

See discussions, stats, and author profiles for this publication at: <https://www.researchgate.net/publication/41759113>

Probing DNA Binding, DNA Opening, and Assembly of a Downstream Clamp/Jaw in Escherichia coli RNA Polymerase– λ P R Promoter Complexes Using Salt and the Physiological Anion Glutamate

ARTICLE *in* BIOCHEMISTRY · MARCH 2010

Impact Factor: 3.02 · DOI: 10.1021/bi100092a · Source: PubMed

CITATIONS

24

READS

33

5 AUTHORS, INCLUDING:



Theodore J Gries

Beloit College

7 PUBLICATIONS 92 CITATIONS

SEE PROFILE



Ruth Saecker

University of Wisconsin–Madison

41 PUBLICATIONS 4,265 CITATIONS

SEE PROFILE

Published in final edited form as:

Biochemistry. 2010 May 25; 49(20): 4361–4373. doi:10.1021/bi100092a.

Probing DNA Binding, DNA Opening and Assembly of a Downstream Clamp/Jaw in E. coli RNA Polymerase – λP_R Promoter Complexes Using Salt and the Physiological Anion Glutamate[†]

Wayne S. Kontur^{‡,¶,1}, Michael W. Capp^{§,¶}, Theodore J. Gries[§], Ruth M. Saecker^{§,*}, and M. Thomas Record Jr.^{‡,§,*}

[‡]Department of Chemistry University of Wisconsin-Madison, Madison WI 53706

[§]Department of Biochemistry University of Wisconsin-Madison, Madison WI 53706

Abstract

Transcription by all RNA polymerases (RNAP) requires a series of large-scale conformational changes to form the transcriptionally-competent open complex RP_o . At the λP_R promoter, E. coli σ^{70} RNAP first forms a wrapped, closed 100 bp complex I_1 . The subsequent step opens the entire 13 base DNA bubble, creating the relatively unstable (open) complex I_2 . Additional conformational changes convert I_2 to the stable RP_o . Here we probe these events by dissecting the effects of Na^+ salts of Glu^- , F^- and Cl^- on each step in this critical process. Rapid mixing and nitrocellulose filter binding reveal that the binding constant for I_1 at 25 °C is ~30-fold larger in Glu^- than in Cl^- at the same $[Na^+]$, with the same log-log [salt] dependence for both anions. In contrast, both the rate constant and equilibrium constant for DNA opening (I_1 to I_2) are only weakly [salt]-dependent, and the opening rate constant is insensitive to replacement of Cl^- by Glu^- . These very small effects of [salt] on a process (DNA opening) which is strongly [salt]-dependent in solution may indicate that the backbones of both DNA strands interact with polymerase throughout the process, and/or that compensation is present between ion uptake and release.

Replacement of Cl^- by Glu^- or F^- at 25 °C greatly increases the lifetime of RP_o and greatly reduces its [salt]-dependence. By analogy to Hofmeister salt effects on protein folding, we propose that the excluded anions Glu^- and F^- drive the folding and assembly of the RNAP clamp/jaw domains in the conversion of I_2 to RP_o , while Cl^- does not. Because the Hofmeister effect of these anions largely compensates for the destabilizing coulombic effect of any salt on the binding of this assembly to downstream promoter DNA, RP_o remains long-lived even at 0.5 M Na^+ in Glu^- or F^- salts. The observation that $E\sigma^{70} RP_o$ are exceedingly long-lived in moderate to high $[Glu^-]$ argues that $E\sigma^{70}$ RNAP do not dissociate from strong promoters in vivo when the cytoplasmic increases during osmotic stress.

[†]This work was supported by NIH grant GM23467 to M.T. R. W. S. K. gratefully acknowledges the support of the Biotechnology Training Program (NIH 5 T32 GM08349); T. G. is the recipient of the William R. and Dorothy E. Sullivan Distinguished Graduate Fellowship.

*To whom correspondence should be addressed: Telephone 608-262-5332, FAX 608-262-3453 mtrecord@wisc.edu, rmsaecker@wisc.edu.

[¶]These authors contributed equally to this work.

¹Current address: Department of Bacteriology, University of Wisconsin-Madison/Great Lakes Bioenergy Research Center, Madison, WI 53706

Supporting Information

Supporting information reports the irreversible rate constants (k_{obs}) for formation of open complexes as a function of $[RNAP]_{total}$ and $[Na^+]$ in Cl^- buffer (STable 1), and in Glu^- and F^- buffers (STable 2). STable 3 presents the fits ($K_{heparin}$, $[R]_{free}$) of relaxation to equilibrium data (k_r) at 25 °C to obtain k_d in Glu^- and F^- buffers. This material is available free of charge via the Internet at <http://pubs.acs.org>.

E. coli cells have the remarkable ability to grow over a very wide range of environmental conditions, including osmotic conditions ranging from very dilute (0.02 osmolal) to concentrated, high [salt] media (> 3 osmolal) (3-5). In vivo, increases in external osmolarity, which increase cytoplasmic $[K^+]$ and $[Glu^-]$, reduce bulk transcription (6) while transcription from genes involved in the osmotic stress response selectively increases (7-9). In the absence of any known osmosensing (e.g. glutamate binding) transcription factor, the simplest hypothesis is that changes in cytoplasmic $[K^+]$ and $[Glu^-]$ alone alter transcription (10) during the initial response to osmotic upshift. What features of RNA polymerase and promoter sequence allow the cell to achieve a dramatic reprogramming to maintain homeostasis?

Understanding how cells regulate their gene expression requires kinetic-mechanistic studies. Although transcription by *E. coli* RNAP has been studied in vivo and in vitro for four decades, we are only beginning to understand the workings of this machine at a molecular level. Key questions are how is DNA opened by RNAP and what significant conformational changes occur during the steps that precede and follow this central DNA opening step? How do variables like the identity of the sigma specificity subunit, the promoter DNA sequence, supercoiling, concentrations of solutes, small ligands, or regulatory proteins affect the rate of formation and lifetime of the transcriptionally-competent open complex, and thereby alter the frequency of productive versus abortive initiation from the promoter?

Because the RNAP machinery and promoter organization are conserved, it likely that the sequence of conformational changes that form the bacterial RP_o is largely invariant (promoter DNA recognition, DNA opening, stabilization of RP_o), but that the rates of interconversion and populations of initiation intermediates differ as a function of the variables noted above. Here we investigate how the substitution of the physiological anion Glu^- for the laboratory anion Cl^- affects each step in the process of open complex formation at the λP_R promoter by the housekeeping form of *E. coli* RNA polymerase (RNAP, subunit composition $\sigma^{70}\alpha_2\beta'\omega$). While our research focuses on this promoter, we expect that these results reflect the salient functional characteristics of many $E\sigma^{70}$ RNAP promoter interactions.

In vitro, changes in salt concentration profoundly affect both the composite second order association rate constant (k_a) and composite dissociation rate constant (k_d) for the formation and dissociation of RP_o at the σP_R promoter, respectively (11,12). One contributor to these salt effects is coulombic: positively charged residues in the promoter-RNAP interface reduce the high negative phosphate charge density of DNA, releasing locally accumulated salt ions (e.g. K^+ or Na^+ from DNA). In addition, the identity of the anion has large effects on the magnitude of k_a : for the σP_R promoter k_a was found to be at least 30-fold larger in glutamate (Glu^-) than in chloride (Cl^-) at 0.20 M K^+ (13). Here we dissect this large salt anion effect. Large effects of substituting another anion for Cl^- are also observed for many other nucleic acid DNA binding proteins, including transcription factors (14-16), helicases (15), T7 RNAP (17) and DNA polymerases. Large Hofmeister effects of salt anions like Glu^- and F^- , relative to Cl^- , are expected on steps in which interface formation or coupled folding buries large amounts of nonpolar (hydrocarbon) surface (18,19).

At the strong λP_R promoter (P), the mechanism of open complex formation consists of a minimum of three steps, proceeding through two kinetically significant intermediates (11, 20-22)



where R represents RNA polymerase $E\sigma^{70}$, P is promoter, and I_1 and I_2 are the first and second kinetically significant intermediates, respectively. Opening of the entire DNA bubble (−11 to +2) occurs in the conversion of I_1 to I_2 ((23); Gries et al., unpublished). No other early intermediates are kinetically significant at the λP_R promoter. Solute upshift studies designed to study the late steps in RP_o formation reveal the existence of an intermediate (I_3) forming after I_2 and directly preceding the formation of RP_o (24). I_3 rapidly equilibrates with I_2 and RP_o under the conditions of this study, and thus can be neglected in the analysis of k_d presented here. Because the kinetics of open complex formation (association) and dissociation are single exponential for almost all conditions examined, we deduce that: i. in association I_1 rapidly equilibrates with free RNAP and promoter DNA on the time scale required for I_1 to convert to I_2 in the forward direction ($k_{-1} \gg k_2$); and ii. I_2 rapidly equilibrates with RP_o on the time scale required for I_2 to convert to I_1 in the reverse direction ($k_3 \gg k_{-2}$). The interconversions of I_1 and I_2 are the bottleneck steps in both the association and dissociation directions.

Using rapid mixing we recently dissected the composite association and dissociation rate constants for RNAP- λP_R interactions into their individual contributions from $K_1 (=k_1/k_{-1})$, k_2 , k_{-2} and $K_3 (=k_3/k_{-3})$, respectively (22,24). We discovered that destabilizing RP_o by a rapid upshift in salt or urea concentration creates a transient burst of I_2 (24). Coupling rapid mixing with filter binding, we determined the kinetics of forming this unstable open complex (I_2) from the stable open complex RP_o , and converting I_2 to the closed complex I_1 over a wide range of salt concentrations at 10 °C and 37 °C. These studies dissected the effect of [salt] on the dissociation kinetics for the first time. The rate constant k_{-2} is independent of salt concentration, while the equilibrium constant for the conversion of I_2 to RP_o decreases strongly with increasing [salt] (24). Permanganate footprinting of I_2 reveals that all reactive thymines in RP_o are also reactive in I_2 , demonstrating that the entire transcription bubble is opened in the rate-determining, single step $I_1 \rightarrow I_2$ (Gries et al., unpublished). Measurements of the time dependence of the decay of permanganate reactivity in conversion of I_2 to I_1 corroborate the rate constant k_{-2} obtained by rapid mixing and filter binding.

The unexpected result that the rate constant k_{-2} characterizing the conversion of I_2 to the transition state (I_1 - I_2) ‡ is independent of salt concentration motivates the work here to dissect the effects of salt concentration on the steps on the first half of the mechanism, namely the formation of I_1 and the transition between I_1 and (I_1 - I_2) ‡ . Melting of DNA in solution is strongly salt concentration dependent (25). How does melting DNA to form the initiation bubble in the isomerization of I_1 to I_2 compare?

To address this question we examine whether the rate constant k_2 characterizing the conversion of I_1 to (I_1 - I_2) ‡ is [salt]-dependent. Using rapid mixing and filter binding, we dissect for the first time the overall second order association rate constant k_a into K_1 and k_2 over a range of NaCl and NaGlu concentrations. In addition, we obtain stringent tests of the single exponential character of both the association kinetic data (in [RNAP] excess) and dissociation kinetic data. These findings demonstrate that the initial steps of association (forming I_1) and of dissociation (forming I_2) do indeed rapidly equilibrate on the time scale of the subsequent slower conformational change (DNA opening or closing). We use manual mixing filter binding to examine the effects of salt concentration (NaX) and choice of anion ($X = Cl^-$, Glu^- , or F^-) on the dissociation of RP_o at 25 °C.

We find that each step exhibits a uniquely different dependence on changes in $[Na^+]$ and choice of anion. While the formation of I_1 and conversion of I_2 to RP_o exhibit large dependences on both $[Na^+]$ and anion, the bottleneck DNA opening step ($I_1 \rightarrow I_2$) is relatively insensitive to either solution variable. Formation of I_1 and especially RP_o are greatly favored in Glu^- relative to Cl^- . Together these results shed additional light on the conformational changes occurring in each step, allowing us to refine our current structural-mechanistic hypotheses regarding open

complex formation by *E. coli* RNAP. In particular, we propose that the very large effects of replacing Cl^- with Glu^- on the steps *after* DNA opening result from burial of hydrocarbon surface of RNAP in coupled folding transitions in late steps (24,26,27). Glu^- and F^- are more highly excluded from nonpolar surface than is Cl^- (18,19). We deduce that these late remodeling events assemble a RNAP clamp/jaw on downstream DNA and contribute significantly to the entropic driving force that stabilizes the open DNA state in RP_0 .

MATERIALS AND METHODS

Buffers

Storage Buffer (SB) for RNA polymerase holoenzyme contained 50% glycerol (v/v), 10 mM Tris-HCl (pH 7.5 at 4 °C), 100 mM NaCl, 0.1 mM dithiothreitol (DTT), and 0.1 mM Na_2EDTA . Binding Buffer for the study of the kinetics of RNAP-promoter binding as a function of univalent salt concentration in the absence of Mg^{2+} (BB) contained 10 mM Na_2HPO_4 buffer (pH 7.4), 6.5% (v/v) glycerol, 1 mM DTT, 100 $\mu\text{g}/\text{mL}$ BSA, 1.3 mM Tris (pH 7.5 at 4 °C; contributed by the Storage Buffer), and variable $[\text{NaCl}]$, $[\text{NaGlu}]$ or $[\text{NaF}]$ to bring $[\text{Na}^+]$ to the desired value. Wash Buffer (WB) contained 0.1 M NaCl, 10 mM Tris-HCl (pH 8.0 at room temperature), and 0.1 mM Na_2EDTA . Addition of NaGlu did not significantly affect the pH of the Tris-HCl buffer.

