The mechanism and high-free-energy transition state of lac repressor–lac operator interaction

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ABSTRACT

Significant, otherwise-unavailable information about mechanisms and transition states (TS) of protein folding and binding is obtained from solute effects on rate constants. Here we characterize TS for lac repressor (R)–lac operator (O) binding by analyzing effects of RO-stabilizing and RO-destabilizing solutes on association (kₐ) and dissociation (k₈) rate constants. RO-destabilizing solutes (urea, KCl) reduce k₈ comparably (urea) or more than (KCl) they increase kₐ, demonstrating that they destabilize TS relative to reactants and RO, and that TS exhibits most of the Coulombic interactions between R and O. Strikingly, three solutes which stabilize RO by favoring burial/dehydration of amide oxygens and anionic phosphate oxygens all reduce k₈ without affecting kₐ significantly. The lack of stabilization of TS by these solutes indicates that O phosphates remain hydrated in TS and that TS preferentially buries aromatic carbons and amide nitrogens while leaving amide oxygens exposed. In our proposed mechanism, DNA-binding-domains (DBD) of R insert in major grooves of O pre-TS, forming most Coulombic interactions of RO and burying aromatic carbons. Nucleation of hinge helices creates TS, burying sidechain amide nitrogens. Post-TS, hinge helices assemble and the DBD-hinge helix-O DNA module docks on core repressor, partially dehydrating phosphate oxygens and tightening all interfaces to form RO.

INTRODUCTION

The lac repressor (R)–lac operator (O) interaction is a classic example of an on–off switch in prokaryotic gene regulation (1–4). Large conformational changes in both repressor and operator occur in forming the highly stable RO complex (5–13). These are reversed by binding of feedback-control inducer ligands, dissociating the RO complex to allow expression of the lac genes (14–17). For an O site embedded in polymeric DNA, the kinetics of RO formation at low pM concentrations of R and O were determined and interpreted in terms of facilitated diffusion of R on DNA after non-specific binding (18–22). To obtain information about the series of conformational changes in RO formation once R has located O and about the high-free-energy transition state (TS) in this mechanism, studies with small O-containing DNA fragments under conditions where diffusion to O is not rate-limiting are needed. In a classic study, Whitson and Matthews (23) determined and compared effects of salt and temperature on rate constants for RO formation and dissociation on a 40 bp fragment and on polymeric DNA.

A plausible mechanism of RO formation can be proposed on the basis of crystallographic (7,8) and nuclear magnetic resonance (NMR) (9–11) structural information. This process starts with insertion of the flexibly-tethered DNA binding domains (DBD) of R into the major grooves of O, followed by folding of the flexible tethers (hinge regions) into α-helices and insertion in the central minor groove, bending O. Folding of the tethers allows the DBD-hinge helices-O DNA assembly to interact with the core of R (24,25). While certainly a logical order of interactions and conformational changes, this structural mechanism has not been tested by kinetic studies. Here we report solute effects on association (kₐ) and dissociation (k₈) rate constants, determined under conditions where diffusion of R to the O site is not rate-determining. Interpretation of these data, using solute-model compound interaction information (α-values) and molecular modelling, provides information about interfaces and conformational changes involved in forming the high-free-energy transition state (TS) that determines these rate constants, and thereby tests the structure-based proposal for the mechanism of RO formation.

A large reduction in water accessible surface areas (ΔASA) of R and O (i.e. a large negative ΔASA) occurs in the RO interaction. Large ΔASA also occur in most other self-assembly interactions of proteins, nucleic acids and their complexes, as well as in the steps of operation of these assemblies as molecular machines. The ASA reduction in RO

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formation occurs as $R$ and $O$ functional groups reduce or eliminate their interactions with water and solutes to form interfaces of the $RO$ complex. Solute effects on rate constants for $RO$ formation and decomposition provide information about the amount and composition of the $\Delta ASA$ of formation of the high-free-energy transition state from reactants, just as effects of temperature on these rate constants provide information about the enthalpic barriers for forming $TS$ from reactants.

