest of the missing base signals on the non-template strand [−41 to −43] coincide with positions protected by RNAP in DMS footprints, suggesting that the effects of missing bases reflect, in part, the loss of minor groove base contacts. However, distortion of the normal structure of the UP element proximal subsite could also account for the effect of the missing bases, since structural changes can occur up to 4 bp from an abasic site (Wang et al. 1997). The proximal subsite contains a long A tract (7–8 residues), and unusual structural features of the A tract [e.g., narrow minor groove width and/or propeller twisting of bases; Shatzky-Schwartz et al. 1997] could play an important role in recognition by α. Thus, at least some of the interference to RNAP binding caused by missing bases could be attributable to effects on a structure important to recognition by σ, rather than to loss of direct base contacts.

In contrast with our results, it was reported that missing bases in the UP elements of the ace or the rRNA P1 promoters stimulated [rather than inhibited] RNAP binding (Negre et al. 1997). We suggest that the A+T-rich UP element region containing an abasic site may have
been recognized as a -10 hexamer rather than as an α-binding site, accounting for the observed stimulation of RNAP binding in these experiments. We and others (Li and McClure 1998) have observed that missing bases
in the −10 hexamer strongly stimulate RNAP binding
[Figs. 2,6]. The results of our footprinting studies are consistent
with the effects on transcription of mutations in the UP
element proximal subsite [Estrem et al. 1999]. Substitution
of C:G for A:T pairs at several UP element positions
(−41 to −45) reduced transcription in vivo and in vitro,
with the strongest effects (>10-fold) occurring at posi-
tions −41, −42, and −43. The 2-amino group of guanine at
these positions could sterically hinder α binding in the
minor groove; alternatively, effects on α binding of a C:G
substitution could result from disruption of A-tract
structure [Koo et al. 1986]. Interestingly, a T:A rather
than an A:T pair at position −42 [in UP element 4542;
Estrem et al. 1999] did not reduce UP element function.
This substitution would preserve hydrogen bonding ca-
pabilities in the minor groove [with the O2 of thymine].

A model for α CTD interaction with an UP element
subsite

Our results indicating that α binds to the UP element
minor groove are consistent with the recent observation
that the α CTD contains an [HhH]2 domain. (Shao and
Grishin 2000). On the basis of this observation, on our
biochemical data, and on our previous genetic identifi-
cation of the α CTD DNA-binding surface [Gaal et al.
1996; Murakami et al. 1996], we suggest a preliminary
model for how α might interact with its binding site [Fig.
7C].

The HhH motif was identified initially in the struc-
ture of the DNA repair enzyme Endonuclease III [Thayer
et al. 1995], and has since been identified in DNA-bind-
proteins involved in replication, recombination, and
repair [e.g., DNA polymerases, ligases, and glycosylases;
Doherty et al. 1996; Shao and Grishin 2000]. The HhH
motif contains a pair of anti-parallel α helices connected
by a hairpin loop that are present in either one or two
copies [the [HhH]2 domain, Shao and Grishin 2000].
DNA binding, in examples considered thus far, is not
sequence specific [Doherty et al. 1996]. In the rat DNA
polymerase β-DNA complex [Pelletier et al. 1996] and
the E. coli RuVA-DNA complex [Ariyoshi et al. 2000],
amino acid residues in the loop and in the amino-termi-
nus of the second α helix make nonspecific hydrogen
bond contacts to the DNA backbone. RuVA binds across
the minor groove of duplex DNA, with each of the re-
peated HhH motifs in a RuVA monomer contacting the
backbone of one of the two DNA strands.