Wild-type $E\sigma^{70}$ RNA polymerase holoenzyme

E. coli K12 wild type RNA polymerase holoenzyme was purified as described in (28) and stored in SB at -70°C . All [RNAP] reported here are concentrations of active holoenzyme (capable of RP_0 formation). Activities were determined at the time of use, either by analysis of forward titrations of promoter DNA with RNAP performed in the 'complete binding of limiting reagent' regime at 37°C as described (11) or by comparison of association rate constant for different samples determined at the same total [RNAP]. Typically activities were ~70%. Systematic comparisons of the different preparations showed that no difference in promoter association kinetics between individual tubes or preps, when compared at the same active [RNAP].

λP_R promoter DNA

A ^{32}P -DNA fragment containing the λP_R promoter was obtained from the plasmid pBR81 as described in (29). *Bss*H II and *Sma* I cleavage of pBR81 centrally positions the λP_R wild-type sequence (−60 to +20) in a fragment which extends from −115 to +76 relative to the transcription start site (+1) of the promoter. The specific activity of the fragment was generally $\sim 10^8$ cpm/mole.

Association Kinetics

For conditions where the kinetics are sufficiently slow (e.g. RNAP concentrations below 10 nM at mid-range salt concentrations; high ($<0.3\text{M}$) salt concentrations), manual mixing and filter binding were used to determine the kinetics of formation of long-lived (open) complexes, resistant to a 10 - 30 s heparin challenge. For conditions where the kinetics are too fast (first order rate constant $k_{\text{obs}} > 0.01\text{ s}^{-1}$) to determine by manual mixing (e.g. low salt and higher RNAP concentrations), rapid mixing (2 ms mixing time; RQF-3, Kintek Corp., Austin TX) followed by filter binding on quenched solutions was used (22). Rapid mixing and manual mixing experiments were performed as described in (22). Concentrations of active RNAP were always in at least 6-fold excess over that of promoter to ensure that the initial bimolecular binding step is pseudo-first order within experimental uncertainty.

Dissociation Kinetics

i. Irreversible dissociation initiated by adding heparin to pre-formed RP_o —

Effects of $[Na^+]$ in Cl^- buffer on the dissociation of preformed λP_R DNA (0.08-0.09 nM) and RNAP (10 – 40 nM) open complexes at 25 °C were performed as described in (27).

ii. Decay to equilibrium assay to determine k_d —The irreversible dissociation of RP_o complexes initiated by the addition of excess heparin is exceedingly slow in Glu^- at 25 °C, yielding poorly determined values of k_d . Thus to determine k_d as a function of Na^+ in Glu^- buffer, we used a decay to equilibrium assay at a low $[RNAP]$ held constant by an excess of the competitor heparin. In this assay, 7 nM RNAP was initially mixed with a vast excess of heparin (0.1 – 1 mg/mL) at the salt concentration of the experiment. After 60 min of incubation, ^{32}P λP_R DNA (12.5 pM) was added, initiating promoter binding which proceeds to equilibrium at the low $[RNAP]$ of the experiment. The amount of λP_R - RNAP open complexes was monitored using nitrocellulose filter binding. Typically $>10^5$ s was required to approach promoter binding equilibrium. At longer times, a very slow decay of the plateau was noted, probably the result of inactivation of RNAP.

Nitrocellulose filter binding assays

The amount of open complexes as a function of time and solution condition was determined using nitrocellulose filter binding as described in (22). Studies in Glu^- and F^- cannot be performed above 0.70 M Na^+ because filter efficiency becomes too low.

DATA ANALYSIS

Data were analyzed using either SigmaPlot 6.0 or 2000 (SPSS, Inc. Chicago, IL) on a Dell Optiplex GX270 workstation running Windows XP Professional v. 2002 (Microsoft, Seattle, WA), or using Igor 5.0.3.0 (WaveMetrics, Inc., Lake Oswego, OR)) on Dell Inspiron 8100.

Determination of k_{obs} , k_a , K_1 and k_2 from the kinetics of formation of open complexes

In excess RNAP, the kinetics of irreversible association of RNAP with λP_R promoter DNA to form open complexes (I_2 , RP_o) are single exponential (see Results) with observed rate constant k_{obs} , determined by fitting the fraction of promoter DNA bound in open complexes at time t (θ_t^{open}) (calculated as described in (22)) versus time to eq 1:

$$\theta_t^{open} = 1 - e^{-k_{obs}t} \quad (1)$$

The rate constant k_{obs} (previously called α_{CR}) is a hyperbolic function of RNAP concentration (22):

$$k_{obs} = \frac{K_1 k_2 [RNAP]}{1 + K_1 [RNAP]} = \frac{k_a [RNAP]}{1 + K_1 [RNAP]} \quad (2)$$

where $K_1 (= k_1/k_{-1})$ is the equilibrium constant for formation of I_1 , k_2 is the microscopic rate constant for the subsequent (rate-determining) conversion of I_1 to I_2 in Mechanism I, and $k_a = K_1 k_2$ is the composite overall second-order association rate constant. The data for k_{obs} as a function of $[RNAP]$ in Figs 3 and 4 were weighted by $1/\sigma^2$, where σ is the standard deviation of k_{obs} , and fit to eq 2 to determine values of k_a , K_1 and k_2 at each salt concentration investigated. $K_1 (= k_1/k_{-1})$ is the equilibrium constant for formation of the first kinetically-significant intermediate (I_1 , in rapid equilibrium with free RNAP and DNA), $k_2 (= k_i)$ is the microscopic rate constant for the subsequent (rate-determining; $k_2 < k_{-1}$) conversion of I_1 to

I_2 in Mechanism I, and $k_a (= K_1 k_2)$ is the composite overall second-order association rate constant. Values of k_{obs} (weighted by $1/\sigma^2$, where σ is the standard deviation of k_{obs}) were plotted versus [RNAP] and fit to eq 2 to determine k_a , K_1 and k_2 .

Above 0.23 M Na^+ in NaCl buffer and 0.29 M Na^+ in NaGlu buffer k_{obs} increases linearly with [RNAP] up to the highest accessible concentration (~120 nM), indicating (eq 2) that $K_1[RNAP] \ll 1$. In this limit, the overall association rate constant was determined using the following equation:

$$k_a = k_{obs} / [RNAP] (1 + K_1 [RNAP]) \approx k_{obs} / [RNAP] \quad (3)$$

The quality of the approximation was estimated by linear extrapolation of the log-log plots of K_1 vs. [RNAP] (Fig 4A below) and found to be appropriate.

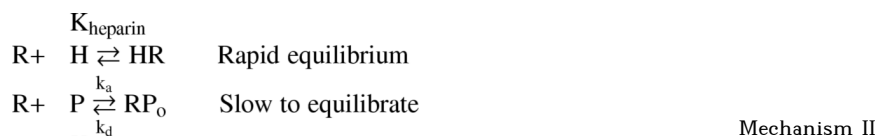
Determination of the Composite Dissociation Rate Constant k_d

- i. The observed rate constant k_d for the irreversible dissociation of open complexes was determined by fitting θ_t^{open} versus time at a given $[Na^+]$ to a single-exponential decay equation

$$\theta_t^{open} = \theta_{t=0}^{open} e^{-k_d t} \quad (4)$$

where $\theta_{t=0}^{open}$ is the value of θ_t^{open} at time $t = 0$.

- ii. The promoter dissociation rate constant (k_d) and the RNAP-heparin binding constant ($K_{heparin}$) at 25 °C in Glu^- and F^- buffers were determined from the observed decay to equilibrium (relaxation) rate constant k_r (see below) and the RNAP-promoter association rate constant k_a (Table1). The data were modeled using the following mechanism:



In mechanism II, H is heparin, P is promoter DNA, R is free RNAP and RP_o is the final open complex (concentrations of intermediates are negligible). $K_{heparin}$ is the equilibrium constant for the formation of a heparin-RNAP complex; this equilibrates rapidly on the time scale of association of RNAP with promoter DNA. The composite rate constants for formation (k_a) and dissociation (k_d) of RP_o are defined by eqs 2 and 4, respectively. Our analysis of Mechanism II assumes one heparin binding site on RNAP and applies to conditions where $[H]_{total} \gg [R]_{total} \gg [P]_{total}$. For this situation, the experimentally determined approach-to-equilibrium rate constant

$k_r = (d \ln (\theta_{t=\infty}^{open} - \theta_t^{open}) / dt) = k_a [R] + k_d$, where $[R]$ is the free RNAP concentration. $[R]$ is constant in these assays (though not in excess over promoter) because the heparin-binding equilibrium buffers its free concentration. The integrated form of this equation is used to determine k_d :

$$\theta_t^{open} = \frac{k_a [R] (1 - e^{-(k_a [R] + k_d)t})}{k_a [R] + k_d} \quad (5)$$

where $[R]$ is determined from eq 6:

$$[R] = \frac{[R]_{total}}{(1 + K_{heparin} [H]_{total})} \quad (6)$$

For the set of relaxation data collected as a function of $[H]_{total}$ at a given Na^+ in Glu^- or F^- buffer, data at each [heparin] were fit to eqs 5-6, using the value of k_a determined from the independent kinetics of association experiments (Table 1). Reported values of $K_{heparin}$ and k_d for each salt condition represent the average and standard deviation from these individual fits.

Results

Single Exponential Kinetics of Formation of E. coli RNA Polymerase – λP_R Promoter Complexes: Effects of Glutamate and Chloride Salts

To compare the effects of the laboratory anion chloride (Cl^-) and the physiological anion glutamate (Glu^-) on the binding of E. coli RNAP to the λP_R promoter to form a closed complex and on the subsequent DNA opening step, we determined the kinetics of association over wide ranges of RNAP concentration (1 nM – 100 nM) and concentrations of chloride and glutamate salts (0.13 M to 0.27 M Na^+ in Cl^- buffer; 0.23 M to 0.64 M Na^+ in Glu^- buffer) at 25 °C. A limited number of association experiments were performed in F^- buffer at low [RNAP] to compare to the results in Glu^- buffer. Sodium salts were used because complexation of potassium by heparin interferes with the heparin challenge (see Methods) at high concentrations of potassium salts (12); Mg^{2+} was omitted because it is complexed by glutamate (30). In most cases the full accessible range of RNAP and of salt concentration were investigated.

Following either manual or rapid mixing, nitrocellulose filter binding was used to determine the kinetics of formation of long-lived, competitor-resistant complexes (RP_o). Representative kinetic data for selected low and high RNAP concentrations over the range of NaCl and NaGlu concentrations are plotted as the fraction of promoter DNA in open complexes (θ_t^{open}) as a function of time in Figures 1 and 2, respectively. The plateau at $\theta_t^{open} = 1$ observed at long times for all salt concentrations investigated demonstrates that open complex formation is irreversible and therefore directly interpretable in terms of the forward rate constant k_{obs} (see eq 1 in Methods), without the need for a decay-to-equilibrium analysis. Irreversibility of the forward kinetics is confirmed by direct measurement of the dissociation rate constant (see below).

Single exponential fits of these data to the irreversible first order rate eq 1 (Methods) are shown in Figs 1 and 2. Values of k_{obs} determined from these fits are given in Supplemental Tables 1 and 2, respectively. To expand the initial (ms) time regime and examine whether the short time data provide any indication of a second (faster) decay, the insets to the panels of Figs 1 and 2 plot the same data as θ_t^{open} vs. $\log t$ (cf. (1,2)). For the λP_R promoter and the conditions investigated previously (cf. (11,12,20,22)) and in the present study, all association kinetic data are well-described as single exponential decays. Single exponential kinetics of open complex

formation indicate that the step forming the first kinetically significant intermediate (I_1) rapidly equilibrates with free promoter DNA on the time scale of the conversion of I_1 to I_2 (21).

The Observed Forward Rate Constant k_{obs} is a Hyperbolic Function of [RNAP] over the Accessible Range (1 nM to 100 nM)

Values of the overall rate constant k_{obs} are plotted versus RNAP concentration in Fig 3A (Cl^-) and B (Glu^-) for the salt concentrations investigated. In most cases these data cover a 100-fold range of RNAP concentrations (~1 nM to 100 nM). At all salt concentrations investigated, except the highest [Glu^-], values of k_{obs} exhibit the expected hyperbolic dependence on [RNAP] (eq 2). Except at the highest salt concentrations, a well-defined plateau is obtained in the accessible range of [RNAP]. There is no evidence of sigmoidicity in any these plots at low [RNAP], as would be predicted if dissociation of holoenzyme into sigma subunit and core polymerase were significant at these concentrations of RNAP and salt (31). The observations of single-exponential association kinetics (Figs 1 & 2) and hyperbolic dependences of k_{obs} on [RNAP] (Fig 3) validate the use of eq 2 (Methods) to analyze the dependence of k_{obs} on [RNAP] and obtain k_2 , the DNA opening rate constant and K_1 , the equilibrium constant for forming I_1 (see Methods).