While structural information about $TS$ of enzyme-catalyzed reactions has been obtained using stable $TS$ analogs (26,27), no similar structural approach is generally available for non-covalent $TS$. To determine what conformational changes occur and what order they occur in, general probes that are sensitive to ASA changes in mechanistic steps are needed. Because solutes affect biopolymer processes by interacting with the biopolymer surface exposed or buried in the process (28), solutes are a natural choice as both thermodynamic and mechanistic (kinetic) probes of ASA changes.

Guinn et al. (29) characterized the $TS$ for folding of 13 proteins by analyzing effects of urea, GuHCl and temperature on folding and unfolding rate constants. In protein folding and unfolding to $TS$, GuHCl primarily probes changes in accessible surface area ($\Delta ASA$) of amide groups, while urea probes both amide and hydrocarbon $\Delta ASA$ and activation heat capacities primarily probe hydrocarbon $\Delta ASA$. Amounts of amide and hydrocarbon ASA buried in $U \rightarrow TS$ and $TS \rightarrow F$ for folding all 13 proteins showed that $TS$ is advanced (because a majority ($50\%$–$90\%$) of the $\Delta ASA$ for $U \rightarrow F$ is buried in $U \rightarrow TS$), and that the $\Delta ASA$ of $U \rightarrow TS$ is preferentially amide. These findings are consistent with a folding mechanism in which all elements of unstable $2^\circ$ structure form early and intervening backbone regions then organize in $TS$ to nucleate coalescence of these $2^\circ$ structure elements post-$TS$. This research served as proof-of-principle of the use of solutes as probes of changes in surface area in biopolymer processes. Solute and salt effects on the thermodynamics and kinetics of RNA polymerase–promoter DNA open complex formation and stabilization were also used to identify and characterize large conformational changes and new interfaces in the key intermediates and in the DNA opening-closing $TS$ in transcription initiation (30–32).

Here, we extend this approach to characterize $TS$ and propose a mechanism for the structurally well-characterized $RO$ interaction. We measure and interpret effects of four solutes (the $RO$ destabilizer urea and $RO$-stabilizers glycerol, proline and glycine betaine (GB)) on the association and dissociation rate constants and equilibrium constant of the $RO$ interaction. We also report and interpret effects of the stabilizer tetraethylene glycol (TEG) on the $RO$ equilibrium constant, and analyze published effects of KCl concentration (23) on $RO$ rate and equilibrium constants. Interactions of these solutes with the different coarse-grained types of unified atoms of proteins have been investigated previously, and strengths of interaction ($\alpha$-values) with these unified atoms and with anionic phosphate oxygens of DNA have been determined (33–36). Approximately 80% of the predicted $\Delta ASA$ of $RO$ dissociation ($4600 \text{Å}^2$ of the total $5800 \text{Å}^2$) is from exposure of these types of unified atoms. Contributions of the remaining DNA atoms are known for urea (34) and are estimated for the other solutes using information for chemically similar protein atoms.

**Background on lac repressor–lac operator interaction**

$R$ is a homo-tetramer (dimer of dimers). Each subunit consists of a N-terminal 49-residue headpiece or DNA binding domain ($DBD$), a 13-residue flexible connector (hinge) region, a core regulatory domain and a C-terminal tetramerization domain (7,8,37). $DBDs$ of two subunits of $R$ bind one palindromic $O$ sequence. The structure of $RO$ in Figure 1 shows that these $DBD$ interact with the $O$ major groove and that the $\alpha$-helices formed by coupled folding of the hinge regions interact with the central minor groove of $O$, bending it away from the body of $R$. The entire assembly of $DBD$ and hinge helices on $O$ DNA binds to the core of $R$. This core–hinge helix-$DBD$ interaction facilitates transmission to the $DBD$ of the conformational change that occurs upon binding of alloolactose or another inducer to the core regions of the $R$ subunits, weakening $O$ binding (14–16,38).

Interpretation of heat capacity and entropy changes in binding of lac $R$ to $O$ and variants, as compared with binding to non-operator DNA, revealed extensive coupled folding to form the specific $RO$ complex but not the non-operator (RD) complex (12,13), and NMR studies identified the two hinge regions as the residues that fold to form $\alpha$-helices in specific but not non-specific binding (9–11). Analysis of the strong destabilizing effect of urea and strong stabilizing effect of GB on the equilibrium constant $K_{\text{obs}}$ for $RO$ complex formation provided additional evidence for coupled folding ($\alpha$-helix formation with burial of amide oxygens) and dehydration of DNA phosphates (39,40).