The RuVα CTD contains a short nonstandard helix
[NSH] followed by four α helices [Leon et al. 1995; Gaal et
al. 1996]. In the [HhH]2 domain interpretation of CTD
[shown in schematic form in Fig. 7B], the first HhH motif
contains the NSH residues 253–257, a loop 258–263,
and a second α helix 264–272 referred to as α helix 1.
The second HhH contains α helix 3 [286–292], a loop
293–297, and α helix 4 [298–309]. The two HhH motifs
are separated by α helix 2. The positions of substitutions
in α CTD that result in strong defects in DNA binding
[Gaal et al. 1996; Murakami et al. 1996] affect residues

that, by analogy to [HhH]2 domains in other proteins
[Rafferty et al. 1998; Ariyoshi et al. 2000], would be pre-
predicted to make DNA contacts (e.g., R265 in the amino
terminal of the second helix in HhH1, G296 in the loop,
and K298, S299 in the amino terminal of the second
helix in HhH2).

Based on these considerations, and by analogy to the
RuVA-DNA structure [Ariyoshi et al. 2000], we suggest
that each of the HhH motifs in the α CTD makes back-
bone contacts with one of the DNA strands, at the posi-
tions that are protected by the α CTD in hydroxyl radi-
cal footprints [Fig. 7C]. The length of the α-binding site is
similar to the length of DNA covered by one RuVA
monomer [Ariyoshi et al. 2000].

Unlike other HhH-motif proteins, the α CTD demon-
strates sequence specificity in DNA binding. We suggest
that this sequence specificity could derive from (at least)
two characteristics of the UP element DNA sequence.
First, the narrow width of the minor groove in an A tract
[the preferred α-binding site] could be important for si-
multaneous binding of α to both DNA strands. It has
been proposed previously that effects of sequence on
DNA structure could influence binding of HhH motif
proteins [Rafferty et al. 1998].

Second, our data suggest that the specificity of α
binding to an A-tract sequence derives in part from base con-
tacts in the minor groove [Fig. 5]. These contacts [to N3A
and potentially to O2T] could occur through hydrogen
bonds with the basic side chains of residues R265 and
K298. Base-specific contacts in the minor groove of A+T-
rich DNA sequences have been demonstrated for a num-
ber of other DNA-binding proteins. Frequently these in-
teractions are made by short peptide motifs containing
arginine residues that contact bases in the minor groove
and the DNA backbone along the minor groove [e.g.,
AT-hooks in the chromatin-associated protein HMG-I(Y)
[Huth et al. 1997], and extended arm sequences in the
Hin recombinase [Feng et al. 1994] and in homeodo-
mains [Kissinger et al. 1990]]. Because several substitu-
tions in the α CTD cause defects in both specific and
nonspecific DNA binding [Savery et al. 1998], we suggest
that the same DNA-binding motif is responsible for in-
teractions with UP element and non-UP element DNA
sequences.

In the model for the α–UP element interaction pre-
sented in Figure 7C, we have chosen a specific orienta-
tion for α CTD with respect to the UP element sequence,
although in other contexts the orientation could be re-
versed. The orientation shown is suggested by the results
of genetic experiments that implicate α residues 259 and
261 in interactions with the σ subunit at promoters con-
taining the UP element proximal subsite [Gourse et al.
2000]. This orientation was also suggested by the iden-
tification of residues potentially involved in σ–α inter-
actions in the lac promoter–CRP–RNAP complex [Busby
and Ebright 1999; Gourse et al. 2000].

α binding to UP elements with two subsites

In the present study, we focused on interactions of α
with a single consensus binding site, the proximal subsite. However, we have also performed footprinting experiments on full (i.e., two subsite) UP elements. The distal subsite in the full UP element responded similarly in protection and interference experiments to the proximal site by itself (W. Ross and R.L. Gourse, data not shown). However, the proximal subsite in the full UP element responded differently from the proximal subsite in the absence of the distal subsite. In the context of the full UP element, missing bases in the proximal subsite did not interfere with RNAP binding (W. Ross and R.L. Gourse, data not shown), and the proximal subsite was not fully protected in hydroxyl radical footprints (Newlands et al. 1991; Estrem et al. 1998).