DNA Opening by RNAP is Relatively Slow and Only Weakly [Salt]-Dependent; Replacing Cl^- by Glu^- Has No Significant Effect on k_2 or on its log-log derivative Sk_2

Rate constants k_2 of the DNA opening step are plotted on a log-log scale as a function of [Na^+] for experiments in Cl^- and Glu^- buffer in Fig 4. Values of k_2 decrease very modestly with increasing salt concentration, from 0.35 s^{-1} at 0.15 M Na^+ in Cl^- buffer to 0.18 at 0.29 M Na^+ in Glu^- buffer, corresponding to time constants of 3-5 s for opening. Neither k_2 at 0.23 M salt (the overlap concentration in our experiments) nor Sk_2 is affected by replacement of Cl^- by Glu^- (Fig 4B); individual fits of these data yield Sk_2 values of -1.1 ± 0.1 and -0.9 ± 0.3 in Cl^- and Glu^- buffers, respectively. As shown in Fig 4A, the combined data fall on a common line with a log-log slope of $\text{Sk}_2 = -1.1 \pm 0.1$. Previously we found from [KCl]-upshift experiments at 10 and 37 °C in transcription buffer (10 mM MgCl_2) that the rate constant k_{-2} for the reverse direction of this step (I_2 to I_1) is independent of [KCl] ($\text{Sk}_{-2} = 0$; (24)). Interpolation between values yields $k_{-2} \approx 1.7 \text{ s}^{-1}$ at 25 °C. Assuming this value of k_{-2} applies to the [salt] conditions investigated here, we obtain equilibrium constants K_2 for DNA opening by RNAP (i.e. $I_1 \rightleftharpoons I_2$) that decrease from 0.2 at 0.15 M salt to 0.1 at 0.29 M salt. I_2 is therefore unstable with respect to I_1 at all salt concentrations, and the [salt]-dependence of the DNA opening step $I_1 \rightleftharpoons I_2$ on the protein is small ($\text{SK}_2 = \text{Sk}_2 - \text{Sk}_{-2} = -1.1 \pm 0.1$).

Formation of Wrapped I_1 Is Strongly [Salt]-Dependent ($\text{SK}_1 = -6.9 \pm 0.3$); Replacing Cl^- by Glu^- Increases K_1 ~30-fold Without Significantly Affecting SK_1

The proposed RNAP-DNA interface in I_1 spans 100 bp; the net positive charge on RNAP in this interface is at least +37 (see Discussion). How [salt]-dependent is this binding interaction? In Cl^- buffer the binding constant K_1 for formation of I_1 decreases from $1.8 \times 10^8 \text{ M}^{-1}$ at 0.15 M Na^+ to $9.9 \times 10^6 \text{ M}^{-1}$ at 0.23 M Na^+ . In Glu^- buffer at 0.23 M Na^+ , K_1 is $3.6 \times 10^8 \text{ M}^{-1}$ and decreases to $8 \times 10^7 \text{ M}^{-1}$ at 0.29 M Na^+ . In both Cl^- and Glu^- , $\log K_1$ decreases linearly with $\log[\text{Na}^+]$ with slopes $\text{SK}_1 = \text{dlog}K_1/\text{dlog}[\text{salt}]$ which are the same within uncertainty ($\text{SK}_1 = -6.8 \pm 0.3$ in Cl^- , $\text{SK}_1 = -7.5 \pm 2.3$ in Glu^- ; see Fig 4A and Table 1). Fitting both data sets together yields $\text{SK}_1 = -6.9 \pm 0.3$. These are large [salt]-dependences, but not nearly as large as expected from consideration of the interface (see Discussion).

At 0.23 M Na^+ , the only salt concentration where the data sets overlap, the value of K_1 in Glu^- buffer exceeds that in Cl^- by a factor of ~30. This is the same factor by which the log-log plots of k_a vs. [Na^+] in Fig 4B are displaced. Such a large effect of replacing Cl^- with

Glu^- on K_1 is likely a Hofmeister salt effect, based on the observation that fluoride and glutamate have very similar effects on $k_a = K_1 k_2$, as well as on k_d (see below).

Replacement of Cl^- by Glu^- Increases the Second-Order Association Rate Constant k_a by ~30-fold at All [Salt]

Above 0.23 M Na^+ in Cl^- buffer and 0.29 M Na^+ in Glu^- buffer, the equilibrium constant K_1 for I_1 formation becomes too small to allow us to dissect k_{obs} into K_1 and k_2 . At higher [salt], we find that k_{obs} increases linearly with [RNAP], as shown in Fig 3B in Glu^- buffer at 0.55 M Na^+ . Linearity indicates that $K_1[\text{RNAP}] \ll 1$ over the accessible range of [RNAP]. In this limit, the second order association rate constant $k_a = k_{\text{obs}}/[\text{RNAP}](1 + K_1[\text{RNAP}]) \approx k_{\text{obs}}/[\text{RNAP}]$, where the quality of the last approximation can be estimated by extrapolation of the log-log plots of K_1 vs. [RNAP] (Fig 4A). Although individual values of K_1 and k_2 cannot be determined in this higher [salt] range, measurements of k_{obs} at low [RNAP] (typically 7 nM) were performed in Cl^- , Glu^- and F^- buffers up to the highest $[\text{Na}^+]$ accessible in each to test for curvature of the log-log plots arising from Hofmeister effects (32). All determinations of k_a are listed in Table 1, and plotted in Fig 4B as a log-log plot together with those obtained from the data of Fig 3. A much wider range of high $[\text{Na}^+]$ can be investigated in Glu^- and F^- buffers because k_d is much smaller in Glu^- and F^- buffers than in Cl^- . Over the full range of $[\text{Glu}^- \text{ (or } \text{F}^-)]$ investigated, k_a decreases by more than three orders of magnitude (from $7.3 \times 10^7 \text{ M}^{-1} \text{ s}^{-1}$ at 0.23 M Na^+ to $2.6 \times 10^4 \text{ M}^{-1} \text{ s}^{-1}$ at 0.64 M Na^+). These log-log plots are linear and parallel within the uncertainty over the entire accessible salt range in both Cl^- ($\text{Sk}_a = -8.0 \pm 0.2$) and Glu^-/F^- ($\text{Sk}_a = -8.3 \pm 0.2$). The offset corresponds to ~30-fold faster association in Glu^- than in Cl^- . This difference is entirely the result of the difference in K_1 .

Disassembly of the RNAP Clamp/Jaw in Dissociation of RP_0 Complexes in Glu^- is Remarkably Slower and Much Less [Salt] Dependent than in Cl^-

Dissociation kinetic studies probe the steps of disassembly of the clamp/jaw (conversion of RP_0 to I_2) and DNA closing (conversion of I_2 to $(\text{I}_1\text{-I}_2)^+$). Irreversible dissociation data at 25 °C in 0.23 M – 0.29 M Na^+ in Cl^- buffer are plotted in Fig 5 as θ_t^{open} vs. time, and fit to a single exponential decay equation (eq 4). To show the quality of these single exponential fits, the inset in Fig 5 plots θ_t^{open} vs $\log t$; essentially 100% of the decay is described by the fit with dissociation rate constant k_d . This indicates that the steps that convert the final open complex RP_0 to the kinetically significant, open intermediate I_2 rapidly equilibrate on the time scale of the dissociation-rate-determining conversion of I_2 to the closed intermediate I_1 . (I_1 is not filter-retainable in the presence of heparin.) Values of k_d increase by an order of magnitude with this 25% increase in salt concentration (from $2.9 \times 10^{-4} \text{ s}^{-1}$ at 0.23 M Na^+ to $2.4 \times 10^{-3} \text{ s}^{-1}$ at 0.29 M Na^+); these are plotted on a log-log scale vs $[\text{Na}^+]$ in Fig 7 below. The plot is linear with slope $\text{Sk}_d = 9.2 \pm 0.1$, consistent with a previous determination for this promoter in its natural context (12).

By contrast, dissociation in Glu^- is extraordinarily slow at 25 °C and much less dependent on salt concentration. Attempts to directly determine k_d at 0.33 M and 0.6 M Na^+ in Glu^- buffer, using a very high competitor concentration (2.5 mg/mL heparin) in an attempt to ensure irreversibility, exhibited very slow dissociation with rate constants in the range 10^{-5} to 10^{-6} s^{-1} , orders of magnitude slower than in Cl^- at lower [salt] (Fig 7). Permanganate footprinting of these 25 °C promoter complexes as a function of $[\text{Na}^+]$ in Glu^- buffer shows them to be open in the vicinity of the transcription start site over the range 0.13 M to 0.6 M Na^+ (TG, unpublished).

As an alternative, faster and more robust method to determine k_d at 25 °C as a function of $[\text{Na}^+]$ in Glu^- buffer, we observed the kinetics of approach to promoter binding equilibrium in experiments performed at very low, constant [RNAP], maintained by using heparin as an

RNAP buffer. In these experiments, RNAP was initially equilibrated with excess heparin, and subsequently mixed with promoter DNA to initiate open complex formation, which is reversible in excess heparin because the free [RNAP] is so low.

Representative results at 0.46 M Na^+ in Glu^- buffer are shown in Fig 6 as plots of θ_t^{open} as a function of time for different heparin concentrations. In these experiments, $[\text{heparin}]_{\text{total}} \gg [\text{RNAP}]_{\text{total}} \gg [\text{promoter}]_{\text{total}}$. The free [RNAP] is very small (generally $[\text{RNAP}]_{\text{free}} < [\text{promoter DNA}]_{\text{total}}$ see supplemental Table 3) because almost all RNAP is bound to heparin under the conditions of these experiments. Though not in excess, the free [RNAP] is constant throughout the time course of reversible association because it is buffered by the heparin-binding equilibrium. The heparin binding equilibrium is found to be established rapidly on the time scale of the promoter binding equilibrium, which takes many hours, so the observed kinetics are entirely the approach to equilibrium of promoter binding. The first order rate constants obtained from single exponential fits to these data (see Methods) are relaxation rate constants k_r , which are the sum of the pseudo-first order (constant RNAP concentration) association rate constant ($k_a[\text{R}]_{\text{free}}$) and the dissociation rate constant k_d (see Supplemental Table 3). Hence these decay-to-equilibrium experiments are less time-consuming than direct dissociation experiments. To obtain an accurate value for k_d from k_r , independently-determined values of k_a at each of these $[\text{Na}^+]$ in Glu^- and F^- buffers (Table 1) were used. Consequently k_d is well-determined, especially for conditions where $k_d \geq k_a[\text{R}]_{\text{free}}$.

Values of k_d obtained from the analyses of the data of Fig 6 and comparable series of reversible polymerase-promoter binding experiments performed over a range of $[\text{Na}^+]$ in Glu^- and F^- buffers are tabulated in Table 2 and plotted in Fig 7 as a log-log plot vs. $[\text{Na}^+]$ to compare with the behavior of k_d in Cl^- buffer. Values of k_d in Glu^- and F^- buffers fall on a common line on this log-log plot, greatly offset from k_d values in Cl^- buffer, and with a much shallower slope ($\text{Sk}_d = 3.2 \pm 0.8$ for Glu^-/F^- vs. $\text{Sk}_d = 9.2 \pm 0.1$ for Cl^-).

At 0.31 M Na^+ , the salt concentration requiring the least extrapolation of the data sets in Fig 7, k_d is $\sim 10^3$ -fold larger in Cl^- than in Glu^- or F^- buffers. This ratio is predicted to increase with increasing salt concentration if the log-log plot remains linear in Cl^- buffer. These differences in magnitude and log-log salt derivative of k_d between Cl^- and F^- or Glu^- are very large, exceeding those previously observed for equilibrium/rate constants of other protein processes, and also contrast with the behaviors seen in Fig 4 for K_1 , k_2 , and k_a (see Discussion), indicating that very different conformational changes and interfaces are involved in these different steps.

For all determinations of k_d from k_r , the constant free [RNAP] at each [heparin] investigated is also obtained. Values of the free [RNAP] range from $\sim 10^{-12}$ M at low $[\text{Na}^+]$ and high [heparin] to $\sim 10^{-10}$ M at high $[\text{Na}^+]$ and low [heparin]. Analysis of these decay to equilibrium experiments as a function of heparin concentration at constant salt concentration provides no evidence for subunit dissociation of RNAP holoenzyme at any condition investigated. From these free [RNAP], values of the heparin binding constant K_{heparin} are obtained at each $[\text{Na}^+]$ investigated in Glu^- and F^- buffers. Values of K_{heparin} are plotted as a log-log plot vs. $[\text{Na}^+]$ in Fig 8. In Glu^- and F^- buffers, the [salt] dependence of K_{heparin} ($\text{SK}_{\text{heparin}} = -13 \pm 0.2$) is similar to that of the RNAP-promoter binding constant ($\text{SK}_{\text{obs}} = \text{Sk}_a - \text{Sk}_d = -(8.3 \pm 0.2) + (3.2 \pm 0.8) = 11.5 \pm 1.0$).

