Interaction of the ocr gene 0.3 protein of bacteriophage T7 with EcoK1 restriction/modification enzyme

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ABSTRACT

The ocr protein, the product of gene 0.3 of bacteriophage T7, is a structural mimic of the phosphate backbone of B-form DNA. In total it mimics 22 phosphate groups over ~24 bp of DNA. This mimicry allows it to block DNA binding by type I DNA restriction enzymes and to inhibit these enzymes. We have determined that multiple ocr dimers can bind stoichiometrically to the archetypal type I enzyme, EcoK1. One dimer binds to the core methyltransferase and two to the complete bifunctional restriction and modification enzyme. Ocr can also bind to the component subunits of EcoK1. Binding affinity to the methyltransferase core is extremely strong with a large favourable enthalpy change and an unfavourable entropy change. This strong interaction prevents the dissociation of the methyltransferase which occurs upon dilution of the enzyme. This stabilisation arises because the interaction appears to involve virtually the entire surface area of ocr and leads to the enzyme completely wrapping around ocr.

INTRODUCTION

It is apparent from the atomic structure of the ocr protein produced by gene 0.3 of bacteriophage T7 that it is a mimic of the general structure of ~24 bp of B-DNA containing a central kink of 34° (Fig. 1) (1). The surface of the protein is decorated with negatively charged groups which are superimposable upon the phosphate backbone of the DNA and the kink angle is similar to that induced by type I DNA restriction enzymes in their DNA targets. This similarity between ocr and the DNA target of these restriction enzymes occurs upon binding of the enzyme. This stabilisation arises because the interaction appears to involve virtually the entire surface area of ocr and leads to the enzyme completely wrapping around ocr.

MATERIALS AND METHODS

Proteins were prepared as described previously (7,8,16,18). Mutant forms of ocr are referred to as, for example, ocr(W94C) for the substitution of Trp94 by Cys and ocr(Cys) refers to any of the mutant forms containing a cysteine substitution. Note that wild-type ocr contains no cysteine. These six mutant proteins have previously been shown to react with a range of reagents (7) and modification (R/M) systems comprising a restriction endonuclease activity which cuts the viral DNA and a modification methyltransferase activity which modifies host DNA (3). The 27 kDa dimeric ocr protein (6–8) knocks out a major subgroup of R/M enzymes, the type I R/M systems. These R/M systems contain both restriction and modification functions in a single enzyme (9–11). The antirestriction activity of ocr is not dependent on the DNA sequence recognised by the type I R/M enzyme. Each of the approximately 20 known type I R/M systems, covering four separate families in euarchea and archaea, recognise a unique DNA target sequence (12). However, ocr will inactivate representatives of all four families irrespective of their DNA target sequence (1,13–15). This is achieved by very strong, stoichiometric binding of ocr to the R/M enzyme and the competitive displacement of DNA from the enzyme (7).

In this paper, we examine the nature of interactions between ocr and the archetypal type I R/M system, EcoK1. This R/M enzyme exists in two forms, R.EcoK1 (440 kDa) and M.EcoK1 (170 kDa) (16,17). R.EcoK1 is a bifunctional restriction endonuclease and modification methyltransferase containing two restriction (R) subunits, two modification (M) subunits and one DNA specificity (S) subunit whilst M.EcoK1 is a monofunctional modification methyltransferase containing two M and one S subunits.

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in all experiments was 20 mM Tris, 20 mM NH₄Cl, 6 mM MgCl₂ buffer, pH 8 at 25°C. Unless otherwise stated, all proteins were desalted using a PD-10 G-25 column (Amersham Pharmacia) equilibrated in buffer prior to use. Data were analysed using Grafit (Erithacus Software) or Origin (Microcal).

Isothermal titration calorimetry (ITC)
Both ocr and M.EcoKI were dialysed overnight into the above buffer. A 64.5 μM stock solution of ocr protein was titrated into 1.7 ml of 4.8 μM M.EcoKI in the VP-ITC cell. Aliquots of 10 μl of ocr were injected after an initial injection of 1 μl. The heat of dilution was obtained by injection of ocr into buffer alone and buffer directly into buffer. The values were subtracted from the ITC titration data.