These results suggest that the proximal subsite plays a different role when alone than it does in a full UP element. These results are also consistent with three lines of genetic evidence: (1) Proximal subsite sequences identified in binding-site selections were not identical in the presence versus absence of the consensus distal subsite (Estrem et al. 1998, 1999). (2) Proximal subsite mutations had larger effects on transcription when the consensus distal subsite was absent (Estrem et al. 1998; S. Estrem and R.L. Gourse, unpubl.). (3) Certain oCTD and αCTD substitutions reduced transcription from promoters containing only a proximal subsite, but not from promoters with full UP elements (W. Ross, A. Mertens, B. Paul, D. Schneider, J. Toplin, M. Burgess, and R.L. Gourse, unpubl.).

These results suggest that the two subsites do not contribute equally to the binding of RNAP in promoters containing full UP elements. Possible explanations (see also Estrem et al. 1999) could include: (1) Binding of αCTD to the proximal subsite could be weaker, less specific, or nonspecific in the context of both subsites. (2) The proximal subsite in the context of a full UP element could serve primarily a structural role, rather than a role in α binding. (3) The longer, symmetric distal subsite sequence could be a binding site for both α subunits in the context of a full UP element, thereby reducing or excluding occupancy of the proximal subsite by α.

RNAP contacts with the −35 hexamer

Our interference footprints yielded considerable information about the role of specific major groove base surfaces in binding RNAP to the core promoter. These data, together with the DMS protection data, confirm and extend previous information about the interaction of the σ subunit with the −35 and −10 elements.

Of the 12 bases present in the −35 hexamer of rrrB P1 [six on each DNA strand], information about 10 bases was obtained with the base-specific reagents used, and of these 10, five appear to be in close proximity to or in contact with RNAP (Figs. 6, 7). Particularly strong interference signals were observed when U was substituted for T at positions −35 and −36, indicating probable interactions of σ with the C5 methyl groups of these thymines that are critical for RNAP binding. Because A* substitutions at the same positions on the template strand did not interfere with RNAP binding, we suggest that the strong conservation of T:A pairs at these positions in promoter −35 hexamers (Hawley and McClure 1983) reflects interactions with the thymine bases.

These results are consistent with previous observations. Mutations of the thymines in the nontemplate strand at the upstream end of the −35 hexamer have strong phenotypes (Youderian et al. 1982), and 5-bromo-uracil substitutions at these positions in lacUV5 are protected by RNAP against UV-induced strand cleavage (Siebenlist et al. 1980). The region of σ70 that interacts with the −35 hexamer (region 4.2) is thought to contain a helix–turn–helix motif (HTH; Record et al. 1996; Gross et al. 1998). Although we do not know which amino acid residues in region 4.2 interact with the thymine methyl groups at −35 and −36, they are likely to be located in the HTH recognition helix, carboxy-terminal to the arginine residues at positions 584 and 588 that are implicated in contacting the central region of the −35 hexamer (for review, see Gross et al. 1998). RS88 was proposed to contact position −34. However, because the major groove surface of −34 G does not appear to be in contact with RNAP (methylation of this G is increased, not reduced, by the presence of prebound RNAP; Fig. 5; Siebenlist et al. 1980), we suggest that the template strand C, rather than the nontemplate strand G, may contact RNAP and be responsible for the conservation of the G:C pair at −34.

Our results suggest that σ also contacts three template strand bases at the downstream end of the −35 hexamer. A* at position −33 strongly stimulated RNAP binding. This is a nonconsensus position in rrrB P1 [T:A rather than A:T], and the presence of the 7-nitro group on the modified base might provide a contact to RNAP that serves as an alternative to that provided by thymine at the same position in a consensus −35 hexamer, even though the nitro and methyl groups are chemically dissimilar. Additional template strand contacts in the −35 hexamer are suggested by partial protection of NTG at −32 against methylation by DMS (Newlands et al. 1991; see also Johnsrud 1978; Duval-Valentin and Ehrlich 1986), and by partial interference resulting from U substitution at −31. Hydroxyl radical footprints indicate that RNAP also interacts with the DNA backbone along the adjacent minor groove in the spacer at the downstream edge of the −35 hexamer (Newlands et al. 1991; Figs. 6, 7). Hydroxyl radical protections (Newlands et al. 1991) and the effects of missing bases at several positions in the spacer region, particularly around −20 (Fig. 6), suggest that the conformation of this region may be important for RNAP binding.