DISCUSSION

Here we propose interpretations of the effects of [salt] and the difference between effects of Glu^- and F^- versus Cl^- on the steps of open complex formation in terms of extant structural and thermodynamic characterizations of each step. Qualitative comparisons are also made with

the effects of these salts on other protein-DNA interactions, on protein folding and DNA melting, and on model processes. Both the similarity of the effects of Glu^- and F^- and the large differences between effects of these anions and of Cl^- on the early and late steps of open complex formation indicate that these are Hofmeister effects and that they arise from greater exclusion of Glu^- and F^- than of Cl^- from nonpolar hydrocarbon surface, as proposed for protein folding (19) and for IHF-H'DNA binding (32). Fig 9 presents an overview of the steps, proposed conformational changes and the effects of [salt] and anion on each. We discuss how the Glu^- results here are predicted to affect σ^{70} holoenzyme interactions at strong promoters *in vivo* during the adaptation to an osmotic upshift.

Opening of the Initiation Bubble (−11 to +2) in the Active Site Cleft is Surprisingly Insensitive to [Salt] and Anion: Implications for Regulation of RP_0 Formation

In many biochemical processes, including enzyme catalysis and active transport, rapidly reversible initial and final steps bracket a bottleneck step that performs the key biological function (e.g. catalysis/transport). In these cases, regulation is largely achieved by effects of ligands or solutes on the rapidly reversible initial and final steps, but not on the key central catalytic/transport step of the mechanism. In RP_0 formation, the first and last steps rapidly equilibrate on the time scale of the cooperative DNA opening step (24). Here and in our previous work, we find large effects of [salt], anion and solutes on the initial and final steps, but *not* on the central bottleneck step that converts the closed complex I_1 to the unstable open complex I_2 . An exception is the action of far upstream DNA as an effector ligand. Truncation of DNA upstream of −47 greatly reduces the opening rate because it reduces the extent of insertion of downstream duplex DNA in the active site cleft in I_1 , thereby changing the nature of this intermediate (33).

Such a regulatory strategy would appear to be well-suited to organisms like *E. coli* that grow over a wide range of solution conditions, or in general to any cell responding to changes in environment. Clearly in order to reprogram genetic expression, DNA opening is required irregardless of solution conditions in the cell. The response to changes in growth conditions utilizes differences in promoter sequence, and/or shifts in the concentrations of sigma factors, “feedback” solutes (KGlu), and stress factors (ppGpp). It will be important to determine whether these factors act to increase or decrease levels of I_1 or I_2 , leaving the opening step unaffected, or whether they affect structural features of I_1 or I_2 which are central for DNA opening.

The small magnitude of the [salt] dependences of both rate constants of the DNA opening step ($\text{Sk}_2 = -1$, $\text{Sk}_{-2} = 0$, $\text{SK}_2 = -1$) contrast with the situation for DNA melting in solution, which is significantly more strongly [salt] dependent (25). Indeed this small [salt] dependence of DNA opening by RNAP appears inconsistent with proposed mechanisms based on structural data in which DNA opening occurs outside (above) the active site channel (34-36). To explain the small salt dependence of opening in the active site cleft, we propose that these very small effects of [salt] on a process (DNA opening) which is strongly [salt]-dependent in solution may indicate that the backbones of both DNA strands interact with polymerase throughout the process, and/or that compensation is present between ion uptake and release. Possibly loading DNA as a duplex in the highly positively charged cleft in I_1 (see above) neutralizes the charge in the region (−11 to +2) opened in forming I_2 . The subsequent movement of the template strand into the active site might then involve an exchange for basic contacts at the top of the cleft to those at the bottom, thereby buffering it from changes in [salt]. Compensations in the exchange between region 1.1 and the single strands in the channel (as well as unwrapping upstream DNA) may lead to small net changes in ASA, which in turn result in a small net change in binding free energy for the formation of I_2 from I_1 . In addition, opening occurs in a

deep cleft that is bounded by highly conserved structural features on the large subunits (37), which may in some way isolate the process from the bulk solution.

For the λP_R promoter at typical [salt], the initial open complex I_2 is unstable relative to I_1 at low temperatures ($<15^\circ\text{C}$), highly unstable relative to RP_o at higher temperatures (24), and therefore is only transiently populated at any temperature. Binding free energy (specifically a favorable entropy change, since the net enthalpy change for conversion of I_1 to I_2 is +25; (22,24)) drives the opening of the DNA in the channel. However, the small magnitude of SK_2 indicates that ion release is not the origin of the favorable entropy change for this step.

While the thermodynamic origins of this driving force remain unclear, we propose the following series of structural events drive opening in the conversion of I_1 to I_2 . Repositioning of σ region 1.1 in response to loading duplex DNA in the cleft in I_1 allows the strands to descend further the channel in $(I_1-I_2)^\ddagger$, increasing the bending distortion at $-12/-11$. This change likely exposes bases in the -10 hexamer on the NT strand (38) which are “captured” by the conserved aromatic side chains on region 2 of sigma (39-42). Additionally, a gradient of positive charge may exist in the channel, increasing from the top to the bottom. Such a distribution of lysines and arginines could help bring the template strand down to the base of the cleft once region 1.1 has moved.

For the entire bubble to open, we propose that a highly conserved region on the downstream lobe of β known as fork loop 2 (T. thermophilus (T. th.) 413-430, E. coli 533-550; (37)) may insert in the minor groove of the DNA duplex, as the DNA descends, prying apart the strands and helping to stabilize the open bubble surrounding +1. By analogy with base flipping enzymes, this insertion would further unwind the helix and create an additional $\sim 90^\circ$ bend near $-2/-1$. In the crystal structure of the T. th. transcription elongation complex, fork loop 2 is positioned between the nontemplate strand and the backbone of the RNA product (43).

Conversion of the Unstable Open Complex I_2 to the Stable Open Complex RP_o : Evidence for Large-Scale Folding and Assembly of a Clamp/Jaw on Downstream DNA

The finding here that replacement of Cl^- by Glu^- (or F^-) greatly increases the lifetime of RP_o at 25°C , especially at higher $[\text{Na}^+]$ is most consistent with a large-scale burial of hydrocarbon surface in the late steps (from which F^- and Glu^- are more completely excluded than is Cl^- (19)). This work along with our previous studies of k_d as a function of temperature, KCl, urea, and glycine betaine all indicate that large-scale rearrangements occur in the late steps of forming RP_o (24,27). These steps exhibit a large negative heat capacity change (-1 kcal/K), comparable to that for folding a 75-100 residue globular protein; the effect of urea is consistent with folding at least 100 residues, while the effects of [salt] and GB are consistent with formation of a significant new RNAP-DNA interface involving 10-15 DNA phosphates. The most plausible scenario is that >100 residues in RNAP fold and bind to the downstream DNA backbone in the conversion of I_2 to RP_o (24,27).

From an analysis which predicts natively disordered regions in E. coli RNAP, we deduced that the regions that fold are parts of the downstream clamp and jaw of β' (27). Highly conserved in bacterial RNAP (37), the following are positioned near the downstream DNA (+5 to +20) in the FRET model of RP_o (44) and the TEC structure (43): (i) a highly positively charged helix-hairpin-helix (T. th. 470-507, E. coli 194-232), (ii) a C-terminal region (T. th. 1378-1424, E. coli 1263-1310) adjacent to (i) and switch 5 (37); and (iii) the mobile downstream jaw (T. th. 1270-1329, E. coli 1142-1214). Regions (i) and (ii) form part of the ‘clamp’ of RNAP. Strong biochemical evidence exists for interactions between the clamp and jaw and downstream DNA in RP_o . Internal deletions in the jaw (45) as well as binding of the negatively charged phage protein Gp2 (46) to the jaw strongly destabilize RP_o at various promoters including λP_R . The helix-hairpin-helix binds near the major groove at +10 in the TEC; deletion

of *E. coli* 210-215 also strongly destabilizes RP_o (47). The lineage-specific sequence insertion 3 (SI3) in *E. coli* β' (48) likely also plays a part in the assembly of the stabilizing clamp/jaw. Comprised of a repeated sandwich-barrel-hybrid motif, β' SI3 connects to the “trigger loop” by a flexible 13 amino acid linker (49). Deletion of β' SI3 increases k_d at λP_R by ~4 fold (27). In a model of the TEC, β' SI3 is positioned near the jaw, the downstream lobe of beta and the downstream DNA from +13 to +18 (49).

We proposed that the clamp and jaw domains are mobile and partially disordered in I_2 and assemble on the downstream DNA in the conversion of I_2 to RP_o (27). Because the regions in the clamp/jaw contain runs of positive residues, their interactions with duplex DNA are predicted to release cations. Importantly, our data indicate that the RNAP machinery clamps onto the downstream DNA during $I_2 \rightarrow RP_o$, *after* the entire transcription bubble has been untwisted and opened in the active site channel ($I_1 \rightarrow I_2$; Gries et al., unpublished).

Predicting the Salt Concentration Dependence of Forming the 100 bp Wrapped DNA Interface in the Closed Intermediate I_1

The first steps of RP_o formation discriminate promoter from nonpromoter DNA by establishing interactions between sigma regions 4 and 2 and conserved six bp sequences at -35 and -10, respectively (41). However, formation of I_1 at the λP_R promoter creates an extensive interface with DNA that extends well beyond the -10 and -35 regions: protection from hydroxyl radical ($\bullet OH$) or DNase I cleavage of the DNA backbone in I_1 extends to at least +20 at λP_R (23,50) (and at the $\lambda P_{RM\ up1}$ promoter (51)); upstream protection from $\bullet OH$ cleavage at λP_R extends to ~-85 (23) (also apparently at the T7A1 promoter (2)). These data, interpreted using structures of the free RNAP (35,52) and a complex with a fork junction DNA (from -7 to -41; (34)), are most simply consistent with a model (23) where upstream DNA is wrapped around RNAP and downstream DNA is bound in the active site channel created by the opposition of the β and β' subunits (footnote 1). What is the relationship between this model of I_1 and SK_1 ?

In our model of I_1 , approximately sixty-two positive (arginine (R), lysine (K), histidine (H)) and twenty-five negative (aspartate (D), glutamate (E)) side chains lie near the DNA phosphate backbone, yielding a net charge of ~+37. With the exception of the C-terminal domain (CTD) of α (modeled using the crystal structure of the *E. coli* subunits (53)), this charge count is based on the *T. th.* RNAP subunits (αNTD , β , β' , ω , σ^A). However, the charged residues involved in forming the interface from -55 to +20 are highly conserved in bacterial RNAP. With the exception of three (R, K, H) and three (D, E) on β predicted to contact DNA from -11 to +5 in the model of I_1 , the remainder from the count above (R, K, H, E, D) are present in *E. coli*. While the one-to-one correspondence for charged residues in the region upstream of -55 is more variable between *T. th.* and *E. coli*, the overall charge distribution near the upstream wrapping track (-70 to -85) appears relatively conserved. This approximate structural estimate provides a reasonable basis for predicting the contribution from neutralization of DNA phosphate charge to the observed [salt] dependence of I_1 formation. For binding of an oligocation ligand with a charge of +37 to DNA, the predicted low-[salt] value of SK_{obs} is approximately -32 (or more negative if ion release from the ligand contributes to SK_{obs}).

Interpretation of the Much Smaller than Expected Effect of [Salt] on I_1 Formation

While interactions with the -35 and -10 hexamers involve base specific interactions with σ^{70} regions 4 and 2, respectively, the interface in I_1 is composed of predominantly nonspecific DNA backbone interactions with the other RNAP subunits (e.g. $\alpha CTDs$, β and β'). Our estimate

While our data do not allow us to determine the order of formation of these interactions, they have been determined by fast hydroxyl radical footprinting of early complexes at the T7A1 promoter (1,2). At T7A1, interactions are first established upstream and then proceed downstream (2). These early intermediates at λP_R rapidly equilibrate with one another and thus are not separated in the kinetic analysis.

of SK_1 (~ -30 (above)) is much larger in magnitude than the observed log-log dependence of K_1 on $[Na^+]$ ($SK_1 \sim -7$ in both Cl^- and Glu^- buffers). This difference suggests that some process involving extensive salt ion uptake by RNAP has evolved to largely compensate for this predicted cation release from the promoter DNA in forming I_1 from free RNAP and promoter DNA. We propose two scenarios involving cation uptake that may occur in I_1 formation: disruption of surface salt bridges on RNAP coupled to forming the interface (32, 54,55), and the placement of duplex DNA next to negatively charged region 1.1 in the channel.

Given the juxtaposition of negative and positive charges in the upstream half of the I_1 interface, we propose that coupled salt bridge disruption acts to reduce the binding constant K_1 and the magnitude of its log-log salt derivative $|SK_1|$, and likely explains the large negative ΔC_p of this step ($-1.4 \text{ kcal M}^{-1} \text{ K}^{-1}$; (22)). However, the walls and floor DNA active site channel of RNAP are highly basic. Given this charge distribution, it seems unlikely that salt bridge disruption could offset the salt ion release from binding >20 bp of downstream duplex in the channel. What does?

