Reactions of Cre with Methylphosphonate DNA: Similarities and Contrasts with Flp and Vaccinia Topoisomerase

Chien-Hui Ma1, Aashiq H. Kachroo1, Anna Macieszak2, Tzu-Yang Chen1, Piotr Guga2, Makkuni Jayaram1*

1 Section of Molecular Genetics and Microbiology, University of Texas at Austin, Austin, Texas, United States of America, 2 Department of Bio-organic Chemistry, Center for Molecular and Macromolecular studies, Polish Academy of Sciences, Lodz, Poland

Abstract

Background: Reactions of vaccinia topoisomerase and the tyrosine site-specific recombinase Flp with methylphosphonate (MeP) substituted DNA substrates, have provided important insights into the electrostatic features of the strand cleavage and strand joining steps catalyzed by them. A conserved arginine residue in the catalytic pentad, Arg-223 in topoisomerase and Arg-308 in Flp, is not essential for stabilizing the MeP transition state. Topoisomerase or its R223A variant promotes cleavage of the MeP bond by the active site nucleophile Tyr-274, followed by the rapid hydrolysis of the MeP-tyrosyl intermediate. Flp(R308A), but not wild type Flp, mediates direct hydrolysis of the activated MeP bond. These findings are consistent with a potential role for phosphate electrostatics and active site electrostatics in protecting DNA relaxation and site-specific recombination, respectively, against abortive hydrolysis.

Methodology/Principal Findings: We have examined the effects of DNA containing MeP substitution in the Flp related Cre recombination system. Neutralizing the negative charge at the scissile position does not render the tyrosyl intermediate formed by Cre susceptible to rapid hydrolysis. Furthermore, combining the active site R292A mutation in Cre (equivalent to the R223A and R308A mutations in topoisomerase and Flp, respectively) with MeP substitution does not lead to direct hydrolysis of the scissile MeP bond in DNA. Whereas Cre follows the topoisomerase paradigm during the strand cleavage step, it follows the Flp paradigm during the strand joining step.

Conclusions/Significance: Collectively, the Cre, Flp and topoisomerase results highlight the contribution of conserved electrostatic complementarity between substrate and active site towards transition state stabilization during site-specific recombination and DNA relaxation. They have potential implications for how transesterification reactions in nucleic acids are protected against undesirable abortive side reactions. Such protective mechanisms are significant, given the very real threat of hydrolytic genome damage or disruption of RNA processing due to the cellular abundance and nucleophilicity of water.

In addition to the tyrosine nucleophile, the Cre and Flp active sites are characterized by a catalytic pentad comprised of two arginines, a lysine, a histidine and a tryptophan [5,6]. The type IB topoisomerases also display a similar catalytic constellation, except that the histidine and tryptophan positions are occupied by lysine and histidine, respectively [3,7,8]. There is a striking difference, though, between Cre and Flp in how they assemble their respective active sites. The Flp active site is shared between two neighboring recombinase monomers, one of which provides the pentad and the other the tyrosine nucleophile [5,9]. As a result, strand cleavage by Flp occurs in a trans fashion, that is, across the strand exchange region from where the tyrosine donating Flp monomer is bound (Figure 1B, D). In Cre, the active site is housed entirely within a single monomer, and strand cleavage takes place in cis [3,6]. A non-shared active site and cleavage in cis are also
characteristic of the type IB topoisomerases. The strand cleavage and strand joining steps of recombination or DNA relaxation are thought to follow an acid/base assisted SN2 in-line nucleophilic substitution via a pentacoordinated transition state [10,11]. The array of hydrogen bond donors surrounding the scissile phosphate helps stabilize the transition state, imparting a significant electrophilic character to the recombination and topoisomerase reactions [3].

Substitution of the negatively charged phosphate (P) by the charge-neutral methylphosphonate (MeP) group has been exploited successfully to investigate the importance of electrostatics in nucleic acid-protein interactions as well as catalytic mechanisms during DNA and RNA transactions [12–18]. Particularly relevant to this study are the effects of MeP modification at the scissile phosphate position, coupled with replacement of one of the two conserved arginine residues of the catalytic pentad, on the activities of vaccinia topoisomerase and Flp [19,20]. Activation of the MeP bond is independent of Arg-223 of topoisomerase and the corresponding Arg-308 of Flp [19,20]. Activation of the MeP bond is independent of Arg-223 of topoisomerase and the corresponding Arg-308 of Flp [19,20].