RNAP contacts with the −10 hexamer

In the −10 hexamer, major groove modifications at several positions interfered with the formation of a stable
open complex at rrrB P1. Because open complex formation involves recognition of the −10 region as duplex DNA, followed by strand separation and sequence-specific interaction with the nontemplate strand (Roberts and Roberts 1996; Marr and Roberts 1997; Qiu and Helmann 1999; Fenton et al. 2000), the interference signals observed in our footprints could result from disruption of contacts with duplex DNA, single-stranded DNA, or both. Our footprinting results should be useful for future interpretation and modeling of the interaction of σ region 2.4 with the −10 hexamer.

Specifically, we found that U substitution at two of the three nontemplate strand thymines reduced RNAP binding, indicating that σ may directly contact the C5 methyl group of these Ts (rrrB P1 positions −14 and −9, corresponding to positions −12 and −7 in lacUV5 and λP_r'; Marr and Roberts 1997; Fenton et al. 2000). A* substitution for five of the six adenines in the −10 hexamer interfered with RNAP binding. Because A* contains a C7-nitro group in place of N7, interference could result from loss of direct contacts or from steric hindrance (Min et al. 1996). In either case, our data indicate that RNAP is in close proximity to these bases.

The inhibitory effect of A* was particularly strong at nontemplate strand position −10 (−8 in other promoters), although this is not one of the most highly conserved −10 hexamer positions (Hawley and McClure 1983). A* at −10 could conceivably sterically hinder the approach of RNAP to the adjacent highly conserved thymine or might interfere with critical base stacking interactions of the separated strand (Qiu and Helmann 1999). Strong effects of A* substitutions were also observed at rrrB P1 template strand positions −14 and −12 (−12 and −10 in other promoters). Although an A at nontemplate strand position −13 (−11 in other promoters) is very highly conserved and plays a critical role in promoter function, the substitution of A* at −13 had a relatively small effect on RNAP binding (Fig. 3). A template strand hexamer positions −13 (−11 in other promoters) is very highly conserved and plays a critical role in promoter function, the substitution of A* at −13 had a relatively small effect on RNAP binding (Fig. 3). A is important for single-stranded oligonucleotide binding and fork-junction binding (Marr and Roberts 1997, Guo and Gralla 1998; Qiu and Helmann 1999) and is thought to nucleate strand opening, perhaps through base stacking interactions with aromatic residues in σ region 2.4 (Fenton et al. 2000; Panaghie et al. 2000). Recognition of −13A by σ apparently does not involve critical contacts to the functional groups altered by the A* modification.

Missing bases at all but two of the positions tested in the −10 hexamer (all except −13A, −9T) stimulated RNAP binding, presumably by facilitating the strand separation process. These results are consistent with those reported previously for other promoters (Li and McClure 1998). In addition, a missing base at template strand +1 strongly interfered with RNAP binding to the rrrB P1 promoter (data not shown). This negative effect reflects the requirement for incorporation of the initiating two nucleotides, ATP and CTP, to stabilize the rrrB P1 open complex, and is not a phenomenon general to promoters with longer-lived open complexes; that is, the effect of a missing base at +1 of rrrB P1 does not indicate a requirement for a specific base contact to RNAP at this position.

Conclusion

The work reported here provides information about the functional consequences of specific DNA–protein contacts in the initiation complex. Such data will be important for interpretation of atomic level structures of the promoter–RNAP complex that may become available in the future. We have shown that the α subunit of RNAP binds to UP element DNA using minor groove as well as backbone contacts, and we have proposed a model for αCTD binding to DNA. These data should also facilitate modeling of αCTD interactions with DNA in promoter complexes containing transcription activators. Our results suggest further that the contacts made by α to the UP element and by σ to the −35 hexamer are in very close proximity. Thus, αCTD–σ interactions could play a role in the function of the UP element proximal subsite. Finally, our results identify functional groups in the −10 and −35 hexamers that are involved in the interaction with the σ subunit and thus increase the resolution of our picture of the RNAP–core promoter interaction.