In free σ^{70} RNAP, the highly negatively charged region 1.1 lies in the active site channel (44). Where is it in I_1 ? Solute probes of changes in anionic (GB) and amide (urea) surface area do not detect any signature of release of sigma region 1.1 in forming I_1 (27). Instead, our estimates of the burial of anionic and amide surface in the I_1 protein-DNA interface based on structural data are completely consistent with the magnitude of effects of glycine betaine and urea on K_1 . As a result, we proposed that it was most likely that region 1.1 is not expelled during the formation of I_1 , remaining bound and masking the active site (27). It seems likely that any cation release from loading of duplex DNA in the channel must be offset by cation uptake as a result of placing the two highly negatively charged regions next to each other.

In the model of I_1 , DNA downstream of +1 lies above region 1.1 (23,27). Region 1.1 has a net charge of -19 and can be modeled as a negatively charged alpha-helical “plug” at the N-terminus (56) followed by an unstructured, negatively charged coil. We speculate that positioning downstream duplex DNA next to this negatively charged single stranded (ss) DNA mimic may be somewhat analogous to DNA triplex formation. Theoretically the limiting law per charge ion association is 0.76 for ss DNA, 0.88 for double stranded (ds) DNA, increasing to 0.92 in a triplex (ts). If this model is appropriate for placing 20 bp of duplex DNA near sigma region 1.1, then we predict approximately five cations are taken up.

Implications of Effects of Glu^- on the Steps of Initiation for Adaptation to Osmotic Stress in vivo

In vitro, Gralla observed that transcription from open complexes at the strong σ^{70} promoter lacUV5 decreases greatly as the K Glu concentration is increased from 0.1 to 0.4 M (9). If Cl^- were the cytoplasmic anionic osmolyte, such increases in K^+ concentrations would cause dissociation of RNAP and many other proteins from nucleic acids. However, in Glu^- , open complexes at the strong λP_R promoter are unexpectedly long-lived over the entire range of [salt] investigated here, ($k_d < 10^{-5} \text{ s}^{-1}$ at 25 °C even at 0.5 M Glu^- , corresponding to a lifetime ($1/k_d$) of open complexes exceeding one day). These data suggest that open complexes at strong σ^{70} promoters in vivo (where negative supercoiling should also favor RP_o formation) do not dissociate after an osmotic upshift, but may have difficulty escaping the promoter to transcribe.

While transcription from many σ^{70} holoenzyme-promoter is reduced at high Glu^- , transcription of osmotically-regulated promoters by σ^S holoenzyme increases (7,8). Intriguingly, in vitro DNase I footprints of σ^S RNAP-osmY promoter complexes reveal that contacts upstream of -30 are lost as $[KGlu]$ increases, allowing escape from the promoter (9). Gralla and coworkers deduced from these data that high concentrations of K^+ and Glu^- are able to release σ^S RNAP that are poised at lower salt ion concentrations, allowing both transcription and reinitiation

(9,57,58). By contrast, σ^{70} RNAP appears to remain bound an open promoter complex as [salt] increases in Glu^- buffer (this work), but its ability to transcribe is progressively reduced (9). Hence in vivo salt ion concentrations themselves appear capable both of switching transcription from one set of genes to another, and of poising RNAP to rapidly start at another set when conditions change (9,58,59).

Gralla and coworkers observe that the ability of an anion to increase σ^S transcription at osmotic genes is correlated with its position in the Hofmeister series (10). Since the exclusion of the Hofmeister anion Glu^- from hydrocarbon surfaces causes it to favor processes that reduce the exposure of nonpolar surface to the solution (e.g. folding, assembly, binding) (18), differences in the amounts and types of surface area changes in biological processes will in turn give rise to differential effects of Glu^- .

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

We thank Alison Huckenpahler for her contributions to the glutamate association kinetic data. We thank the editor and reviewers for their help in revising the manuscript.

Abbreviations

ASA	water-accessible surface area
GB	glycine betaine
PNAI	protein nucleic acid interactions
RNAP	RNA polymerase
σ^S	stationary phase sigma factor

References

1. Rogozina A, Zaychikov E, Buckle M, Heumann H, Sclavi B. DNA melting by RNA polymerase at the T7A1 promoter precedes the rate-limiting step at 37 °C and results in the accumulation of an off-pathway intermediate. *Nucleic Acids Res* 2009;37:5390–5404. [PubMed: 19578065]
2. Sclavi B, Zaychikov E, Rogozina A, Walther F, Buckle M, Heumann H. Real-time characterization of intermediates in the pathway to open complex formation by Escherichia coli RNA polymerase at the T7A1 promoter. *Proc Natl Acad Sci USA* 2005;102:4706–4711. [PubMed: 15738402]
3. Cayley S, Record MT Jr. Roles of cytoplasmic osmolytes, water, and crowding in the response of Escherichia coli to osmotic stress: biophysical basis of osmoprotection by glycine betaine. *Biochemistry* 2003;42:12596–12609. [PubMed: 14580206]
4. Wood JM. Osmosensing by bacteria: signals and membrane-based sensors. *Microbiol Mol Biol Rev* 1999;63:230–262. [PubMed: 10066837]
5. Cayley DS, Guttman HJ, Record MT Jr. Biophysical characterization of changes in amounts and activity of Escherichia coli cell and compartment water and turgor pressure in response to osmotic stress. *Biophys J* 2000;78:1748–1764. [PubMed: 10733957]
6. Gralla JD, Vargas DR. Potassium glutamate as a transcriptional inhibitor during bacterial osmoregulation. *EMBO J* 2006;25:1515–1521. [PubMed: 16541105]
7. Weber A, Jung K. Profiling early osmotic stress-dependent gene expression in Escherichia coli using DNA microarrays. *J Bacteriol* 2002;184:5502–5507. [PubMed: 12218039]

8. Cheung KJ, Badarinarayana V, Selinger DW, Janse D, Church GM. A microarray-based antibiotic screen identifies a regulatory role for supercoiling in the osmotic stress response of *Escherichia coli*. *Genome Res* 2003;13:206–215. [PubMed: 12566398]
9. Lee SJ, Gralla JD. Osmo-regulation of bacterial transcription via poised RNA polymerase. *Mol Cell* 2004;14:153–162. [PubMed: 15099515]
10. Gralla JD, Huo YX. Remodeling and activation of *Escherichia coli* RNA polymerase by osmolytes. *Biochemistry* 2008;47:13189–13196. [PubMed: 19053283]
11. Roe JH, Burgess RR, Record MT Jr. Kinetics and mechanism of the interaction of *Escherichia coli* RNA polymerase with the λP_R promoter. *J Mol Biol* 1984;176:495–522. [PubMed: 6235375]
12. Roe JH, Record MT Jr. Regulation of the kinetics of the interaction of *Escherichia coli* RNA polymerase with the λP_R promoter by salt concentration. *Biochemistry* 1985;24:4721–4726. [PubMed: 2934084]
13. Leirimo S, Harrison C, Cayley DS, Burgess RR, Record MT Jr. Replacement of potassium chloride by potassium glutamate dramatically enhances protein-DNA interactions in vitro. *Biochemistry* 1987;26:2095–2101. [PubMed: 2887198]
14. Barkley MD, Lewis PA, Sullivan GE. Ion effects on the lac repressor-operator equilibrium. *Biochemistry* 1981;20:3842–3851. [PubMed: 7272280]
15. Lohman TM, Chao K, Green JM, Sage S, Runyon GT. Large-scale purification and characterization of the *Escherichia coli* rep gene product. *J Biol Chem* 1989;264:10139–10147. [PubMed: 2524489]
16. Dragan AI, Li Z, Makeyeva EN, Milgotina EI, Liu Y, Crane-Robinson C, Privalov PL. Forces driving the binding of homeodomains to DNA. *Biochemistry* 2006;45:141–151. [PubMed: 16388589]
17. Maslak M, Martin CT. Effects of solution conditions on the steady-state kinetics of initiation of transcription by T7 RNA polymerase. *Biochemistry* 1994;33:6918–6924. [PubMed: 7911327]
18. Pegram LM, Record MT Jr. Thermodynamic origin of Hofmeister ion effects. *J Phys Chem B* 2008;112:9428–9436. [PubMed: 18630860]
19. Pegram LM, Wendorff T, Erdmann R, Shkel IA, Bellissimo D, Felitsky DJ, Record MT Jr. Why Hofmeister effects of many salts favor protein folding but not DNA helix formation. *Proc Natl Acad Sci USA*. 2010 in press.
20. Roe JH, Burgess RR, Record MT Jr. Temperature dependence of the rate constants of the *Escherichia coli* RNA polymerase- λP_R promoter interaction. Assignment of the kinetic steps corresponding to protein conformational change and DNA opening. *J Mol Biol* 1985;184:441–453. [PubMed: 3900414]
21. Tsodikov OV, Record MT Jr. General method of analysis of kinetic equations for multistep reversible mechanisms in the single-exponential regime: application to kinetics of open complex formation between $E\sigma^{70}$ RNA polymerase and σP_R promoter DNA. *Biophys J* 1999;76:1320–1329. [PubMed: 10049315]
22. Saecker RM, Tsodikov OV, McQuade KL, Schlax PE Jr, Capp MW, Record MT Jr. Kinetic studies and structural models of the association of *E. coli* σ^{70} RNA polymerase with the λP_R promoter: large scale conformational changes in forming the kinetically significant intermediates. *J Mol Biol* 2002;319:649–671. [PubMed: 12054861]
23. Davis CA, Bingman CA, Landick R, Record MT Jr, Saecker RM. Real-time footprinting of DNA in the first kinetically significant intermediate in open complex formation by *Escherichia coli* RNA polymerase. *Proc Natl Acad Sci U S A* 2007;104:7833–7838. [PubMed: 17470797]
24. Kontur WS, Saecker RM, Capp MW, Record MT Jr. Late steps in the formation of *E. coli* RNA polymerase- λP_R promoter open complexes: characterization of conformational changes by rapid [perturbant] upshift experiments. *J Mol Biol* 2008;376:1034–1047. [PubMed: 18191943]
25. Bloomfield, V.; Crothers, DM.; Tinoco, I., Jr., editors. *Nucleic Acids: Structures, Properties, and Functions*. University Science Books; Sausalito, CA: 2000.
26. Spolar RS, Record MT Jr. Coupling of local folding to site-specific binding of proteins to DNA. *Science* 1994;263:777–784. [PubMed: 8303294]
27. Kontur WS, Saecker RM, Davis CA, Capp MW, Record MT Jr. Solute probes of conformational changes in open complex (RP_o) formation by *Escherichia coli* RNA polymerase at the λP_R promoter: evidence for unmasking of the active site in the isomerization step and for large-scale coupled folding in the subsequent conversion to RP_o . *Biochemistry* 2006;45:2161–2177. [PubMed: 16475805]