Tryptophan fluorescence
An aliquot of 100 μl of 0.3 μM M.EcoKI was placed in a 3 mm path length cuvette and titrated with a 22 μM solution of ocr. Samples were excited at 295 nm and the emission spectra recorded between 300 and 500 nm. Excitation and emission slit widths of 5 and 10 nm, respectively, were used. The fluorescence of the buffer was subtracted from the emission spectra. Spectra were corrected for dilution and inner filter effects. Similar procedures were applied to the titrations using other components of the EcoKI R/M enzyme.

Experiments with 5-((2-iodoacetyl)amino)ethylamino) naphthalene-1-sulfonic acid (1,5-IAEDANS)
Aliquots of 100 μl of 50 μM ocr(Cys) proteins were incubated overnight at 4°C in the dark with a 50-fold molar excess of 1,5-IAEDANS (Molecular Probes). The incubation was stopped by the addition of 100 mM 2-mercaptoethanol and unreacted probe removed by gel filtration on a PD-10 column. The concentration of AEDANS bound to the proteins was calculated from the absorption spectra at 280 nm after subtracting the AEDANS absorbance at this wavelength (the molar extinction coefficient of 1,5-IAEDANS at 280 nm is 1060 M⁻¹ cm⁻¹).

To determine the stoichiometry of binding between ocr(Cys)±AEDANS and M.EcoKI, 100 μl of 1.0 μM ocr(Cys)±AEDANS solutions were titrated with increasing amounts of M.EcoKI. Excitation was at 330 nm and emission intensity monitored at 480 nm with slit widths of 5.0 and 10 nm, respectively. The cuvette path lengths were 3 mm.

The fluorescence intensity of ocr(Cys)±AEDANS proteins was quenched with acrylamide in the absence or presence of M.EcoKI. Solutions used were either 100 μl of 1.0 μM ocr(Cys)±AEDANS or 100 μl of 1.0 μM ocr(Cys)±AEDANS plus 2.0 μM M.EcoKI. The excess of M.EcoKI ensures the formation of a 1:1 complex. The samples were titrated with acrylamide to a final concentration of 0.5 M.

Labelling with 5-iodoacetamidofluorescein (5-IAF)
Ocr(N43C) protein was labelled with 5-IAF (Molecular Probes) overnight at 4°C in the dark using the same procedures as for 1,5-IAEDANS. The unbound 5-IAF was removed using a PD-10 column. The concentration of the label bound to the protein was calculated from the absorbance at 492 nm using an extinction coefficient of 71 000 M⁻¹ cm⁻¹. The protein concentration was determined from the absorbance at 280 nm after subtracting the absorbance of IAF at this wavelength.

An aliquot of 1 ml of 0.1 μM ocr(N43C)–AF was placed in a 4 × 10 mm cuvette (excitation path 10 mm) and titrated with R.EcoKI or M.EcoKI. The slit width used for excitation and emission was 5.4 nm. Labelled protein was excited at 492 nm, the emission spectra were integrated between 510 and 570 nm and the values obtained fitted to a single ligand binding site equation to derive a dissociation constant, K_d.

Labelling with N-(1-pyrene)maleimide (PM)
A stock solution of PM (Molecular Probes) was made in DMSO. Ocr(Cys) proteins were desalted into 100 mM sodium phosphate buffer, pH 7.0. An aliquot of 100 μl of 50 μM
protein was incubated with a 50-fold excess of PM overnight at 4°C, in the dark. The labelling reaction was stopped by adding 2-mercaptoethanol to 100 mM final concentration. Unreacted fluorophore was removed using the PD-10 G-25 gel-filtration column equilibrated in the phosphate buffer (20). Labelling of the ocr(N4C) was performed in 100 mM sodium phosphate, 7 mM 2-mercaptoethanol, pH 7.0, buffer as this protein contains a proportion of disulphide bonds. The degree of labelling was determined spectrometrically using a value of 40,000 M⁻¹ cm⁻¹ for PM at 345 nm (21). The protein concentration was calculated from the absorbance at 280 nm after subtracting the absorbance of PM. The extinction coefficient of PM at this wavelength is 25,000 M⁻¹ cm⁻¹.

To determine the stoichiometry of binding between ocr(Cys)±PM and M.EcoK1, 100 μl of 1.0 μM ocr(Cys)±PM solution in the Tris buffer was titrated with increasing amounts of M.EcoK1. Excitation was at 342 nm and emission intensity monitored at 379 nm with slit widths of 5.0 and 10 nm, respectively. The cuvette path lengths were 3 mm.