Figure 1. Reaction pathways and active site configurations for tyrosine site-specific recombination promoted by Cre and Flp. A, B. Recombination by Flp and Cre proceeds through two steps of single strand exchanges via a Holliday junction intermediate. During strand cleavage, the pair of recombinase monomers in green activates the scissile phosphodiester bonds in cis for nucleophilic attack by the catalytic tyrosine. The pair of Flp monomers in magenta donates the tyrosine nucleophile in trans. The corresponding Cre monomers are thought to allosterically confer chemical competence on the green monomers that perform strand cleavage in cis. B, C. Cross-sections of the Cre and Flp active sites containing the catalytic pentad and the tyrosine nucleophile are shown [5,6]. In Flp, this tyrosine (shown in magenta) is donated by the neighboring Flp monomer. doi:10.1371/journal.pone.0007248.g001

The activities of Flp and topoisomerase, as well as their mutant derivatives, on P and MeP containing DNA substrates are suggestive of a potential role for phosphate or active site electrostatics in blocking water nucleophile during phosphoryl transfer in these two systems [19–21]. In the topoisomerase reaction, the strand joining step is inherently more vulnerable to potential hydrolysis than the strand cleavage step. Protection against this aberrant reaction is provided by the negative charge on the phosphate group, as inferred from the rapid hydrolysis of the MeP-tyrosyl intermediate by topoisomerase or Topo(R223A) under
physiological pH. In the Flp reaction, it is the strand cleavage step that faces greater risk of hydrolysis than the strand joining step. Here, the positively charged Arg-308 appears to deter water, as indicated by the hydrolytic activity of Flp(R308A) on MeP-DNA.

The vaccinia topoisomerase and Flp paradigms may be relevant to several mechanistically similar phosphoryl transfer reactions in biological systems. They include DNA relaxation by type I and type II topoisomerases, conservative site-specific recombination, DNA transposition, telomere resolution in spirochetes, RNA splicing and retrohoming by mobile introns [10,22–24]. The active nucleasephiles in these reactions are a tyrosine or serine derived from the topoisomerase or recombinase or telomere resolvase active site or a 2', 3' or 5' hydroxyl group derived from a nucleic acid or a nucleoside. It would seem logical that the evolutionary design of active sites performing these reactions, in the context of an overwhelming excess of water (~56M), must have evolved strategies to avoid futile hydrolysis of the nucleic acid substrates or reaction intermediates.

As already noted, topoisoaserase and Flp differ from each other in active site assembly (cis versus trans) and engagement of the scissile phosphate. Furthermore, the DNA relaxation complex and the recombination synapase are organized quite distinctly. The conformational dynamics associated with the two reactions are also discrete. These features may account for the characteristic responses of the two enzymes towards MeP containing DNA substrates. Cre is akin to topoisoaserase, and contrary to Flp, in its mode of active site assembly and strand cleavage. However, Cre resembles Flp in the organization of its recombination synapase and the conformational dynamics of strand exchange [3,4]. Two questions therefore become pertinent. Will Cre or Cre(R292A) unleash a strong hydrolytic activity, as topoisoaserase and Topo(R223A) do, on the cleaved MeP-tyrosyl intermediate? Will Cre(R292A), like Flp(R308A), render the MeP bond in DNA susceptible to direct attack by water? Results presented in this report answer these questions. Hydrolysis of the MeP-tyrosyl bond by Cre or Cre(R292A) is not accelerated markedly compared to that of the P-tyrosyl bond by Cre. In this regard, Cre resembles Flp rather than topoisoaserase. However, Cre(R292A) does not promote direct hydrolysis of the scissile MeP bond in DNA. In this respect, Cre is strikingly different from Flp, and quite like topoisoaserase.

**Results**

**General experimental designs for reactions of Cre on MeP substituted substrates**

MeP substituted DNA substrates are particularly instructive in assessing the role of phosphate and active site electrostatics in the mechanisms of type IB topoisoaserase, Flp, and related phosphoryl transfer reactions [19–21]. Substitution of either of the two non-bridging oxygen atoms of the scissile phosphate by the uncharged methyl group reduces the nominal negative charges in the ground and transition states from −1 to 0 and −2 to −1, respectively. The $R_P$ and $S_P$ stereoisomers of the chiral MeP with respect to the achiral P are diagrammed in Figure 2A. According to convention, replacement of the pro-"R" oxygen yields the $S_P$ form MeP; that of the pro-"S" oxygen results in the $R_P$ form MeP. The majority of reactions reported here were carried out with a mixture of the two MeP diastereomers; a subset of reactions was performed with stereochemically pure $R_P$ or $S_P$ form.