Materials and methods

Plasmid DNAs

DNA fragments containing rrrB P1 promoters with different sequences in the UP element region were inserted as EcoRI (−66 with respect to the transcription start site) to HindIII (+50) fragments into pRLG770 (Estrem et al. 1999). Plasmids contained consensus proximal subsite 4547 [−46] 5′-AAAAAGTA-3′ [−38] in pRLG4213, consensus proximal subsite 4549 [−46] 5′-AAAAAGAA-3′ [−38] in pRLG3289, full consensus UP element 4192 [−59] 5′-GGAAAAATTTTTTTTCAAAAGTA-3′ [−38] in pRLG3278, or an upstream sequence lacking UP element function (SUB) in pRLG4210. DNA sequences flanking the promoters were described previously (Estrem et al. 1998, 1999).

RNAP and α purification

Wild-type native RNAP was a gift from Dr. Robert Landick (University of Wisconsin). Wild-type α subunit was purified as described (Gaal et al. 1996). RNAPs containing wild-type or mutant α subunits lacking the carboxy-terminal domain (αΔ35S) and containing an amino-terminal hexahistidine tag were reconstituted from isolated subunits (Gaal et al. 1996), purified by nickel chromatography (Qiagen) and by centrifugal filtration (Centricon YM-100, 100,000 MW cutoff). RNAPs heterodimeric for α containing one wild-type α and one αΔ35S were purified as described (Estrem et al. 1999).

Missing base interference footprinting

Promoter fragments were prepared by 15 cycles of PCR amplification using primers that flanked the promoter in the vector sequence (upstream primer 5′-CGCGGATCCGATATCACGGGCCTTTCG-3′; downstream primer 5′-GGCTGACTGCGTTTCCACTTC-3′). PCR reactions (50 μL) contained 0.4 μg DNA fragments and 0.1 μL Taq polymerase (Promega).
of plasmid DNA, 20 pmol of each primer, 200 µM amounts of each dNTP [Pharmacia], 1.5 mM MgCl₂, and 2.5 units of Taq DNA polymerase (Promega) in a buffer supplied by the manufacturer. For preparation of fragments containing missing thymine bases, the PCR reactions also contained 2 µM dUTP (Phar- 

macia; Devchand et al. 1993).

PCR products were purified by use of a Qiagen PCR Purifica-

tion Kit (Qiagen), digested with the appropriate restriction enzyme [HindIII (+50) for nontemplate strand labeling or BamHI [primer-encoded] for template strand labeling], and 3'-end-la-

beled with Sequenase [USB] and [α-32P]dATP for HindIII sites, or [α-32P]dGTP for BamHI sites. Labeled fragments were gel puri-

fied, eluted by diffusion into 0.2 M NaCl, 20 mM Tris-Cl (pH 7.4), 1 mM EDTA, and concentrated by use of Elutip D columns [Schleicher and Schuell]. To prepare fragments with, on average, one missing thymine base per molecule, end-labeled fragments containing dUTP were then treated with uracil DNA glyco-

sylase (UDG; NEB; 0.75 units for 10 min at 37°C in 100 µL with 

buffer supplied by the manufacturer) to remove the uracil base, leaving the DNA backbone intact. To prepare fragments with, on average, one missing purine base per molecule, end-labeled fragments [prepared without modified nucleotides] were treated with formic acid as described previously [Maxam and Gilbert 1980].