HPLC gel filtration

Gel filtration of M.EcoK1, ocr and 1:1 complexes of ocr and M.EcoK1 were performed under conditions and with buffer solutions as described (18). The chromatography column was calibrated with a range of proteins at several different protein concentrations down to 10 nM. No dependence of elution time upon concentration was observed for the standards. A concentrated stock of each protein or protein mixture was diluted to the appropriate concentration immediately prior to injection onto the column.

RESULTS

Isothermal titration calorimetry data showing 1:1 binding to M.EcoK1

Figure 2 shows a typical ITC measurement of the heat of binding of ocr to M.EcoK1. The reaction was exothermic and each addition of ocr lowered the amount of heat released until M.EcoK1 was saturated. After integration of each injection peak and subtraction of the heats of dilution, the values of the heat of interaction normalised for ocr concentration were plotted versus the molar ratio of ocr to M.EcoK1. These experimental data were fitted to a theoretical titration curve to determine the enthalpy of reaction, ΔH, and the number of binding sites, n. The interaction between ocr and M.EcoK1 is very strong and exhibits essentially stoichiometric binding in the concentration range used in the ITC experiment. Therefore, the binding affinity cannot be determined. The stoichiometry of binding was 0.95 ± 0.05 ocr dimer molecules per M.EcoK1 molecule. The reaction was exothermic under the conditions used with a ΔH of −76.3 ± 0.8 kJ mol⁻¹.

Stoichiometry of binding of labelled mutants to M.EcoK1 observed by extrinsic fluorescence

Upon titration of the labelled ocr protein with M.EcoK1, the fluorescence intensity of the fluorescent label changed. After correction for dilution, a plot of intensity against the molar ratio [M.EcoK1]/[labelled ocr] showed stoichiometric binding for each different labelled ocr protein (Fig. 3 and Table 1). The fluorescence emission intensity increased for all of the labelled proteins except for ocr(D62C) when labelled with pyrene, which showed a decrease. While all of the titrations showed an end-point at a ratio of one ocr dimer per M.EcoK1, several of the titrations with AEDANS-labelled ocr showed an extra end-point corresponding to two ocr dimers per M.EcoK1. The different titration behaviour of the fluorescent labels depending upon location on the surface of the ocr mutant protein may indicate interaction with different parts of M.EcoK1. Alternatively, since the ocr(Cys) mutant proteins contain two labelling sites, the two stoichiometries may reflect a difference between ocr molecules with one or two extrinsic labels.
Stoichiometry of binding to M.EcoKI using intrinsic fluorescence

Wild-type ocr contains only one tryptophan residue at position 94 in each monomer (22). Fluorescence spectroscopy and X-ray crystallography indicate that Trp94 is located on the surface of ocr (1,7). EcoKI contains multiple tryptophan residues and displays an emission maximum around 340 nm (16). The addition of ocr to a solution of M.EcoKI causes an increase in total emission intensity from tryptophan. However, this increase is less than expected by merely adding the intensity of M.EcoKI and ocr together. This quenching effect can be attributed to the interaction between the two proteins affecting the environment of some of the tryptophan side chains and reducing their contribution to the total emission. A graph of the difference between the sum of the individual intensities of ocr and M.EcoKI at 340 nm and the observed fluorescence intensity of the ocr±M.EcoKI mixture plotted against the molar ratio [ocr]/[M.EcoKI] once again shows saturation behaviour (Fig. 4). The stoichiometry of the ocr±M.EcoKI complex by this method is 1.5 ocr dimers per M.EcoKI. This non-integral ratio is unexpected and may be a reflection of the fact that M.EcoKI can dissociate into an M1S1 dimer and an M subunit with a dissociation constant of 97 nM (see below). Both of these fragments of M.EcoKI can bind ocr, as described later.