P- or MeP-‘half-site’ substrates contain one Cre binding element and one cleavage site (Figure 2B, C). In the native full-site target of Cre, the spacer between the two Cre binding elements is 8 bp long, with the scissile phosphates on the two strands being 6 bp apart. Although a single monomer of Cre harvests a complete active site, crystal structures suggest that its functionality may require allosteric activation by a second monomer [6,25]. Hence reactions are likely to occur in the context of at least two Cre bound half-sites. Cleavage of a half-site by Tyr-324 of Cre, with formation of the Cre-linked DNA intermediate (Figure 2B), was assayed by SDS-polyacrylamide gel electrophoresis (SDS-PAGE). Potential hydrolysis in a half-site, with either the DNA or the tyrosyl intermediate as target for water attack, was assayed by denaturing urea-polyacrylamide gel electrophoresis (denaturing...
In most reactions, the 5′-OH on the bottom strand of the half-site was blocked by phosphorylation (Figure 2B) to prevent the strand joining reaction. A subset of the reactions utilized a half-site with a modified design in which this 5′ end was free (Figure 2C). In principle, the bottom strand could fold back to assist the hydroxyl group perform nucleophilic attack on the tyrosyl intermediate. This ‘pseudo strand exchange’ reaction within a half-site would produce a hairpin recombinant. Strand exchange is also possible between two half-sites to form a ‘pseudo-full-site’. The recombinant strands resulting from these reactions are identical in sequence, and will migrate as one band during electrophoresis under denaturing conditions. The potential Cre assisted hydrolytic reactions and the corresponding outcomes are indicated in Figure 2D. The target of hydrolysis could be the phosphotyrosyl intermediate (type I) or the scissile phosphodiester bond in DNA (type II). Previous studies have shown that both vaccinia topoisomerase and Flp exhibit the type I activity, whereas the type II activity is unique to Flp(R308A).

Site-specific endonuclease activity of wild type Cre on native and MeP-half-sites

As pointed out, the presence of MeP at the scissile position is sufficient to turn vaccinia topoisomerase, but not Flp, into a strong

![Figure 2](https://example.com/figure2.png)
endonuclease that targets the MeP-tyrosyl intermediate [19,21]. Does MeP substitution induce Cre to behave like topoisomerase or like Flp?

The P- or MeP-half site (a racemic mixture of the R, and S, diastereomers) was treated with wild type Cre under standard conditions employed for recombination assays [26] at pH 7.5. Samples were split into halves, and were analyzed for the DNA-protein covalent adduct and the 24-mer hydrolysis product formed from the strand cleavage and potential endonuclease activities, respectively, of Cre (Figure 3A, B). Here and elsewhere, ‘strand cleavage’ refers exclusively to the reaction catalyzed by Tyr-324 of Cre to distinguish it from strand breakage mediated by water acting as an alternative nucleophile.

With the P-half-site, the covalent adduct formed early, peaking around 6–8 hr, and then steadily declined over the remainder of the reaction course (Figure 3A, left). Formation of the hydrolysis product (24-mer) trailed strand cleavage, increasing continuously and finally accounting for nearly quantitative conversion of the substrate (Figure 3A, right). Yields of the cleaved intermediate and the 24-mer from the MeP-half-site were much lower (Figure 3A and B, right) compared to the P half-site. However, the correlation between the kinetics of 24-mer formation and the decline of the tyrosyl intermediate was more or less similar between the two half-sites. To provide a frame of reference, the outcomes of reactions (1 hr time points) carried out with Flp on its cognate P- and MeP-half-sites are displayed alongside the corresponding Cre reactions.

The Cre reaction patterns suggest that the cleaved P-tyrosyl or MeP-tyrosyl intermediate is the precursor of the corresponding 24-mer hydrolysis product. This precursor-product relationship was also evinced by assays carried out with a P-full-site substrate (Figure S1, Supplemental Material). Consistent with this notion, neither the tyrosyl intermediate nor the 24-mer was formed in half-site reactions performed using the Cre(Y324G) mutant protein (Figure S2).