28. Burgess RR, Jendrisak JJ. A procedure for the rapid, large-scale purification of *Escherichia coli* DNA-dependent RNA polymerase involving Polymin P precipitation and DNA-cellulose chromatography. *Biochemistry* 1975;14:4634–4638. [PubMed: 1101952]
29. Craig ML, Suh WC, Record MT Jr. HO and DNase I probing of $E\sigma^{70}$ RNA polymerase- λP_R promoter open complexes: Mg^{2+} binding and its structural consequences at the transcription start site. *Biochemistry* 1995;34:15624–15632. [PubMed: 7495790]
30. Leirmo, S. The Mechanism of Interaction of *E. coli* RNA Polymerase with Bacteriophage and Bacterial Promoters. Department of Biochemistry, University of Wisconsin-Madison; Madison: 1989.
31. Glaser BT, Bergendahl V, Anthony LC, Olson B, Burgess RR. Studying the salt dependence of the binding of σ^{70} and σ^{32} to core RNA polymerase using luminescence resonance energy transfer. *PLoS One* 2009;4:e6490. [PubMed: 19649256]
32. Vander Meulen KA, Saecker RM, Record MT Jr. Formation of a wrapped DNA-protein interface: experimental characterization and analysis of the large contributions of ions and water to the thermodynamics of binding IHF to H' DNA. *J Mol Biol* 2008;377:9–27. [PubMed: 18237740]
33. Davis CA, Capp MW, Record MT Jr. Saecker RM. The effects of upstream DNA on open complex formation by *Escherichia coli* RNA polymerase. *Proc Natl Acad Sci U S A* 2005;102:285–290. [PubMed: 15626761]
34. Murakami KS, Masuda S, Campbell EA, Muzzin O, Darst SA. Structural basis of transcription initiation: an RNA polymerase holoenzyme-DNA complex. *Science* 2002;296:1285–1290. [PubMed: 12016307]
35. Vassilyev DG, Sekine S, Laptenko O, Lee J, Vassilyeva MN, Borukhov S, Yokoyama S. Crystal structure of a bacterial RNA polymerase holoenzyme at 2.6 Å resolution. *Nature* 2002;417:712–719. [PubMed: 12000971]
36. Murakami KS, Darst SA. Bacterial RNA polymerases: the whole story. *Curr Opin Struct Biol* 2003;13:31–39. [PubMed: 12581657]
37. Lane WJ, Darst SA. Molecular evolution of multisubunit RNA polymerases: structural analysis. *J Mol Biol* 2010;395:686–704. [PubMed: 19895816]
38. Schroeder LA, Gries TJ, Saecker RM, Record MT Jr. Harris ME, DeHaseth PL. Evidence for a tyrosine-adenine stacking interaction and for a short-lived open intermediate subsequent to initial binding of *Escherichia coli* RNA polymerase to promoter DNA. *J Mol Biol* 2009;385:339–349. [PubMed: 18976666]
39. Tomsic M, Tsujikawa L, Panaghie G, Wang Y, Azok J, deHaseth PL. Different roles for basic and aromatic amino acids in conserved region 2 of *Escherichia coli* σ^{70} in the nucleation and maintenance of the single-stranded DNA bubble in open RNA polymerase-promoter complexes. *J Biol Chem* 2001;276:31891–31896. [PubMed: 11443133]
40. Schroeder LA, Karpen ME, deHaseth PL. Threonine 429 of *Escherichia coli* σ^{70} is a key participant in promoter DNA melting by RNA polymerase. *J Mol Biol* 2008;376:153–165. [PubMed: 18155246]
41. Helmann JD, deHaseth PL. Protein-nucleic acid interactions during open complex formation investigated by systematic alteration of the protein and DNA binding partners. *Biochemistry* 1999;38:5959–5967. [PubMed: 10320321]
42. Fenton MS, Lee SJ, Gralla JD. *Escherichia coli* promoter opening and -10 recognition: mutational analysis of σ^{70} . *Embo J* 2000;19:1130–1137. [PubMed: 10698953]
43. Vassilyev DG, Vassilyeva MN, Perederina A, Tahirov TH, Artsimovitch I. Structural basis for transcription elongation by bacterial RNA polymerase. *Nature* 2007;448:157–162. [PubMed: 17581590]
44. Mekler V, Kortkhonja E, Mukhopadhyay J, Knight J, Revyakin A, Kapanidis AN, Niu W, Ebright YW, Levy R, Ebright RH. Structural organization of bacterial RNA polymerase holoenzyme and the RNA polymerase-promoter open complex. *Cell* 2002;108:599–614. [PubMed: 11893332]
45. Ederth J, Artsimovitch I, Isaksson LA, Landick R. The downstream DNA jaw of bacterial RNA polymerase facilitates both transcriptional initiation and pausing. *J Biol Chem* 2002;277:37456–37463. [PubMed: 12147705]
46. Camara B, Liu M, Reynolds J, Shadrin A, Liu B, Kwok K, Simpson P, Weinzierl R, Severinov K, Cota E, Matthews S, Wigneshweraraj SR. T7 phage protein Gp2 inhibits the *Escherichia coli* RNA

- polymerase by antagonizing stable DNA strand separation near the transcription start site. *Proc Natl Acad Sci USA* 2010;107:2247–2252. [PubMed: 20133868]
47. Bartlett MS, Gaal T, Ross W, Gourse RL. RNA polymerase mutants that destabilize RNA polymerase-promoter complexes alter NTP-sensing by *rrn* P1 promoters. *J Mol Biol* 1998;279:331–345. [PubMed: 9642041]
 48. Lane WJ, Darst SA. Molecular evolution of multisubunit RNA polymerases: sequence analysis. *J Mol Biol* 2010;395:671–685. [PubMed: 19895820]
 49. Chlenov M, Masuda S, Murakami KS, Nikiforov V, Darst SA, Mustaev A. Structure and function of lineage-specific sequence insertions in the bacterial RNA polymerase β' subunit. *J Mol Biol* 2005;353:138–154. [PubMed: 16154587]
 50. Craig ML, Tsodikov OV, McQuade KL, Schlax PE Jr, Capp MW, Saecker RM, Record MT Jr. DNA footprints of the two kinetically significant intermediates in formation of an RNA polymerase-promoter open complex: evidence that interactions with start site and downstream DNA induce sequential conformational changes in polymerase and DNA. *J Mol Biol* 1998;283:741–756. [PubMed: 9790837]
 51. Li XY, McClure WR. Characterization of the closed complex intermediate formed during transcription initiation by *Escherichia coli* RNA polymerase. *J Biol Chem* 1998;273:23549–23557. [PubMed: 9722594]
 52. Murakami KS, Masuda S, Darst SA. Structural basis of transcription initiation: RNA polymerase holoenzyme at 4 Å resolution. *Science* 2002;296:1280–1284. [PubMed: 12016306]
 53. Benoff B, Yang H, Lawson CL, Parkinson G, Liu J, Blatter E, Ebright YW, Berman HM, Ebright RH. Structural basis of transcription activation: the CAP- α CTD-DNA complex. *Science* 2002;297:1562–1566. [PubMed: 12202833]
 54. Holbrook JA, Tsodikov OV, Saecker RM, Record MT Jr. Specific and non-specific interactions of integration host factor with DNA: thermodynamic evidence for disruption of multiple IHF surface salt-bridges coupled to DNA binding. *J Mol Biol* 2001;310:379–401. [PubMed: 11428896]
 55. Saecker RM, Record MT Jr. Protein surface salt bridges and paths for DNA wrapping. *Curr Opin Struct Biol* 2002;12:311–319. [PubMed: 12127449]
 56. Schwartz EC, Shekhtman A, Dutta K, Pratt MR, Cowburn D, Darst S, Muir TW. A full-length group 1 bacterial sigma factor adopts a compact structure incompatible with DNA binding. *Chem Biol* 2008;15:1091–1103. [PubMed: 18940669]
 57. Huo YX, Rosenthal AZ, Gralla JD. General stress response signalling: unwrapping transcription complexes by DNA relaxation via the σ^{38} C-terminal domain. *Mol Microbiol* 2008;70:369–378. [PubMed: 18761624]
 58. Rosenthal AZ, Kim Y, Gralla JD. Poising of *Escherichia coli* RNA polymerase and its release from the σ^{38} C-terminal tail for *osmY* transcription. *J Mol Biol* 2008;376:938–949. [PubMed: 18201723]
 59. Rosenthal AZ, Hu M, Gralla JD. Osmolyte-induced transcription: –35 region elements and recognition by σ^{38} (rpoS). *Mol Microbiol* 2006;59:1052–1061. [PubMed: 16420371]

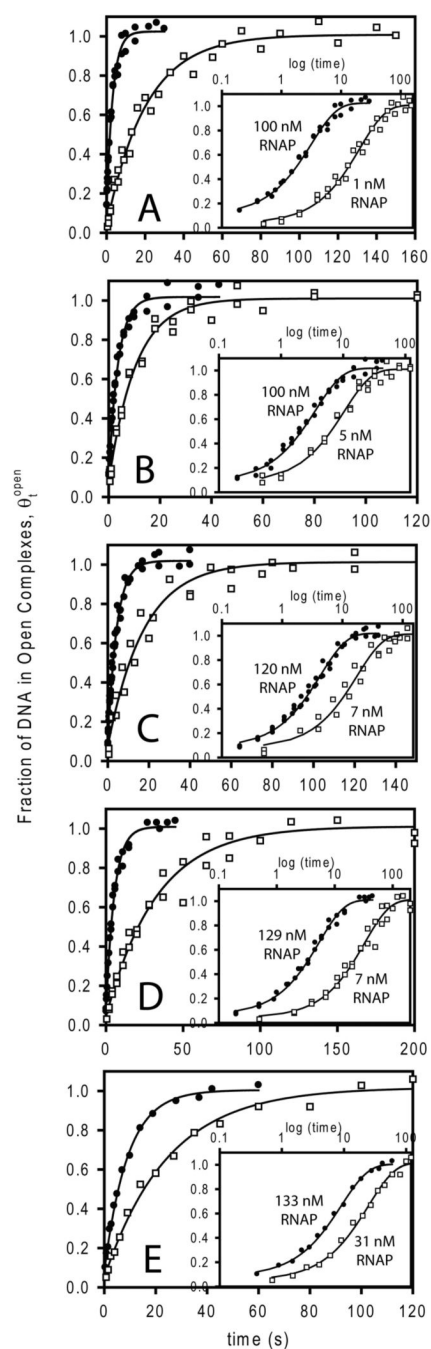


Figure 1.

Representative kinetic data for formation of open complexes at the λP_R promoter at high (closed circles) and low (open squares) concentrations of active RNAP (where RNAP is in excess over promoter DNA) at five different $[Na^+]$ at 25 °C in NaCl buffer. All data shown here were obtained using rapid quench mixing; the fraction of promoter DNA in the form of open complexes θ_t^{open} as a function of time (t) was determined using nitrocellulose filter binding after association reactions were quenched with the competitor heparin. Theoretical curves through the data are the best fits to eq 1; corresponding first order irreversible rate constants k_{obs} for each [RNAP] are given in Supplemental Table 1. Insets replot the data as a function of $\log t$,

demonstrating the single exponential character of the kinetics over the entire time range. (A) 0.15 M Na⁺. (B) 0.17 M Na⁺. (C) 0.19 M Na⁺. (D) 0.21 M Na⁺. (E) 0.23 Na⁺.

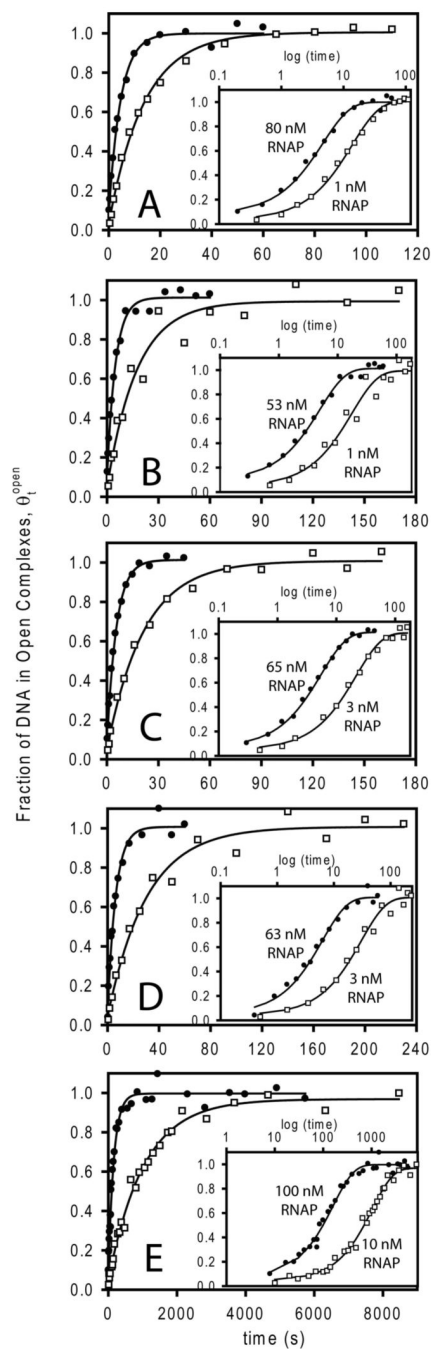


Figure 2.

Representative kinetic data for formation of open complexes at the λP_R promoter at representative high (closed circles) and low (open squares) concentrations of active RNAP at five different $[\text{Na}^+]$ at 25 °C in NaGlu buffer. (Details and insets as in Fig 1; values of k_{obs} for each $[\text{RNAP}]$ are given in Supplemental Table 2.) With the exception of (E), all data were obtained using rapid quench mixing. (A) 0.23 M Na^+ . (B) 0.25 M Na^+ . (C) 0.27 M Na^+ . (D) 0.29 M Na^+ . (E) 0.55 M Na^+ .

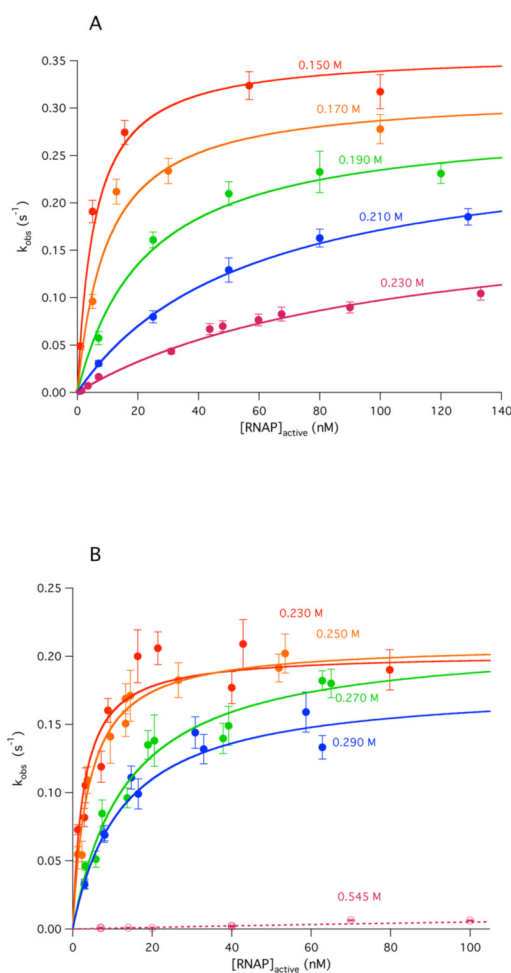


Figure 3.