To carry out the missing base interference footprint, RNAP [0.5 nM] was incubated with a 20-fold excess of end-labeled 

DNA fragment (10 nM) containing either a missing thymine base or a missing purine, for 5 min at 22°C in 100 µL of Tris-Cl (pH 7.9), 30 mM KCl, 10 mM MgCl₂, 1 mM DTT, 500 

µM ATP, 50 µM CTP. Heparin (final 10 µg/mL) was added, and RNAP-promoter complexes (~5% of input DNA) were sepa-

rated from unbound DNA by electrophoresis on a 5% acryl-

amide gel in 0.5 

Tris-Borate-EDTA (TBE). Because DNA was 

kept in excess to promote competition for RNAP, the unbound 

promoter fraction did not provide useful information and was 

not analyzed. Control DNA used for comparison to the RNAP-

bound population was the input DNA population without in-

cubation with RNAP.

RNAP-bound DNA was eluted by diffusion, and purified by use of Elutip D columns as described above. Then, strand cleav-

age was carried out at the abasic sites by incubation in 100 µL of 1 M piperidine at 90°C for 30 min. Fragments were ethanol precipitated, then separated by electrophoresis on 9.5% acryl-

amide–7 M urea gels. Dried gels were scanned by PhosphoIm-

aging using a Storm system (Molecular Dynamics). Data points 
in each lane were normalized to correct for differences in load-

ing. Normalization factors were derived from comparisons of 
portions of the profiles that were distant from the promoter and 
assumed not to be in contact with RNAP. Normalized profiles 
were graphed using Sigma Plot [Jandel Scientific].

Modified base interference footprinting
Promoter fragments for modified base interference footprinting were prepared by PCR as described above, but reactions also contained either 2 µM dUTP (Pharmacia) for modified thymine 

containing fragments, or 30 µM 7-deaza-7-nitro dATP (da *TP; 

see below for synthesis) for modified adenine containing frag-

ments. Fragments were 3'-end-labeled and gel purified as de-

scribed above.

RNAP-promoter fragment complexes were formed under conditions described above for missing base interference foot-

printing, gel isolated and the DNA eluted and purified as de-

scribed. Strand cleavage at modified adenine [A*] bases was then carried out by incubation with 1 M piperidine, as described above. DNA fragments containing dUTP were treated with 

UDG [above] to remove the uracil base prior to the strand cleav-

age at the resulting abasic sites (see above). Then, fragments 
were analyzed by denaturing gel electrophoresis as described.

Synthesis of 7-deaza-7-nitro-2'-deoxyadenosine 

triethylammonium 5'-triphosphate (da *TP)

7-Deaza-7-nitro dATP (da *TP) was synthesized using modifi-
cations of the procedure described previously [Min et al. 1996]. A suspension of 7-deaza-7-nitro-2'-deoxyadenosine [Min et al. 1996; Secla et al. 1997; 23.2 mg, 78.4 µmoles, coevaporated twice from pyridine and dried under high vacuum overnight] in 

trimethylphosphate [0.32 mL] was treated at 0°C with POCl₃ 

(8.1 µL, 86.3 µmoles), and stirred for 20 min at 0°C to produce a yellow solution. This solution was treated dropwise with 0.5 M 

([Bu₃NH]H₂P₂O₇, solution in dimethylformamide [0.78 mL, 

39.0 mL] of ice-cold 1.5 M triethylammonium bicarbonate (TEAB at pH 7.5), and lyophilized. The residue was dissolved in 10 mL of water and purified by ion-exchange chromatography [DEAE Sephadex A25, 9 × 2.5 cm, linear gradient of TEAB (0.05 to 1.5 M at pH 7.5, 1500 mL) at a flow rate of 5 mL/min]. The 

triphosphate was eluted at 0.6-0.7 M TEAB. Lyophilization (three times from water) gave da *TP (40.6 mg, 55%) as a yellow solid. Analytical HPLC: t₁ = 18.27 min. UV [H₂O]: 364, 276. 31P-

NMR [162 MHz, D₂O]: -7.19 [d, J = 20.1, P(II)]; -10.62 [d, J = 20.9, P(II)]; -22.07 [t, J = 19.5, P(II)]. MS-ES [negative mode]: 

C₁₇H₂₃N₅O₈P₃ (535): 534 (100, [M – H]).