Table 1. Stoichiometry of the ocr–EcoKI interaction

<table>
<thead>
<tr>
<th>EcoKI component</th>
<th>Ocr protein</th>
<th>Method</th>
<th>Observed endpoint stoichiometries</th>
</tr>
</thead>
<tbody>
<tr>
<td>M.EcoKI</td>
<td>Wild type</td>
<td>ITC</td>
<td>0.95 ± 0.05</td>
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<td></td>
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<td>R.EcoKI</td>
<td>Wild type</td>
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<tr>
<td>R subunit</td>
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Stoichiometry of binding to M.EcoKI using intrinsic fluorescence

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Binding affinity of ocr and M.EcoKI and R.EcoKI

The measurements of binding of ocr to M.EcoKI described above were performed at concentrations considerably greater than the dissociation constant for binding of ocr to M.EcoKI. This constant was estimated as being at least 100 pM (7) but the fluorescence anisotropy method used was at the limits of its sensitivity. The strong extinction coefficient and high quantum yield of fluorescein make it a suitable probe for determining the dissociation constant for binding of ocr to EcoKI in the low concentration regime required to examine this tight binding inhibitor although, as usual, one has to assume that the probe does not alter the interaction between the two proteins. Ocr(N43C) and ocr(S68C) were labelled with fluorescein and titrated with M.EcoKI or R.EcoKI. The fluorescence emission intensity of the fluorescein label decreased as EcoKI was added (Fig. 5). This decrease was fitted to a ligand binding curve to give dissociation constants. The titration of labelled ocr(N43C) gave dissociation constants of 39 ± 3 and 53 ± 4 pM for M.EcoKI and R.EcoKI, respectively. Labelled ocr(S68C) protein gave dissociation constants of 79 ± 12 and 99 ± 24 pM for M.EcoKI and R.EcoKI, respectively. The different locations of the fluorescein label and differences between different labelling reactions may account for the different dissociation constants observed. These dissociation constants are 20- to 100-fold lower than for the binding of EcoKI to DNA (23,24), indicating a strong preference for EcoKI to bind to ocr rather than DNA, and they may still be overestimates of the true dissociation constants as the extrinsic label may interfere with the binding interaction. The 39 pM dissociation constant obtained from the M.EcoKI titration experiment was used to calculate the Gibbs free energy (ΔG) from the equation ΔG = RTlnKd, where R is the gas constant and T is the temperature in Kelvin. The value obtained for ΔG was −59.4 kJ mol⁻¹. The Gibbs free energy is also expressed as ΔG = ΔH − TΔS and using the value of ΔH of −76.3 kJ mol⁻¹ from the calorimetry
Enthalpy contributes favourably to the Gibbs free energy of association constant estimated to lie between 15 and 80 nM (18).

Enthalpy contributes favourably to the Gibbs free energy, while the contribution of entropy is unfavourable at 25°C. The association of native ocr with M.EcoKI was further investigated using gel filtration chromatography. It was previously observed that M.EcoKI dissociates at low protein concentration during the gel filtration process with a dissociation constant estimated to lie between 15 and 80 nM (18).

Figure 6 shows the change in elution volume of M.EcoKI dissociated at low protein concentration using tryptophan fluorescence for detection. This is more sensitive than the absorption method used previously and allows an upper limit for the $K_D$ of 97 ± 9 nM to be determined. Ocr, in common with M.EcoKI, has unusual elution behaviour on the gel filtration column, appearing to be considerably larger than a dimer. However, equilibrium analytical ultracentrifugation (7) and dynamic light scattering (8) showed that ocr is a dimer in this concentration range so the unusual gel filtration behaviour must be due to the elongated shape of the ocr dimer rather than some aggregation to form larger species. The behaviour of a 1:1 mixture of M.EcoKI and ocr dimer during gel filtration is also shown in Figure 6. It is apparent that the ocr–M.EcoKI complex does not dissociate during the gel filtration process at any detectable protein concentration. Thus ocr stabilises the trimeric structure of M.EcoKI. This experiment further indicates that once ocr is bound, it remains bound to M.EcoKI for periods significantly longer than the 15 min duration of the gel filtration experiments. Therefore, the association between ocr and M.EcoKI must be very rapid given the high binding affinity and slow dissociation rate for the complex. This behaviour has been observed for other strong protein–protein associations (see for example 25,26).