**Key functional groups in R-R and R-O interfaces formed in the RO complex.** Analysis of the structure of $RO$ reveals that aromatic, amide and cationic side chains of the $DBD$ and hinge regions, together with amide backbone groups and non-polar (hydrophobic) side chains, are centrally involved in forming the new $R$-$R$ and $R$-$O$ interfaces in $RO$ (7–11,41,42). In each half-site, two tyrosine residues (Y17, Y7) on the surface of the $DBD$ interact with DNA bases in the major groove of $O$. Positively-charged groups (e.g. K2, R22, N-terminal $-\text{NH}_3^+$) on each $DBD$ appear positioned to interact favorably with (i.e. within 6 Å of) negatively charged DNA phosphate oxygens, as is R118 of the core repressor (43).

The hinge regions (residues 50–62) are disordered in unbound $R$ but fold into a pair of interacting $\alpha$-helices in the process of $RO$ formation. These insert in the central minor groove of $O$, burying side-chain amide groups of N-terminal residues N50 and Q54 in the minor groove interface and the Q55 amide in an interface with the other hinge region (backbone amide of V52) and the core (R118). Non-polar side chains of residues A53, L56, A57 contact sugars of bases 10–13 in minor groove, and residues V52, A53, Q55, L56 form apolar contacts where hinge helices pack against each other. Interactions of the hinge helices with the minor groove bends $O$ DNA away from repressor by $\sim40^\circ$ (7–11,41,42). Analysis of the effect of glycine betaine on the $RO$ equilibrium constant revealed that anionic oper-
atator phosphate oxygens are significantly dehydrated in the specific RO interface (39).

The mechanism of the RO interaction can be deduced from information about which of these key functional groups are buried in forming TS from R and O (RO → TS), and which ones are buried subsequently in conversion of TS to RO (TS → RO).

MATERIALS AND METHODS

Chemicals and buffers

Urea, proline, glycerol, glycine betaine (GB) and tetraethylene glycol (tetraEG or TEG), purchased from Fluka and Sigma Aldrich were all of the highest available purity (>98%). Solutions of these solutes were prepared in the standard binding buffer (BB), which is 10 mM K2HPO4, 0.1 mM ethylenediaminetetraacetic acid (K2), 1 mM dithiothreitol (DTT), 100 μg/ml bovine serum albumin, 5% glycerol. KCl was added to obtain a final [K+] of 220 mM (for some tetraEG experiments) or 260 mM (for urea, proline, glycerol, GB and other tetraEG experiments), and pH was adjusted with HCl to pH 7.5 at assay temperature (25°C). The molar concentration of K+ was held constant in experiments as a function of solute concentration.

Lac repressor and lac operator DNA

Lac repressor tetramer was prepared, stored and diluted for use as previously described (40). Synthetic SymL DNA top strand (bases 1–36) and bottom strand (bases 1–40) were purchased in purified form from Integrated DNA Technologies (IDT) and annealed as described previously (13). The overhang was filled in by DNA polymerase Klenow fragment (44) using α P32 dATP and cold dCTP, followed by a 2 μM cold dATP chase. The resulting double-stranded 40 bp duplex was purified on a 2 ml Sephadex G25 column run in TE buffer, and DNA concentration was determined in a microcell using a Beckman DU640 spectrophotometer. The symmetric lac operator sequences are as follows.

O Sym36

5′-CACATATACC AATTGTGAGC GCTCACAATT CCAGAT

O Sym40

5′-GTGTATCTGG AATTGTGAGC GCTCACAATT GGTATATGTG

Nitrocellulose filter binding assays

Filter binding assays were used to determine the amount of operator DNA retained on a nitrocellulose filter membrane in a complex with lac repressor protein (30,32). All thermodynamic and kinetic assays were performed in BB. For all assays, repressor was in sufficient excess over operator so only 1:1 RO complexes were formed. The fraction of operator DNA in the form of RO complexes (θ) was determined from the radioactive counts per minute (cpm) of 32P-labeled operator DNA retained on the filter using the relationship

θ = \frac{cpm - cpm_{bgd}}{E (cpm_{tot} - cpm_{bgd})} \tag{1}

where cpm_{bgd} is the cpm of 32P-labeled operator DNA retained on the filter in the absence of lac R, cpm_{tot} is the total cpm of 32P-labeled operator DNA filtered in each assay and E is the filter efficiency (the fraction of RO complexes in solution that are retained on the filter membrane), determined as previously described (30). Average filter efficiency was 0.61 ± 0.09. For the range of solute concentrations studied here, we observe no systematic variation of filter efficiency.
with solute concentration within the ±15% uncertainty in these determinations.