The 24-mer product from the P-half-site reaction migrated as a roughly equal intensity doublet band (Figure 3B, left). As shown by further characterization (see below), the heterogeneity of the 24-mer was due to distinct 3’ ends, either a free hydroxyl group or a phosphate group. The hydrolysis product from the MeP-half-site was primarily a single band, corresponding to a 3’ methylphosphate end, although a fainter upper band was detectable at very late time points after prolonged phosphorimaging (Figure 3B, right). The 24-mer formation from the MeP-half-site was more abundant in reactions with a Cre mutant lacking the active site Arg-292 (see below; Figure 4).

The results summarized above demonstrate that the tyrosyl intermediate formed during Cre recombination is potentially

---

**Figure 3. Strand cleavage and endonuclease activities of Cre in P- and MeP-half-site substrates.** In the schematic representations of half-site substrates, the asterisk indicates 32P-label at the 5’-end, ‘p’ the scissile phosphate and ‘mp’ the scissile methylphosphonate. Reactions were split into halves to analyze the covalent Cre-DNA adduct (line ending in a circular knob) by SDS-PAGE (A) and the potential hydrolysis product(s) by denaturing PAGE (B). Reactions of Flp with its cognate half-sites in similar assays [19] are provided for comparison. The end-labeled half-site (A) or its labeled strand (B) is denoted by ‘S’ and the hydrolysis product by ‘HP’. The labeled strands in the Cre and Flp half-sites were 27 and 26 nucleotides long, respectively. HP in the Cre reaction was a 24-mer and that in the Flp reaction a 23-mer. The plots below represent mean values from three separate experiments with Cre.

doi:10.1371/journal.pone.0007248.g003
subject to attack by water. However, hydrolysis does not become significant until several hours into the reaction. Under the conditions employed here, recombination between two P-full-sites goes to near completion in less than 30 min (data not shown). In a P-half-site containing a free 5'-hydroxyl group on its bottom strand, the predominant outcome was strand joining, not hydrolysis. The latter was not noticeable until about 3 hr into the reaction (Figure S3). There must be a built-in protective mechanism, other than phosphate electrostatics, to ensure that the 5'-hydroxyl group has a strong advantage over water as a nucleophile during strand joining (see under ‘Discussion’).

Cre is similar to Flp, and distinct from vaccinia topoisomerase, in that MeP substitution does not strongly enhance its endonucleolytic activity directed against the tyrosyl intermediate [19,21] (Table 1). The efficiency of strand cleavage by Cre is significantly reduced in the MeP-half-site compared to the P-half-site. Once formed, the MeP-tyrosyl adduct does not turn over rapidly, but is hydrolyzed over hours (Figure 3B, [19]). By contrast, the rate of hydrolysis of the MeP-tyrosyl intermediate formed by topoisomerase is rapid compared to its P-tyrosyl counterpart [21] (Table 1). The half-life of the former at physiological pH is less than 2 minutes; that of the latter is roughly 36 days.

Replacement of Arg-292 of Cre by alanine enhances strand cleavage in the MeP-half-site and correspondingly hydrolysis of the MeP-tyrosyl intermediate

Neutralizing the charge on the scissile phosphate by its replacement with MeP alleviates the requirement of Arg-223 in vaccinia topoisomerase and the corresponding Arg-308 in Flp for activating the MeP target for nucleophilic attack [19,20]. Topo(R223A) forms the MeP-tyrosyl intermediate, and hydrolyzes it with roughly the same efficiency as the wild type enzyme. The reductions in the rates of strand cleavage and hydrolysis due to this mutation are within a factor of two (see Table 1). Flp(R308A), by contrast, directly hydrolyzes the MeP bond in DNA, preempting Tyr-343 mediated strand cleavage. Is Cre(R292A), the counterpart of Topo(R223A) and Flp(308A), competent for MeP activation? If it is, does it promote nucleophilic attack by Tyr-324 as in the topoisomerase case or direct hydrolysis (type II endonucleolytic activity) as in the Flp case?