Ocr–M.EcoKI interface

The titration of fluorescently labelled ocr mutant proteins showed a change in fluorescence intensity upon interacting with the M.EcoKI irrespective of the location of the labelling site. Furthermore, the titration of the AEDANS-labelled ocr also showed that the emission spectrum of AEDANS shifted to a lower wavelength when the ocr was saturated with M.EcoKI (data not shown). A shift to lower wavelengths would imply a less aqueous environment for the label. This change in solvation could arise if all six labelling locations are at the binding interface between ocr and M.EcoKI or if these sites change conformation when the M.EcoKI binds elsewhere on ocr. Similar changes were noticed when ocr labelled with pyrene was titrated with M.EcoKI (data not shown).

Fluorescence quenching was used to determine whether the labelling sites were in the M.EcoKI–ocr interface or merely affected by some long-range conformational change. Acrylamide, a polar uncharged, electron-deficient molecule efficiently quenches the AEDANS fluorophore when in direct contact with it (Fig. 7). It was observed that acrylamide was able to quench the AEDANS emission from all six labelled forms of ocr only in the absence of saturating amounts of M.EcoKI (Fig. 7). The simplest interpretation of this result is that all six labelling sites on each ocr monomer are part of the interface with M.EcoKI. The labelling sites are dispersed all over the ocr surface thus it is extremely unlikely that M.EcoKI binding causes a conformational change in ocr that indirectly buries all six locations. This result, in conjunction with the crystal structure of ocr (1), indicates that the surface area on ocr involved in binding to M.EcoKI must be very large. These results suggest that M.EcoKI virtually engulfs ocr.

Interaction of ocr with other components of EcoKI

The interaction of ocr with other components of the EcoKI R/M system was also examined. These components were the M subunit, the M₁S₁ dimer and the R subunit, all of which are inactive, and the complete restriction enzyme R.EcoKI (R₂M₂S₁).

Binding of ocr to the complete R.EcoKI nuclease was examined using the change in total tryptophan fluorescence for wild-type R.EcoKI and ocr. This titration indicated a stoichiometry of 1.9 ocr dimers per R.EcoKI (Fig. 8). The same stoichiometry was also observed using pyrene-labelled ocr(D25C) in a titration with R.EcoKI (data not shown).
conclude that two ocr dimers can bind tightly to a single R.EcoKI enzyme.

The change in tryptophan fluorescence was also used as a probe of the interaction of ocr with partially assembled forms of the EcoKI enzyme (Fig. 8). Only weak binding of native ocr to the M subunit alone was observed and a stoichiometry was not clear. The dissociation constant for the interaction between ocr and the M subunit was determined to be 234 ± 69 nM. The M₁S₁ dimer was able to bind tightly to ocr and showed a stoichiometry of 1:1 ocr dimer per M₁S₁. Surprisingly, we also found that ocr bound tightly to the R subunit of the EcoKI R/M system. It appears that saturation is reached when two ocr dimers are bound to a single R subunit.

**DISCUSSION**

**Ocr binding to M.EcoKI**

Our experiments show a strong 1:1 interaction between ocr and M.EcoKI and we believe that the primary mode of binding is one ocr dimer per M.EcoKI. Although there are other modes of binding between ocr and EcoKI, as discussed later, it is of particular importance that the calorimetric titration shows only one binding site as this method provides the most direct determination of the interaction stoichiometry. Calorimetry avoids the uncertainties associated with the use of fluorescence where the experimental observable may result from multiple tryptophan residues within the proteins or from extrinsic labels which may have altered the protein conformation. The 1:1 ratio of ocr dimer to M.EcoKI was also observed using the extrinsic fluorescence from AEDANS-labelled ocr(S68C) and ocr(N43C) and pyrene-labelled ocr(N43C), ocr(D25C), ocr(N43C), ocr(D62C) and ocr(W94C). The slight deviations of these fluorescence measurements from the 1:1 ratio observed in the ITC experiment can be attributed to experimental error in the calculation of the protein concentrations caused by interfering absorption from the labels at 280 nm.