To test if the low concentration of RO₂ could contribute significantly to r-values, we analyzed TEG equilibrium titration data (Supplementary Figure S1), where the |R|:|O| total concentration ratio (≈2:1) is smaller than in our other thermodynamic and kinetic studies. These titration data were fitted including and excluding 1:2 complexes and r-values were compared. The change in r-value from inclusion of 1:2 complexes is <5%, which is negligible given the experimental uncertainty. For all other conditions, the |R|:|O| total concentration ratio is larger, and the effect will be even smaller.

Binding activity assays were performed to convert total repressor concentrations to active concentrations (44); 40–45% of repressor tetramers were active in binding operator.

Equilibrium titrations with solutes. For solute equilibrium titrations, Lac R and Lac O concentrations were selected to give initial fractional occupancies θ (Equation 1) in the range 0.22 < θ < 0.38 for stabilizing solutes, and θ = 0.79 – 0.83 for the destabilizing solute urea. The Lac O concentration ([O]_{total}) was 20 pM except for experiments with tetraEG, where 10 pM was used. Total concentrations of active Lac R ([R_{site,active}]_{total}) expressed on a dimer (operator binding site) scale, were usually in the range 68–85 pM, except for urea (1.7 nM). Series of samples with the same total repressor and operator concentrations and increasing solute concentration in the range 0 to 1 molar (1.2 molar for tetraEG) were prepared, incubated for 1–3 h and filtered. Values of θ as a function of solute concentration (see Supplementary Figure S1A) were determined (Equation 1) and used to calculate equilibrium constants K_D as a function of solute molarity from Equation 2:

\[ K_D = \left( 1 - \theta \right) \left( \frac{[R_{site,active}]}{\theta} - [O]_{total} \right)_{eq} \] (2)

Values of lnK_D were fitted linearly with fixed intercepts, as functions of solute molarity. The intercepts (lnK_D^0; Supplementary Table S1) obtained from data in the absence of solutes, were used to obtain normalized plots of lnK_D/lnK_D^0 versus solute concentration (Figure 2A). Propagated uncertainties were calculated for each data point and representative uncertainties are shown in figures.

Decay to equilibrium kinetic studies. Kinetic experiments were performed as a function of solute concentration by mixing 10–20 pM 32P-labeled operator DNA with excess repressor ([R_{site,active}]_{total}) in the range 80–406 pM for all solutes. Repressor concentrations were chosen so that the binding reaction decayed to equilibrium (θ_{eq} ≤ 0.9). Single-exponential decay-to-equilibrium kinetics (examples of which are shown in Supplementary Figure S1B) were analyzed using background corrected cpm (i.e. cpm – cpm_{kgd}) and the Equations (3,4).

\[ \text{cpm}_i = \text{cpm}_{eq} \left( 1 - e^{-k_{obs} t} \right) \] (3)

where,

\[ k_{obs} = k_a \left( [R_{site,active}]_{eq} + [O]_{eq} \right) + k_d \approx k_a \left( 1 + \frac{[R_{site,active}]}{K_D} \right) \] (4)