Cre(R292A) was virtually nonreactive towards the P-half-site. Only a trace of the P-tyrosyl intermediate was detected (Figure 4A, left), and the 24-mer formation was also correspondingly inefficient. These results are consistent with the essential role of Arg-292 in the phosphate activation step. The MeP-half-site readily gave rise to the MeP-tyrosyl intermediate as well as the 24-
mer in the Cre(R292A) reaction (Figure 4A and B, right). The MeP-tyrosyl intermediate reached its maximal levels at approximately 16–24 hr, and thereafter showed a slow, steady decay. The 24-mer continued to accumulate throughout the entire reaction course. In the Flp(R308A) reaction on its cognate MeP-half-site, shown here for comparison, the hydrolysis product (23-mer in this case) was formed directly without mediation of the MeP-tyrosyl intermediate [19].

The kinetic data in Figure 4 are consistent with the 24-mer being the hydrolytic breakdown product of the MeP-tyrosyl intermediate. In accordance with their precursor-product relationship, treatment of the MeP-half-site with Cre(R292A,Y324G) did not result in the formation of either the cleaved intermediate or the 24-mer (Figure 5).

The endonucleolytic activity of Cre(R292A) on the scissile MeP bond is mechanistically similar to that of vaccinia topoisomerase, Topo(R223A) [20,21] and the corresponding activity of wild type Flp, termed type 1 Flp endonuclease [19]. The Cre(R292A) activity is distinct from the type II endonuclease activity of Flp(R308A) whose target is the scissile MeP bond in DNA rather than the cleaved MeP-tyrosyl intermediate [19].

Kinetic parameters for strand cleavage and hydrolysis of the cleaved intermediate by Cre, Flp and topoisomerase in P- and MeP-half-sites

Under our assay conditions, the first order rate constant for strand cleavage in the P-half-site by Cre was slower by a factor of ~100 compared to Flp ($k_1$ of $1.2 \times 10^{-5}$ s$^{-1}$ versus $1.8 \times 10^{-7}$ s$^{-1}$) (Table 1). Because of the very low yields of strand cleavage and hydrolysis products from the MeP-half-site in Cre reactions, the corresponding rate constants were not determined. However, values of $k_4$ for the MeP-half-site were comparable between Cre(R292A) and Flp ($3.2 \times 10^{-3}$ s$^{-1}$ and $2.4 \times 10^{-4}$ s$^{-1}$, respectively) (Table 1). The rate constant for hydrolysis of the MeP-tyrosyl intermediate by Cre(R292A) was ~7 times that of Flp ($k_{\text{hydro}}$ of $0.5 \times 10^{-4}$ s$^{-1}$ versus $1.3 \times 10^{-5}$ s$^{-1}$) (Table 1). Note, though, that there was only about 3 fold enhancement in $k_{\text{hydro}}$ of the MeP-tyrosyl intermediate by Cre(R292A) relative to $k_{\text{hydro}}$ of the P-tyrosyl intermediate by Cre ($8.8 \times 10^{-5}$ s$^{-1}$ and $2.8 \times 10^{-5}$ s$^{-1}$, respectively) (Table 1). The corresponding relative $k_{\text{hydro}}$ enhancement by the vaccinia topoisomerase or Topo(R223A) was much more dramatic, 1.6 to $3.2 \times 10^4$ ($k_{\text{hydro}}$ of P-tyrosyl intermediate by topoisomerase = $2.2 \times 10^{-7}$ s$^{-1}$; $k_{\text{hydro}}$ of MeP-tyrosyl intermediate by topoisomerase and Topo (R223A) = $7.0 \times 10^{-5}$ s$^{-1}$ and $3.5 \times 10^{-3}$ s$^{-1}$, respectively) (Table 1).

Taken together, the results from Figures 3 and 4 and the kinetic constants in Table 1 argue against a significant role for either phosphate or active site electrostatics in guarding the strand cleaved intermediate formed by Cre against hydrolysis. They highlight the effect of eliminating Arg-292, critical for activity on the scissile phosphodiester bond, in enhancing the strand cleavage activity of Cre on the scissile MeP bond.

It should be noted that the methylphosphonates are not intrinsically prone to hydrolysis at neutral pH. Oligonucleotides containing MeP linkages are stable in concentrated ammonium hydroxide at 36°C for up to 16 hr [27]. Direct hydrolysis of the scissile MeP bond by Flp (R308A) is significant against the absence of this reaction with Topo(R223A) and Cre(R292A). Similarly, the striking rate enhancement in the hydrolysis of the MeP-tyrosyl bond by topoisomerase and Topo (R223A) sharply differs from the lack of such enhancement by Cre, Cre(R292A), Flp and Flp(R308A).