Footprinting with DMS
Promoter fragments used in DMS protection footprints were obtained by HindIII digestion of pRLG213 or pRLG3289 and 

labeling of the 3' end of the nontemplate strand using Sequenase 

[USB] and [α-32P]dATP [Newlands et al. 1991]. Then, labeled 

DNA was digested with AatII [vector-encoded site is 75 bp up-

stream of EcoRII], and the fragment was gel isolated, eluted, and 
purified by use of Elutip D columns as described above.

Promoter fragments (~1 nM) were pre-incubated with wild-
type or α mutant RNAP (~10 nM) or with purified α subunit (~2 

µM) for 10 min at 22°C in 50 µL of the buffer used above for interference footprinting. Then, the DNA was methylated by addition of 1 µL of DMS for 30 sec at 22°C, followed by addition of 25 µL of cold 1.5 M β-mercaptoethanol and 0.3 M NaOAc, extraction with 75 µL of cold phenol, precipitation with etha-

nol, and reprecipitation from 0.3 M NaOAc. The DNA pellet was 

resuspended in 30 µL of 10 mM sodium phosphate [pH 6.8], 1 

mM EDTA, and methylated purines [A,G] were released by 

incubation at 92°C for 15 min [Maxam and Gilbert 1980]. Strand 

cleavage at abasic sites was carried out by addition of 3 µL of 1 

N NaOH and further incubation at 92°C for 30 min. Samples 

were precipitated with ethanol, and examined by electrophore-

sis, gel scanning, and quantitation as described above.

Hydroxyl radical footprinting to detect distamycin-binding sites
An end-labeled promoter fragment containing rrnB P1 with UP 
element 4547 [prepared as described for DMS footprinting 

above) was incubated at 22°C for 10 min with 125, 250, or 500 

nM distamycin A [Sigma] in 150 mM NaCl, 20 mM Tris-Cl (pH 7.4), 1 mM EDTA, and 1 µg/mL sonicated calf thymus DNA. 

OH cleavage was carried out as described previously [Newlands
et al. 1991), and gel electrophoresis and scanning of gels was performed as described above.

**Gel mobility shift assay with the α subunit**

An end-labeled promoter fragment (∼0.1 nM; rrnB P1 with proximal subsite 4547) was preincubated for 10 min at 22°C with 15.6–1000 nM distamycin A in 10 µL of buffer containing 150 mM NaCl, 20 mM Tris–Cl (pH 7.4), 1 mM EDTA, 10% glycerol, and 1 µg/mL sonicated calf thymus DNA. Purified α subunit (2 µM) was then added for 10 min, and electrophoresis was carried out at 4°C on prechilled 6% acrylamide gels containing 0.5x TBE and 10% glycerol. Bands containing the α–DNA complex were quantified by PhosphorImaging, and values were expressed as a percentage of the fragment population bound in the absence of distamycin.

**In vitro transcription**

Plasmid DNAs (∼60 ng of pRLG4213 or pRLG4210) were incubated with distamycin A (15.6–1000 nM) for 10 min at 22°C in 25 µL of transcription buffer containing 150 mM NaCl, 10 mM Tris–Cl (pH 7.9), 10 mM MgCl₂, 1 mM DTT, 100 µg/mL bovine serum albumin, 500 µM ATP, 50 µM CTP, 50 µM GTP, 10 µM UTP, and 4 µCi [³²P]UTP [NEN]. Transcription was initiated by addition of RNAP (2 µM) for 15 min, and the reaction was terminated by addition of 25 µL of stop solution [Ross et al. 1998]. Samples were analyzed by electrophoresis on 5% acrylamide–7 M urea gels and quantified after gel scanning with a PhosphorImager.

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Fine structure of *E. coli* RNA polymerase-promoter interactions: α subunit binding to the UP element minor groove

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