Here \([R_{site,active}]_{eq} = [R_{site,active}]_{total} - \theta_{eq}[O]_{total}\) and \([O]_{eq} = (1 - \theta_{eq})[O]_{total}\), so that \([R_{site,active}]_{eq} + [O]_{eq} = [R_{site,active}]_{total} + [O]_{total} (1 - 2\theta_{eq})\). Since \([O]_{total} <\) \([R_{site,active}]_{total}\), the term \([O]_{total} (1 - 2\theta_{eq})\) can be neglected. Dissociation rate constants k_d were obtained from decay to equilibrium rate constants k_{obs} using the appropriate \([R_{site,active}]_{total}\) and the K_D obtained from the corresponding solute titration, and association rate constants k_a were obtained from k_d = k_a/K_D. Values of k_d and k_a obtained from tetraEG experiments at low salt concentration (220 mM), were corrected for salt effects on the two rate constants using experimentally determined values of S_{K_i}(23). Normalized plots of lnk_d and lnk_a as functions of solute molarity (Figure 2B and C) were obtained by using lnk_d^0 and lnk_a^0 values obtained from experiments in absence of solutes (Supplementary Table S1). Contribution of the term \([O]_{total} (1 - 2\theta_{eq})\) in comparison to \([R_{site,active}]_{total}\) was estimated from experimental concentrations and values of θ and found to be on average 4%, which is negligible. Propagated uncertainties (σ) calculated for each data point using Equation 5, are shown in figures.

\[ \sigma_{rK}^2 = \left( \frac{\partial f}{\partial x} \right)^2 \sigma_x^2 + \left( \frac{\partial f}{\partial y} \right)^2 \sigma_y^2 + \left( \frac{\partial f}{\partial z} \right)^2 \sigma_z^2 \] (5)

Equation (5) shows the generic method for calculating propagated uncertainty (σ_r) for any given function, f(x, y, z) using the product of partial derivatives of the function (f) with respect to the independent variables (x, y, z) and the uncertainties associated with each variable (σ_x, σ_y, σ_z).

RESULTS

Solute and salt effects on the RO dissociation equilibrium constant (K_D). Effects of probe solutes on the RO dissociation equilibrium constant K_D are shown in Figure 2A (and Supplementary Figure S2), in which the logarithm of K_D, obtained from solute titrations, is plotted versus molal solute concentration (m_3). These plots are linear over the range studied (0.1 to ~1.5 m). Effects of solutes are characterized by slopes dlnK_D/dm_3, here called r_K-values (Figure 2A). These r_K-values, reported with uncertainties in Supplementary Table S2, range from 2.7 for urea to ~1.1 for glycerol, ~1.7 for proline and ~1.8 for GB. Data for a fourth stabilizing solute (tetraethylene glycol, TEG), obtained at a somewhat lower KCl concentration, are reported in Supplementary Figure S2. The TEG r_K-value (~2.0) is similar to that of GB. These r_K-values are proportional to thermodynamic m-values (m-value = dΔG^0/dm_3 = −RTdlnK_D/dm_3 = −RT(r_K-value)) = Δμ_{23}, where Δμ_{23} is the difference in chemical potential-derivatives μ_{23} = δμ_{23}/dm_3 at constant pressure, temperature and m_2, between the RO complex and reactants (28). Values of Δμ_{23}, and hence r_K-values, are interpreted as preferential interactions of the solute with the functional groups exposed in the process (28–32,45) (see Analysis below and Supplementary Equations S4 and 5).

Figure 2A shows that addition of urea favors RO dissociation, increasing K_D by 15-fold at 1 molal urea. This is expected because urea is found to interact favorably with all protein and nucleic acid functional groups except...
cations and under different conditions (28,39,40). Unlike urea, aliphatic (sp³) C (36). Unfavorable interactions of these so-

alicylic N (Supplementary Table S3) (33,34,46). Other sol-

tuates with oxygens and also (except for TEG) with hydrocar-

bon groups exposed in RO must be dominant.

These findings indicate that the entire effect of the RO-stabilizing solutes is on the conversion of the high free-

energy TS to RO, while approximately half the destabilizing effect of urea is on conversion of TS to RO. Consistent with this, Figure 2C reveals that association kinetic $k_a$-values for the RO-stabilizing solutes are zero within uncertainty, while the urea $r_K$-value is about half as large as the urea $r_K$-value.

Literature values of the rate constants $k_a$ and $k_d$ as func-
tions of [KCl] for the RO interaction involving 40 bp O are shown in Figure 3 as log-log plots (23). At 0.15 M salt, the association rate constant ($\sim 10^9$ M⁻¹ s⁻¹ or larger) and activation energy of association ($\sim 4$ kcal (23)) are consistent with a diffusion-limited reaction. An increase in [KCl] destabilizes RO by reducing the association rate constant greatly ($Sk_d = -3.5$) and increasing the dissociation rate constant slightly ($Sk_d = 0.7$) (Supplementary Table S4).