Stereospecificity of the Cre(R292A) endonucleolytic activity: characterization of the 3′-terminus of the hydrolysis product from P- and MeP-half-sites

Cre crystal structures indicate potential hydrogen bonding interactions between the two arginine residues of the catalytic pentad and the nonbridging oxygen atoms of the scissile phosphate [3]. Substitution of either one of these oxygen atoms by a methyl group is likely to affect MeP reactivity. Furthermore, the magnitude of this effect could be different in the two cases. We therefore tested the potential stereochemical specificity of the Cre(R292A) endonucleolytic reaction using pure SP and RP forms of the MeP-half-site substrate. The hydrolysis product of Cre reaction on the P-half-site migrates as a doublet during electrophoresis whereas that of Cre(R292A) reaction on the MeP-half-site is almost exclusively a single band (Figures 3A and 4B). We wished to characterize these products with respect to their 3′ ends.

Reactions of Cre(R292A) revealed a strong preference for the SP MeP-half-site over the RP form (Figure 6A). The SP reaction showed an initial rapid phase of ~30 min, followed by a slower phase. The RP reaction followed a more uniform pace. For the early phase, the rate of product formation was roughly 7-fold higher for the SP form. The product yield at the 24 hr time point favored the SP over the RP form by a factor of approximately 3. This bias in 24-mer formation was qualitatively the same as that reported for vaccinia topoisomerase and Flp. Since strand cleavage by the Tyr-343 nucleophile is accompanied by inversion of configuration at the phosphate center, the hydrolysis reaction targets the MeP-tyrosyl intermediate whose chirality is opposite to that of the starting SP MeP-half-site.

The upper 24-mer band from Cre endonuclease activity (lane 2, Figure 6B) on the P-half site was determined to contain a hydroxyl

Figure 5. Lack of endonucleolytic activity in MeP-half-site reactions of Cre(R292A, Y324G). Reactions and product analysis were performed as described under Figure 3. The results of SDS-PAGE and denaturing PAGE analyses are shown in A and B, respectively. The predicted positions of migration of the covalent intermediate (line ending in a circular knob) and the hydrolysis product (HP) are indicated. There was a faint band of background hydrolysis in the control reaction, which did not increase in intensity with increasing time of incubation with Cre. doi:10.1371/journal.pone.0007248.g005
group at the 3′ end by the following criteria. It migrated slightly above the appropriate rung of a Maxam-Gilbert sequencing ladder derived from the labeled strand of the half-site (lane 1 (MG), Figure 6B), and co-migrated with a synthetic 24-mer of the predicted sequence bearing a 3′-OH (lane 3 (M), Figure 6B). Furthermore, this hydrolysis product and the synthetic 24-mer served as primers in a sequenase primer extension reaction (lanes 5, 7, Figure 6B). By analogous reasoning, the lower 24-mer band was deemed to contain a 3′-phosphate end. It co-migrated with the appropriate band of the sequencing ladder and slightly below the synthetic 24-mer (lanes 1–3, Figure 6B), and failed to serve as a primer in the chain extension reaction (lanes 5, Figure 6B). The relative mobility of the principal 24-mer band from Cre(R292A) reaction on the MeP-half-site (lane 2, Figure 6C) with respect to the sequence ladder (lane 1 (MG), Figure 6C) and the synthetic 24-mer marker (lane 3 (M), Figure 6C), as well as its failure to be extended by sequenase (lane 5, Figure 6C) in contrast to M (lane 7, Figure 6C), was consistent with its harboring a 3′-methylphosphate end. Note that chemical cleavage at the MeP position during the sequencing reaction yielded a doublet (lane 1, Figure 6C). The lower band was inferred to contain a 3′-methylphosphate end and the upper one a 3′-OH end, based on their behavior in the sequenase reaction (lane 9, Figure 6C).