Irreversible forward rate constant k_{obs} plotted as a function of the total, active RNAP concentration at five different $[\text{Na}^+]$ in Cl⁻ buffer (cf. Fig 1) (A) and in Glu⁻ buffer (cf. Fig 2) (B). With the exception of data obtained by manual mixing at 0.55 M Na⁺ in Glu⁻, data were obtained by rapid mixing and nonlinearly fit to eq 2, yielding values of K_1 , k_2 and k_a (see Table 1) for each $[\text{Na}^+]$ investigated.

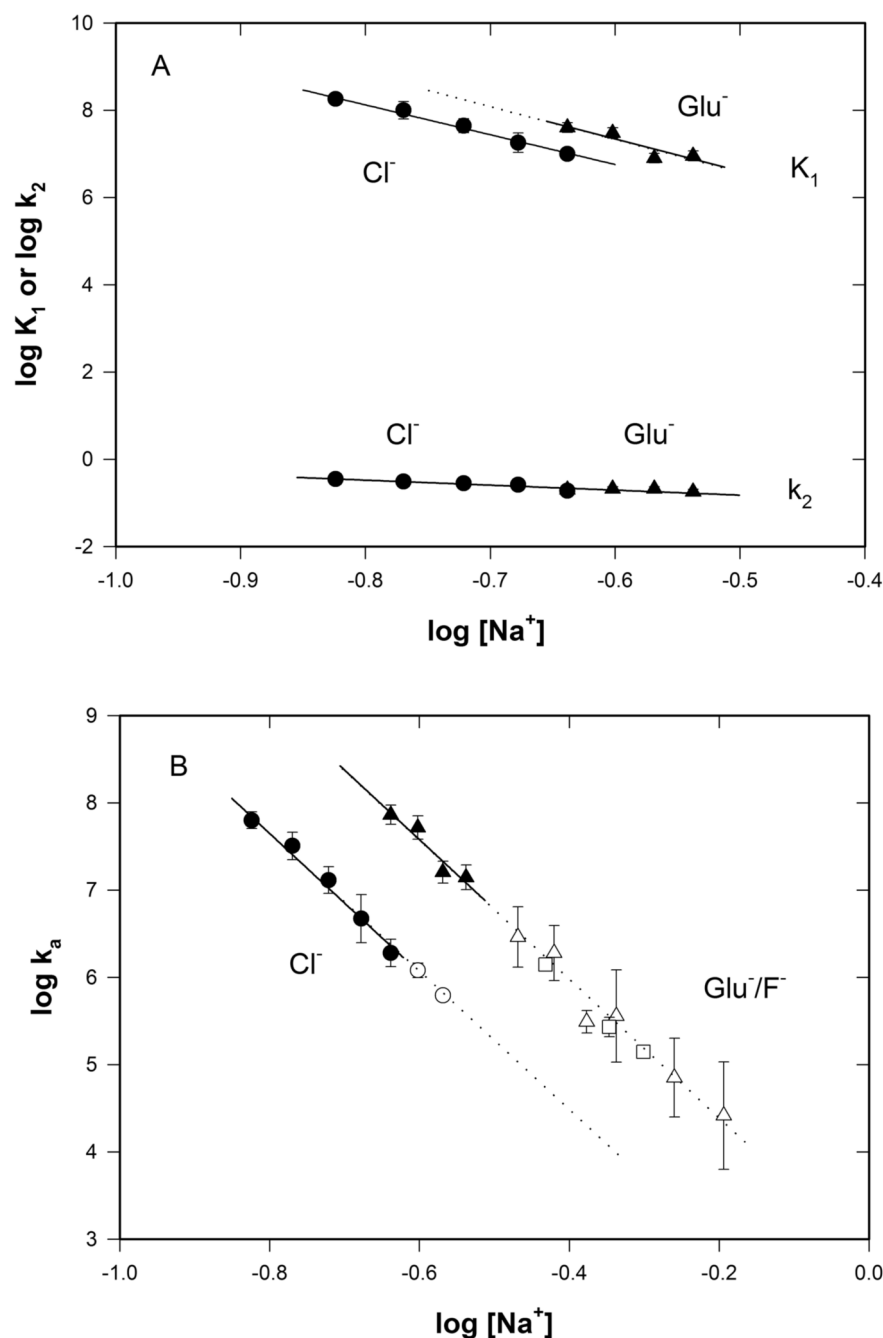


Figure 4.

Log-log $[\text{Na}^+]$ -dependence of K_1 , k_2 and k_a as a function of anion. (A) Data from rapid mixing experiments (cf. Fig 3) fit to eq 2 allow a dissection of $k_a (=K_1k_2)$ into individual values of K_1 and k_2 in NaCl (circle) and NaGlu (triangle) buffers. (B) Values of $\log k_a$ as a function of $\log [\text{Na}^+]$ in NaCl (circle), NaGlu (triangle), and NaF (square) buffers. Closed and open symbols represent values of k_a obtained using rapid and manual mixing, respectively. Dashed lines represent linear extrapolations of the log-log fits to the individual data sets, drawn for purposes of comparing the slopes for the different anions. Values of the log-log slopes Sk_a , SK_1 and Sk_2 are given in Table 1 (see Methods for details of the fitting).

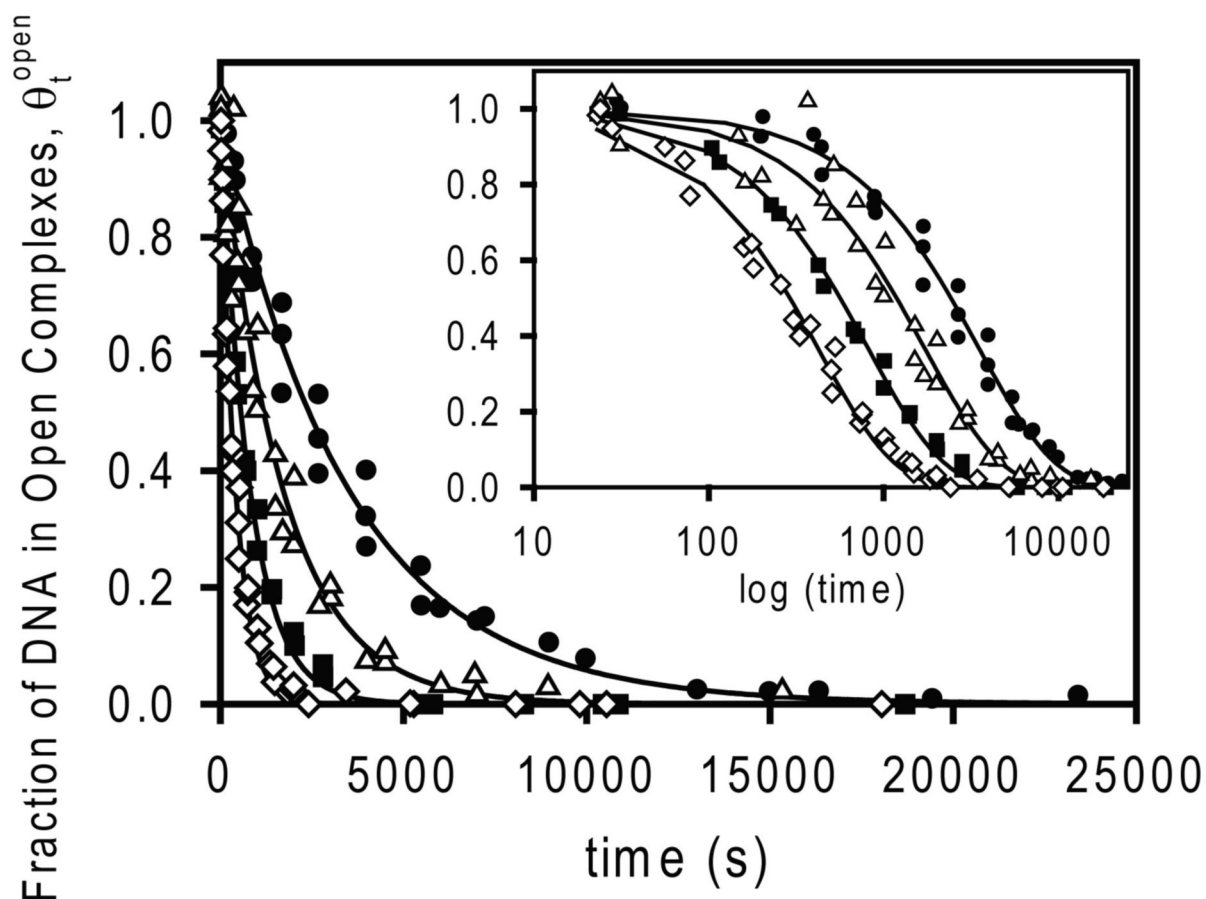


Figure 5.

Kinetics of the irreversible dissociation of open complexes at the λP_R promoter in NaCl buffer at 25 °C. Representative values of θ_t^{open} as a function of time (t) determined using nitrocellulose filter binding at four different $[Na^+]$: 0.23 M (filled circles), 0.25 M (open triangles), 0.27 M (filled squares) and 0.29 M Na^+ (open diamonds). Theoretical curves through the data are the best fits to eq 4; corresponding first order dissociation rate constants k_d for each [RNAP] are given in Table 2. Insets replot θ_t^{open} as a function of $\log t$, demonstrating the single exponential character of the kinetics over the entire time range.

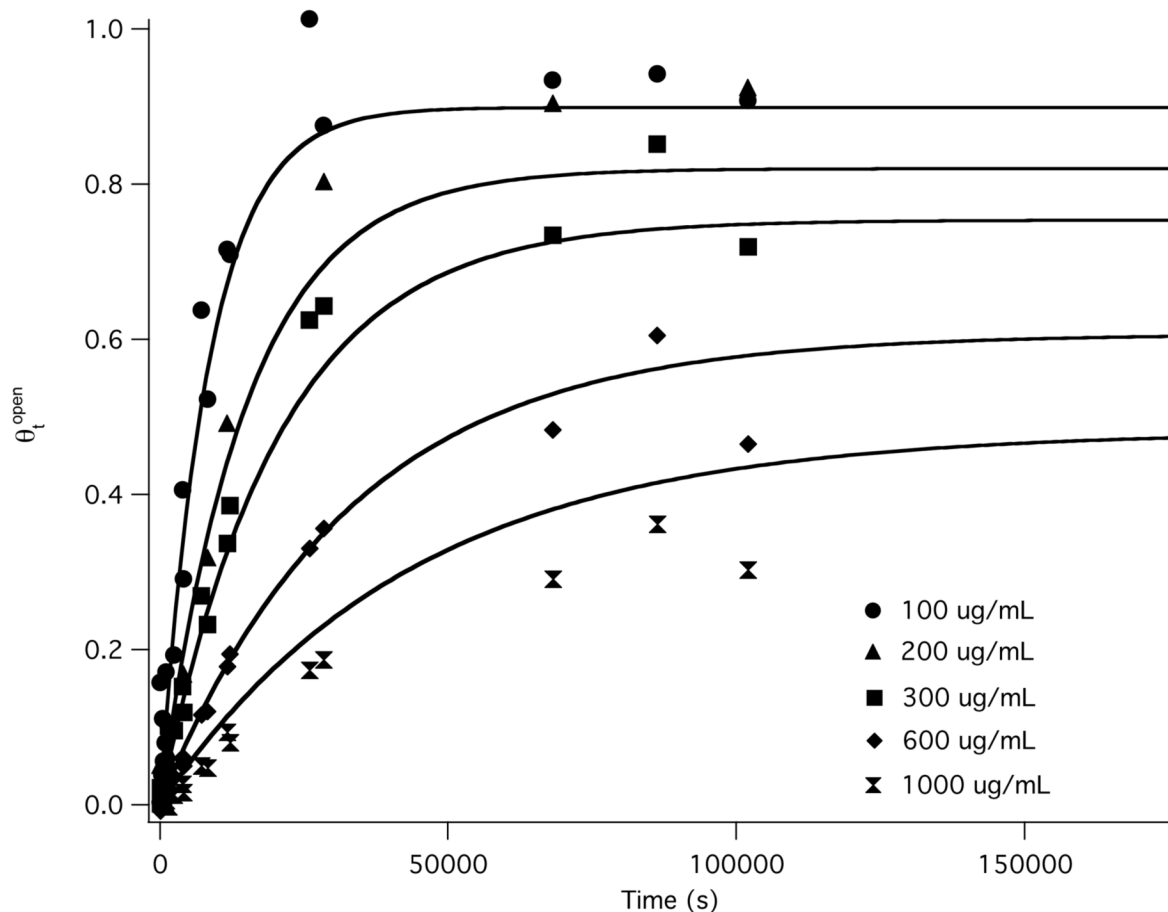


Figure 6.