At 0.26 M KCl, the salt concentration investigated here, we find $k_a = 4 \times 10^7$ M⁻¹ s⁻¹ in the absence of perturbing solutes, two orders of magnitude less than the diffusion limit
DISCUSSION

Analysis of thermodynamic $r_K$-values for RO dissociation using ∆ASA information and solute α-values

Thermodynamic ($K_D$) $r_K$-values quantifying effects of solutes on dissociation of RO are determined by preferential interactions of these solutes with the different types of unified atoms on the R and O surfaces exposed in dissociation. Using the RO structure (1EFA (7)) and models for free R and O (models described in Supplementary Methods), the amount and composition of R and O surfaces exposed in RO dissociation were estimated. The overall change in ASA (∆ASA) is predicted to be approximately 5800 Å² (Figure 4A). The contribution to ∆ASA from R in dissociation (4100 Å²) greatly exceeds that from O (1700 Å²). Exposure of 3000 Å² of aliphatic (sp³) and aromatic (sp²) C (70% from R, 30% from O) accounts for half of the overall ∆ASA of RO dissociation. The predicted contribution of anionic DNA phosphate oxygens to the ∆ASA is 467 Å², one-fourth of the DNA contribution to ∆ASA. Other significant contributions to the ∆ASA come from Ramide O, N and cationic N atoms and O sugar sp³ C.

Predicted contributions to the observed solute $r_K$-values from solute interactions with all types of R and O atoms exposed in dissociation of RO are shown in the bar graph of Figure 4B. These are calculated from the ASA information of Figure 4A and the interaction potentials (α-values) of Supplementary Table S3. These α-values quantify solute interactions with a unit area of the seven major types of unified atoms of protein (aliphatic and aromatic C; amide and cationic N; amide, hydroxyl and carboxylate O) and with anionic phosphate O of DNA (28,33–36). Supplementary Table S3 also lists α-values for interaction of urea with other types of DNA atoms. α-Values for interactions of other solutes with these types of DNA atoms are assumed to be the same as those for chemically-similar types of protein atoms. Inspection of urea α-values shows this is a reasonable first approximation.

Figure 4 reveals that while aliphatic carbon (C) makes the largest contribution to ∆ASA (Figure 4A), exposure of DNA anionic phosphate oxygens in RO dissociation is predicted to make the largest contribution to the overall $r_K$-value for the three solutes (TEG, GB and proline) that most stabilize RO (Figure 4B), because of the large positive α-values and consequent strong unfavorable interactions of these solutes with phosphate oxygen. For GB and TEG, the two solutes that are most highly excluded from anionic phosphate oxygens, predicted contributions to $r_K$-values from anionic phosphate oxygen ∆ASA appear too large in magnitude, causing magnitudes of overall predicted $r_K$-values to exceed observed $r_K$-values by 60 to 70% (Supplementary Table S5). A likely interpretation of these discrepancies is that anionic phosphate oxygens are not completely dehydrated when they are incorporated into the RO interface, retaining ~1/3 of their original water of hydration. Hence the amount of water that is taken up upon exposure of these phosphate oxygens in RO dissociation is only 2/3 of that calculated from the structure. The thermodynamically-detected residual hydration of phosphates in the interface is most simply parameterized for analysis by reducing the anionic phosphate oxygen ∆ASA for RO dissociation by approximately one-third, from 467 Å² to 302 ± 32 Å².

Structural evidence for residual water in the interface of a complex between one DBD and a half operator site was obtained previously (41), though more recent structures of the RO complex formed by two DBD do not resolve any bound...
NLPB prediction of $SK_D$ quantifying the KCl effect on $K_D$ of RO dissociation

Coulombic free energies $G_{coul}^\circ$ of R, O and the RO complex at 25°C were obtained as functions of 1:1 salt (KCl) concentration from the non-linear Poisson-Boltzmann equation as described in Supplementary Data. These energies $G_{coul}^\circ$ are plotted versus ln[KCl] in Supplementary Figure S4A. The corresponding Coulombic free energy changes in RO dissociation, $\Delta G_{coul}^\circ = G_{coul}^\circ R + G_{coul}^\circ O - G_{coul}^\circ RO$ are plotted versus ln[KCl] in Supplementary Figure S4B. Individual values of free energies $G_{coul}^\circ$ decrease non-linearly with increasing ln[KCl], but $\Delta G_{coul}^\circ$ decreases linearly with ln[KCl] in the range of interest. $\frac{d\Delta G_{coul}^\circ}{d\ln[KCl]} = -RT SK_D$ yields a parameter-free predicted value of $SK_D = 6.5$, which agrees well with the experimental $SK_D = 5.6 \pm 0.4$ (Figure 3; Supplementary Figure S3C and Table S4; ref (23)).