This 1:1 binding of ocr to M.EcoKI has a very favourable free energy change, even greater than that observed between DNA and M.EcoKI, and is strongly exothermic. Given the coverage of the ocr surface with negative charges to mimic the arrangement of phosphates on DNA, the negative enthalpy change is presumably the result of the formation of numerous electrostatic interactions between ocr and M.EcoKI plus further contributions from hydrogen bonding and non-polar contacts. The electrostatic interactions between M.EcoKI and the ocr DNA mimic will be considerable as each monomer of ocr mimics 11 phosphate groups on B-DNA. These interactions may be optimal for the formation of the ocr–M.EcoKI complex as found for the complex between the RNase barnase and its inhibitor barstar (26,27). Unfavourable enthalpy contributions due to the introduction of DNA bending by M.EcoKI, as observed by atomic force microscopy (1), are avoided in the formation of the ocr–M.EcoKI complex as ocr is already bent to an appropriate angle to facilitate M.EcoKI binding. It is likely that most of the free energy difference between M.EcoKI binding to DNA and binding to ocr is due to the enthalpic penalty of bending the DNA substrate. The entropy change in forming the M.EcoKI–ocr complex is unfavourable and such a change is usually attributed to the loss of translational and rotational freedom of the macromolecules, trapping of solvent molecules within the interface between the macromolecules, and the introduction of additional constraints on the conformational dynamics and folding of the macromolecules (28). The magnitude of these contributions to the entropy change are the subject of considerable discussion (see for example 29,30). However, the constraints on flexibility and folding and hence the entropy of the ocr–M.EcoKI complex are likely to be significant as large conformational changes in the average structure of type I methyltransferases (31), with concomitant decreases in flexibility, have been observed upon binding of DNA.

However, as mentioned earlier, it is apparent that the interaction between EcoKI and ocr is more complex than a simple 1:1 complex in which the inhibitor blocks the site on EcoKI which would normally bind to the DNA target sequence. A subset of our titrations show that a second ocr dimer can bind to M.EcoKI. This secondary site appears to be of high affinity as clear saturation points for the site appear in some of the fluorescence titrations. This secondary site is most clearly observed for the ocr(N4C), ocr(D25C) and ocr(D62C) proteins when labelled with AEDANS and for ocr(S68C) when labelled with pyrene. It is likely that the label subtly alters the nature of the interaction between EcoKI and the ocr protein. A stoichiometry of 1.5 ocr per M.EcoKI is also seen when observing the combined tryptophan fluorescence from both ocr and M.EcoKI. However, as mentioned earlier, one might expect such a titration to show a non-integral ratio of ocr to M.EcoKI, firstly because M.EcoKI can dissociate into components each of which binds ocr differently, and secondly as the total number of tryptophan residues in M.EcoKI and ocr is considerable, the signal observed is a composite of many different fluorescent signals. We conclude that this secondary binding site for ocr on M.EcoKI is different in character from the one detected by calorimetry and at least partly reliant upon some change induced in the ocr molecule by the extrinsic label.

**Interaction of ocr with other components of EcoKI**

The interaction of ocr with other components of EcoKI shows that one dimer can bind to M₁S₁, two dimers bind to the R

**Figure 8.** Change in tryptophan emission intensity at 340 nm upon the addition of ocr to R.EcoKI (filled circle), R subunit (open circle), M₁S₁ (filled triangle) and M subunit (open triangle). The change in fluorescence is calculated from the sum of the fluorescence of the individual components minus the fluorescence observed from the mixture of the components. The intensity change for R.EcoKI has been scaled by one-third. The change in tryptophan fluorescence was also used as a probe of the interaction of ocr with partially assembled forms of the EcoKI enzyme (Fig. 8). Only weak binding of native ocr to the M subunit alone was observed and a stoichiometry was not clear. The dissociation constant for the interaction between ocr and the M subunit was determined to be 234 ± 69 nM. The M₁S₁ dimer was able to bind tightly to ocr and showed a stoichiometry of 1:1 ocr dimer per M₁S₁. Surprisingly, we also found that ocr bound tightly to the R subunit of the EcoKI R/M system. It appears that saturation is reached when two ocr dimers are bound to a single R subunit.
subunit and two dimers bind to the complete R\textit{EcoKI} enzyme. If one \textit{ocr} dimer binds per \textit{M.EcoKI} and two \textit{ocr} dimers per R subunit, then one might expect five dimers to bind to the complete \textit{EcoKI} enzyme. This is not observed and we conclude that the multiple binding sites presented by the \textit{M.EcoKI} and R subunits must partially occlude each other in the complete enzyme. The observation of strong binding of \textit{ocr} by the R subunit is noteworthy. \textit{M.EcoKI} alone has a preference for binding to its DNA target sequence rather than to a random DNA sequence (23,24). However, in the absence of cofactors, the complete enzyme binds strongly to all DNA sequences and this was attributed to the interaction of the R subunits with DNA (32). It appears that \textit{ocr} targets not only the sequence-specific binding capability of the \textit{M.EcoKI} core protein but also the non-sequence-specific DNA binding capability of the R subunit.