Decay to equilibrium kinetics of promoter association-dissociation at high glutamate (25 °C, 0.46 M Glu⁻) and low, constant RNAP concentrations buffered by heparin. The fraction of promoter DNA in open complexes θ_t^{open} is plotted as a function of $\log t$, where t is the time after addition of promoter DNA (12.5 pM final) to a solution of RNAP (10 nM total concentration) and heparin at total concentrations of 100 $\mu\text{g/mL}$, 200 $\mu\text{g/mL}$, 300 $\mu\text{g/mL}$, 600 $\mu\text{g/mL}$ and 1000 $\mu\text{g/mL}$. Data points are from two independent experiments at each heparin concentration are plotted, except for 200 $\mu\text{g/mL}$ which was done once. Fits shown are to the single-exponential decay to equilibrium kinetic equations derived for this situation (eqs 5-6 in Methods). From the observed first order rate constant, analyzed using the best fit value of the association rate constant k_a for these conditions (Fig 4B), the dissociation rate constant k_d , the free RNA polymerase concentration, and the equilibrium binding constant K_{heparin} for binding RNAP to heparin are determined (see Table 3 and supplemental Table 3).

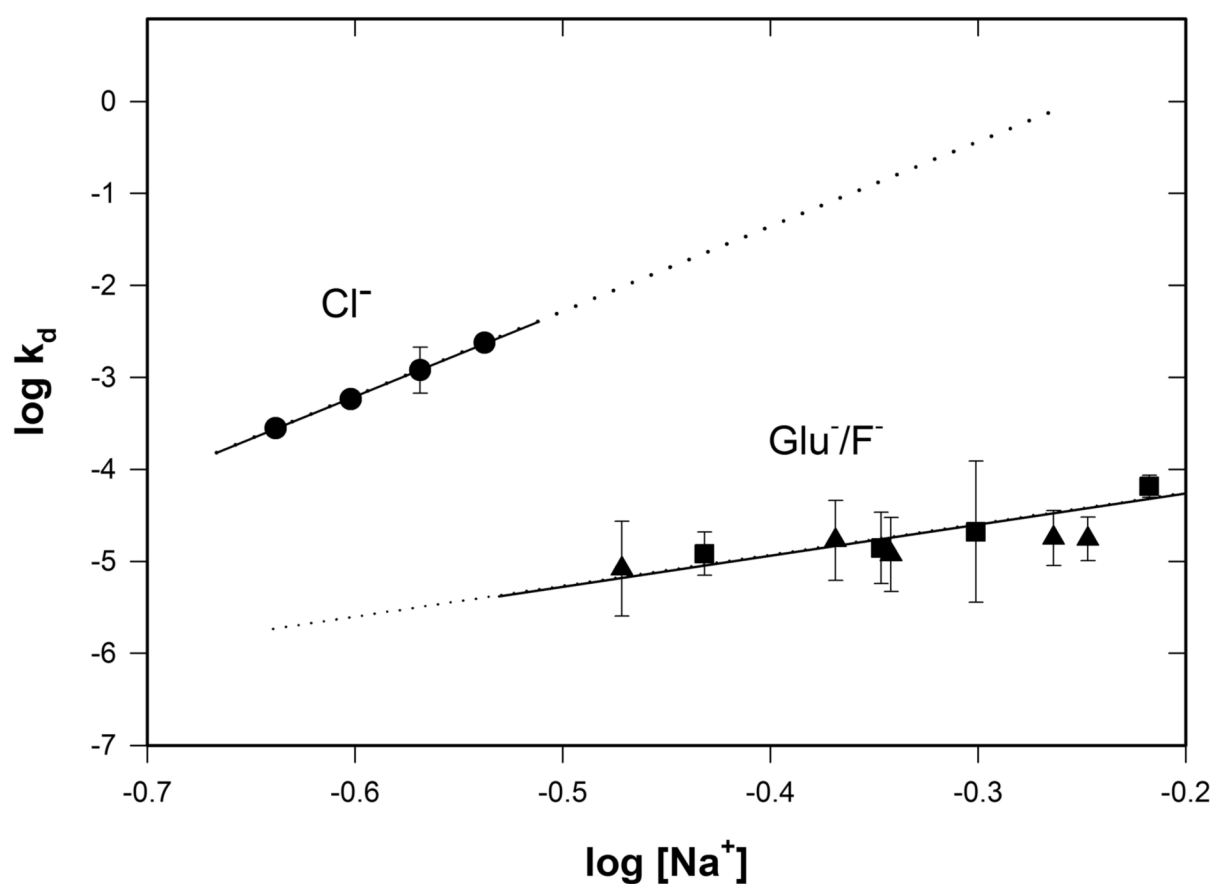


Figure 7.

Log-log plot of dissociation rate constants k_d of RNAP- $\lambda\sigma_R$ promoter open complexes as a function of $[\text{Na}^+]$ in Cl^- and in Glu^- or F^- buffers at 25 °C. Values of k_d in Cl^- are from irreversible dissociation experiments of Fig 5; values of k_d in Glu^- and F^- are from reversible dissociation experiments like those in Fig 6. Values of k_d are reported in Table 2.

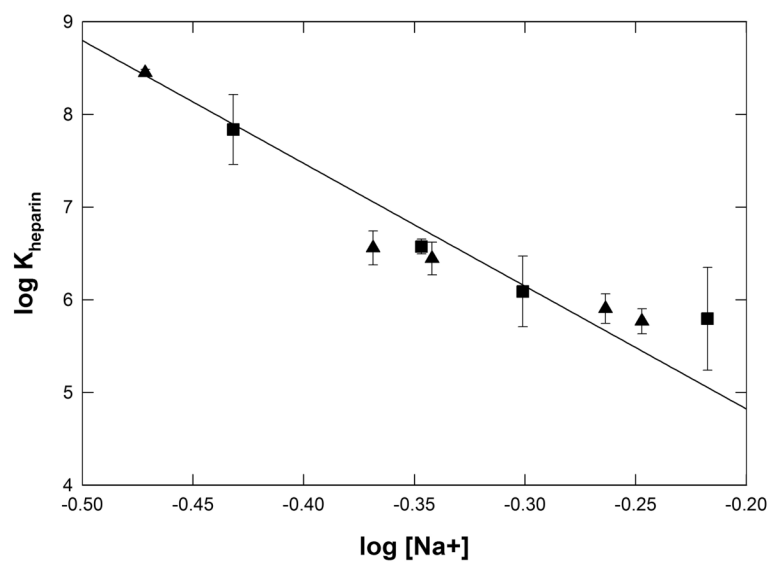


Figure 8.

Log-log plot of the heparin binding constant K_{heparin} as a function of $[\text{Na}^+]$ for experiments in Glu^- and F^- buffers at 25 °C. A linear fit of these data yields $SK_{\text{heparin}} = (\text{dlog}K_{\text{heparin}}/\text{dlog}[\text{Na}^+]) = -13.2 \pm 0.8$ and an extrapolated 1 M Na^+ intercept of 2.2 ± 0.3 .

Open Complex Formation: Proposals for Large-Scale Conformational Changes

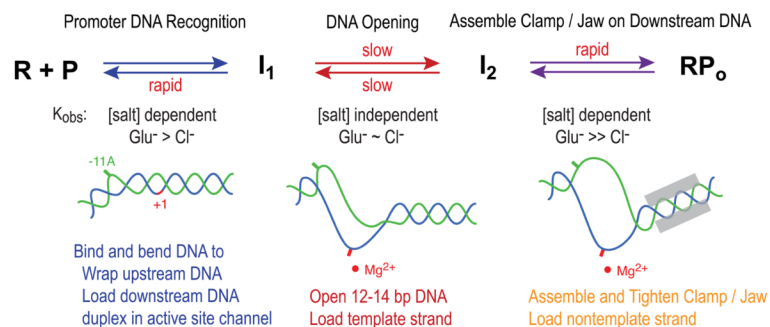


Figure 9.

Overview of the steps of RP_0 formation, proposed conformational changes and the effects of [salt] and anion on each. Schematic based on results in (23,24,27,33, Gries et al., unpublished).

Table 1

Values of k_a , K_1 and k_2 as a function of Na^+ and anion (Cl^- , Glu^- or F^-) for RP_0 formation at the λPR promoter at 25 °C^a

	Na^+ (M)	k_a ($\text{M}^{-1}\text{s}^{-1}$)			K_1 (M^{-1})			k_2 (s^{-1})	
		Cl^-	Glu^-	F^-	Cl^-	Glu^-	Cl^-	Cl^-	Glu^-
A	0.15	$(6.3 \pm 0.6) \times 10^7$			$(1.8 \pm 0.2) \times 10^8$			$(3.5 \pm 0.1) \times 10^{-1}$	
	0.17	$(3.2 \pm 0.5) \times 10^7$			$(1.0 \pm 0.2) \times 10^8$			$(3.1 \pm 0.1) \times 10^{-1}$	
	0.19	$(1.3 \pm 0.2) \times 10^7$			$(4.4 \pm 0.7) \times 10^7$			$(2.8 \pm 0.2) \times 10^{-1}$	
	0.21	$(4.7 \pm 1.3) \times 10^6$			$(1.8 \pm 0.4) \times 10^7$			$(2.6 \pm 0.3) \times 10^{-1}$	
	0.23	$(1.9 \pm 0.3) \times 10^6$	$(7.3 \pm 0.8) \times 10^7$		$(9.9 \pm 1.2) \times 10^6$	$(3.6 \pm 0.4) \times 10^8$	$(1.9 \pm 0.2) \times 10^{-1}$		$(2.0 \pm 0.1) \times 10^{-1}$
	0.25		$(5.2 \pm 0.7) \times 10^7$			$(2.5 \pm 0.3) \times 10^8$		$(2.1 \pm 0.1) \times 10^{-1}$	
	0.27		$(1.6 \pm 0.2) \times 10^7$			$(7.3 \pm 0.8) \times 10^7$		$(2.1 \pm 0.1) \times 10^{-1}$	
	0.29		$(1.4 \pm 0.2) \times 10^7$			$(8.0 \pm 0.9) \times 10^7$		$(1.8 \pm 0.1) \times 10^{-1}$	
SK ^b		-8.0 ± 0.6	-8.0 ± 1.7		-6.8 ± 0.3	-7.5 ± 2.3	-1.1 ± 0.2	-0.9 ± 0.3	
B	0.25	$(1.2 \pm 0.1) \times 10^6$							
	0.27	$(6.2 \pm 0.3) \times 10^5$							
	0.34		$(2.9 \pm 1.0) \times 10^6$						
	0.37			$(1.4 \pm 0.1) \times 10^6$					
	0.38		$(1.9 \pm 0.6) \times 10^6$						
	0.42		$(3.1 \pm 0.4) \times 10^5$						
	0.45			$(2.7 \pm 0.3) \times 10^5$					
	0.46		$(3.6 \pm 1.9) \times 10^5$						
	0.50			$(1.4 \pm 0.1) \times 10^5$					
	0.55		$(7.1 \pm 3.2) \times 10^4$						
	0.64		$(2.6 \pm 1.6) \times 10^4$						
SK ^c		-8.0 ± 0.2	-8.3 ± 0.2						

^aFor 0.15-0.23 M Na^+ in NaCl buffer and 0.23-0.29 M Na^+ in NaGlu buffer (part A of the table), values of k_a , K_1 and k_2 determined from fits of rapid quench mixing data to eq 2. For 0.34 – 0.64 M Na^+ (part B of the table), the value of k_a was determined from fits of manual mixing data to eq 3; values of k_{obs} as a function of [RNAP] in this [salt] range do not determine K_1 and k_2 (see text).

^bSK = $-\text{dlog}X/\text{dlog}[Na^+]$ where X = k_a , K_1 or k_2 ; each SK value determined from fits of the data in section A of the table.

^cValues of SK determined from fitting data from section A and B.

Table 2

Values of k_d as a function of Na^+ and anion (Cl^- , Glu^- or F^-) for the dissociation of RP_O at the λP_R promoter at 25 °C

Na^+ (M)	k_d (s^{-1})		
	Cl^-	Glu^-	F^-
0.23	$(2.8 \pm 0.1) \times 10^{-4}$		
0.25	$(5.8 \pm 0.4) \times 10^{-4}$		
0.27	$(1.2 \pm 0.3) \times 10^{-3}$		
0.29	$(2.37 \pm 0.09) \times 10^{-3}$		
0.34		$(8 \pm 4) \times 10^{-6}$	
0.37			$(1.2 \pm 0.3) \times 10^{-5}$
0.43		$(1.7 \pm 0.7) \times 10^{-6}$	
0.45			$(1.4 \pm 0.5) \times 10^{-5}$
0.46		$(1.2 \pm 0.5) \times 10^{-5}$	
0.50			$(2.1 \pm 0.2) \times 10^{-5}$
0.55		$(1.8 \pm 0.5) \times 10^{-5}$	
0.57		$(1.8 \pm 0.4) \times 10^{-5}$	
0.61			$(6.5 \pm 0.8) \times 10^{-5}$
Sk_d	9.2 ± 0.1	3.2 ± 0.8	

Values of k_d in Cl^- buffer determined by direct measure of the irreversible rate of dissociation of RP_O initiated by the addition of heparin. Values of k_d in Glu^- and F^- buffer determined using a decay to equilibrium experiment (eqs 5-6, see Methods) at 25 °C. $\text{Sk}_d \equiv (\partial \log k_d / \partial \log [\text{Na}^+])$. Values of k_d in Glu^- do not differ from those in F^- outside of experimental uncertainty and were fit together to determine Sk_d .