Qualitative interpretation of effects of KCl and solutes on rate constants

At 0.26 M KCl, the second order association rate constant $k_a$ for RO formation is approximately two orders of magnitude less than the diffusion limit. From Figure 2B and C, the effect of urea concentration on the dissociation equilibrium constant $K_D$ is distributed roughly equally between the rate constants $k_a$ and $k_d$. From Figure 3, most of the large effect of KCl concentration on $K_D$ is on $k_a$; $k_d$ is relatively independent of KCl concentration. Taken together, these findings indicate that at 0.26 M KCl the rate determining step in RO formation is subsequent to the initial diffusion–collision interaction of R and O. This initial binding step is deduced to be in rapid equilibrium on the time scale of subsequent conformation changes, and the high-free-energy transition state for the rate-determining conformational change is somewhere in mid-mechanism of RO formation.

Effects of the three RO-stabilizing solutes on the kinetics are largely on the dissociation rate constant $k_d$. Comparison of Figure 2B and C with Figure 2A shows that these solutes have very similar effects on $k_a$ and $k_d$ and have negligibly small effects on $k_a$. The most likely interpretation of the small-magnitude $k_d$-r-values for RO-stabilizing solutes is that (i) little if any dehydration of anionic phosphate oxygens occurs in forming TS from R and O, and (ii) there is near-complete compensation between TS-stabilizing and TS-desstabilizing interactions of these RO-stabilizing solutes. If $k_a$ were closer to the diffusion limit and if KCl and urea also had small effects on $k_a$ and large effects on $k_d$, an alternative interpretation would be that TS is very early, involving few interactions between R and O charges or functional groups and no dehydration of interfaces. But the KCl and urea results, and the fact that $k_a$ is much less than the diffusion limit, show that this interpretation is incorrect. The only atoms that these RO-stabilizing solutes interact favorably with are aromatic carbon (sp2 C) and amide nitrogen (N). We therefore assume that anionic phosphate oxygens remain hydrated in TS, and propose that the most likely TS candidates are ones that preferentially bury sp2 C and amide N, but not amide O.
Evidence for the nature of the high-free-energy transition state from comparison of predicted and observed solute and salt effects on rate constants $k_a$, $k_d$

Structural models of four possible intermediates in $RO$ complex formation from free $R$ and $O$ (labeled $I_1$ to $I_4$; see Figure 5) were generated starting from published coordinates of non-specific ($RD$) and specific ($RO$) complexes (see Supplementary Methods). These models correspond to steps in the proposed mechanism (see ‘Introduction’ section), deduced from structural information. Intermediate $I_1$ is based on the structure of the non-specific ($RD$) complex (10) (using the $O$ DNA sequence), which we assume is an appropriate model of the least advanced intermediate formed with $O$ DNA. Modeled intermediate $I_2$ is more advanced than $I_1$, having the specific interactions of the two $R$ DBDs with the major grooves of $O$ that are present in the stable $RO$ complex. In addition to the $DBD$-major groove interactions of $I_2$, intermediate $I_3$ has nucleated $\alpha$-helix formation at the N-termini of the two hinge regions, contacting and bending the central minor groove of $O$. The hinge helices are completely formed but not yet inserted in the bent $O$ minor groove in more advanced intermediate $I_4$, in which the $DBD$-hinge helix- $O$ DNA module is also not yet docked on the core of repressor to form the stable $RO$ complex.

Solute $r_K$-values for formation of each intermediate from free $R$ and $O$ and from $RO$ are predicted from the $\alpha$-values of Supplementary Table S3 and the results of ASA analysis of these models of possible intermediates (Supplementary Table S6). Comparison of these predictions (Supplementary Figure S5) with observed association and dissociation kinetic $r_k$-values and with the $r_K$-value of $RO$ reveals when DNA phosphates lose their outer layers of hydration in $RO$ formation and which, if any, of these proposed intermediates might be a good model for the high-free-energy transition state that determines $k_a$ and $k_d$.