**Structural implications**

Using the superimposition of \textit{ocr} and DNA shown in Figure 1 and an earlier model of \textit{M.EcoKI} bound to DNA (33), a model of the \textit{M.EcoKI}–\textit{ocr} complex can be constructed by the following procedure: (i) determine the correspondence of nucleotides in Figure 1 with the nucleotides in the target site sequence of \textit{EcoKI}; (ii) superimpose the DNA from Figure 1 with the DNA from the \textit{M.EcoKI}:DNA model; (iii) replace the DNA from Figure 1 with the \textit{ocr} structure and remove the DNA from the \textit{M.EcoKI} model. This then leaves \textit{ocr} bound to \textit{M.EcoKI}. The nature of type I DNA recognition sequences, described below, allows further simplification as we have to consider only one \textit{ocr} monomer and one DNA dodecamer from Figure 1 and half of the \textit{M.EcoKI}–DNA model.

In common with all type I R\textit{M} enzymes, \textit{EcoKI} methylates two adenine bases on complementary DNA strands within a bipartite recognition sequence. The recognition sequence for \textit{EcoKI} is 5'\text{-}AACNNNNNNGTGC-3' (methylation locations underlined). The two adenine bases targeted for methylation by \textit{EcoKI} are separated by an intervening 8 bp. From Figure 1, it is apparent that an extra 2 bp of DNA can be placed between the two dodecamer duplexes to form a continuous stretch of double-stranded DNA. The insertion of these 2 bp requires the two dodecamer duplexes to form a continuous stretch of DNA (36). The only orientation which gives the correct placement of major and minor grooves for the DNA in Figure 1 and the DNA in the model of the TRD (35) has the C9 in the dodecamer, which overlaps with D92 of \textit{ocr}, corresponding to the methylatable adenine in the \textit{EcoKI} target. This produces the alignment shown in Figure 9. This alignment indicates that \textit{ocr} is primarily a mimic of DNA outside the recognition sequence of \textit{EcoKI} and would explain the ability of \textit{ocr} to inhibit type I enzymes with different recognition sequences (1). Given this alignment, one can then replace the DNA from Figure 1 with the \textit{ocr} protein, remove the DNA from the model of the TRD and thereby derive a model of \textit{ocr} against the TRD. As in earlier models (33,38,39), the catalytic domain of the M subunit can then be placed next to the TRD to form an approximate model of part of \textit{EcoKI} bound to ocr (Fig. 10). The models of the TRD and catalytic domain can be superimposed upon each monomer of \textit{ocr} to produce a more complete model of \textit{M.EcoKI} but this elaboration is not necessary to assist with the interpretation of the fluorescence quenching experiments shown in Figure 7.

The physical location of the six sites of protein labelling on \textit{ocr} are well dispersed over the 11 238 \text{Å}² surface area of \textit{ocr} (1). The inability to quench the fluorescence of labels attached to these locations in the presence of \textit{M.EcoKI} (Fig. 7) indicates that the interface between \textit{ocr} and \textit{M.EcoKI} must be substantial. The burial of such a large surface area of \textit{ocr} upon interaction with \textit{M.EcoKI} would be unusual for a protein–protein interface but not without precedent (40,41). Large interfaces typically require substantial conformational changes as observed for the binding of type I methyltransferases to DNA (32). If one highlights the location of the six mutation sites on \textit{ocr} (Fig. 10) one can see that N43 is partially covered by the TRD and catalytic domains, N4 and D62 are completely covered and D25, S68 and W94 are not covered at all. This implies that other parts of the methyltransferase must cover these exposed locations on \textit{ocr} to effectively prevent the quenching of fluorescent probes at these locations. The N-
C-terminal parts of the M subunits would seem to be the most likely parts of the methyltransferase to cover these exposed residues on ocr as these are the largest parts of the methyltransferase for which we possess no structural information.

This work provides a foundation for further analysis of the thermodynamic, kinetic and structural basis for the extra-

ordinary ability of ocr to mimic DNA and thereby inactivate the methyltransferase for which we possess no structural information.

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