When in $RO$ formation do DNA phosphates lose their outer layers of hydration? Dehydration of the outer layers of hydration of DNA anionic phosphate oxygens (from three layers of water to one layer) must occur post-$TS$ in $RO$ formation. Conversely, hydration of anionic phosphate oxygens from one layer to three layers of water occurs pre-$TS$ in $RO$ dissociation. If substantial dehydration of anionic phosphate oxygen surface occurred pre-$TS$ in association, $RO$-stabilizing solutes would of necessity also stabilize $TS$ and therefore increase $k_d$. For $RO$-stabilizing solutes to have the observed minimal effect on $k_a$, the anionic phosphate oxygens that are involved in coulombic interactions with cationic groups of $R$ must remain hydrated in associa-
tion to TS. Loss of two layers of hydration water of anionic phosphate oxygens in converting TS to RO makes a major (dominant) contribution to the $k_3$ $r_3$-values, and therefore is a large part of the explanation for why $k_3$ $r_3$-values and $K_3$ $r_3$-values are so similar for these RO-stabilizing solutes. Conversion of TS to RO therefore tightens the interfaces between $R$ domains and $O$ phosphates, squeezing out two of the three layers of water of hydration of anionic phosphate oxygens.

Nucleation of hinge helix formation in I3 makes it the most plausible candidate for TS. For the range of possible TS models considered here (I species I1 to I4 in Figure 5), Supplementary Figure S6 predicts contributions to the association kinetic $r_k$-value from the various atom-types buried in forming these TS candidates from $R$ and $O$ and compares predicted and observed association kinetic $r_k$-values. In all cases, anionic phosphate oxygens are assumed to remain completely hydrated in TS (0% dehydration). For comparison, a ‘late I’ model for TS is also considered, with the structure of RO but with 0% dehydration of anionic phosphate oxygens. The circled results in Supplementary Figure S6, representing the best agreement between predicted and observed association kinetic $r_k$-values, indicate that the most likely TS structure is more advanced than I1 but less advanced than I4, and could be either I2 or I3. The urea data are most definitive, favoring I3 as TS though not excluding I2. Results for the RO-stabilizing solutes in Supplementary Figure S6 show a small but significant preference for I1 or I2 as TS over I3 and I4.

Nucleation of hinge helix formation in I3 makes it an attractive candidate for TS in RO formation. Nucleation steps generally are highly unfavorable. Nucleation of tertiary structure formation appears to be the high free energy TS in protein folding (29). Moreover, complications between favorable and unfavorable interactions of RO-stabilizing solutes with the ΔASA of forming I3 (also I2) from R and O, shown in Figure 6 and quantified in Supplementary Figure S6, result in small-magnitude predicted association kinetic $r_k$-values for these RO-stabilizing solutes, consistent with experiment.

Figure 6 shows our proposal that nucleation of hinge helix formation in the conversion of I2 to I3 buries side-chain amide N of three asparagines and glutamine residues (N50 (dark pink), Q54 and Q55 (light pink)) at the N-terminus of each hinge region, without much burial of amide O. Indeed, half of the amide N ASA buried in RO formation occurs in forming I3, our TS candidate, while only a third of total amide O burial occurs in forming TS (see Supplementary Table S6). Earlier in formation of this TS model from R and O, aromatic C atoms of tyrosines Y7 and Y17 (cyan) of each DBD are buried in the major grooves of O DNA (Figure 6), accounting for all the R aromatic C buried in the stable RO complex.

Post-TS, the favorable propagation steps of hinge helix formation occur. We propose that the pair of interacting hinge helices insert into the central minor groove of the O DNA, bending the DNA away from the protein and forming intermediate I4. In the final step, the assembly of DBD, O DNA and hinge-helices in I4 docks on the C-terminal core of R, tightening and partially dehydrating the R-O-in-terface to form the final RO complex. Figure 5 places the various I-species and the proposed TS on an activation free energy diagram and summarizes this non-covalent self-assembly (nucleation-propagation) mechanism of RO formation.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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