Cre, Flp and vaccinia topoisomerase are mechanistically equivalent in that their hydrolytic actions on the respective MeP-tyrosyl intermediates generate products with a 3′-methylphosphate end, as the nearly exclusive or at least the major output. In these reactions, the water nucleophile is oriented opposite a tyrosine leaving group. Hydrolysis of the phosphotyrosyl intermediate by Flp and Cre yields products with either a 3′-OH or a 3′-phosphate end. It is possible that the hydroxyl end is generated by dephosphorylation. However, incubation of the isolated 24-mer with Cre for up to 8 hr did not remove the phosphate group from the 3′ end (Figure S4). By contrast, treatment with T4 polynucleotide kinase in the absence of ATP quantitatively dephosphorylated the 3′ end in 30 min. An intriguing alternative to dephosphorylation of the 24-mer during Cre reaction is for the attacking water molecule to line up opposite the 3′-hydroxyl end.
group. This would require considerable conformational flexibility within the active site. Both Flp and Cre have been shown to utilize a vicinal 2'-OH to attack the phosphotyrosyl bond when the scissile position is substituted with a ribonucleotide [28,29].

Discussion

We have examined the activities of the Cre recombinase and its active site variant Cre(R292A) on DNA half-site substrates containing MeP replacement of the scissile phosphate. We shall discuss the current results and their implications in the context of similar analyses carried out previously on vaccinia topoisomerase and Flp recombinase [19–21].

The requirement of a conserved catalytic arginine in Cre, Flp and vaccinia topoisomerase is mandated by phosphate electrostatics

Arg-292 of Cre, like the corresponding conserved Arg-308 of Flp and Arg-223 of topoisomerase, becomes dispensable in activating the scissile phosphodiester bond when the negatively charged oxygen is replaced by the uncharged methyl group. In fact, Arg-292 poses serious impediment to MeP activation by Cre, as evinced by the ability of the R292A mutation to strongly stimulate strand cleavage in the MeP-half-site. Arg-223 of topoisomerase is more neutral with respect to MeP activation. The R223A mutation does not appreciably alter the rate of formation of the MeP-tyrosyl intermediate or of its subsequent hydrolytic breakdown [20,21]. Unlike Cre and topoisomerase, Flp does not yield strand cleavage in the MeP substrate by Tyr-343 nucleophile when it harbors the R308A mutation. This is not due to lack of MeP activation; Flp(R308A) targets the MeP bond for direct hydrolysis (type II endonucleolytic activity) [19].

Effect of MeP substitution on recombinase and topoisomerase activities: more than just active site perturbation?

The rates of strand cleavage by wild type topoisomerase and Flp in their MeP-substrates are reduced roughly by factors of $3 \times 10^2$ and $10^3$, respectively, relative to the native P-substrates. [19,20][see Table 1]. The magnitude of this effect for Cre is likely to be in this range or perhaps even greater. Because of weak activity, we have not determined the rate constant for MeP-half-site cleavage by Cre.

The lower reactivity of the wild type recombinases and topoisomerase towards MeP may, at least in part, be accounted for by active site perturbations induced by the methyl modification. The larger van der Waals radius of the methyl group ($\sim 1.8 \, \text{Å}$) compared to the oxygen atom ($\sim 1.5 \, \text{Å}$), together with its non-polar character, could lead to steric clashes with the guanidinium group of Arg-292. Repositioning of the scissile MeP and/or active site residues to accommodate this conflict may thus result in disruption or weakening of contacts required for normal catalytic power. The stimulation of strand cleavage by the R292A mutation in Cre fits this explanation.

Nevertheless, removal of the arginine side chain may do more to catalysis than merely rectify the steric problem posed by MeP. For topoisomerase and Flp, the salutary effects of R223A and R308A mutations, respectively, on catalytic efficiency per se are not dramatic. The rate constants for strand cleavage in the MeP-substrate are within a factor of two for wild type topoisomerase ($1.4 \times 10^{-3} \, \text{s}^{-1}$) and Topo(R223A) ($7 \times 10^{-4} \, \text{s}^{-1}$) [20] (Figure 1). Strikingly, both proteins target the MeP-tyrosyl intermediate for rapid hydrolysis. Flp cleaves the MeP bond using the Tyr-343 nucleophile ($2.4 \times 10^{-4} \, \text{s}^{-1}$) and Flp(R308A) hydrolyzes it directly ($3.7 \times 10^{-3} \, \text{s}^{-1}$) with comparable rate constants [19] (Table 1). Thus methyl substitution by itself (as revealed by topoisomerase) or this substitution plus the removal of a catalytic arginine residue (as revealed by Flp/R308A) can dramatically alter the native reactions qualitatively.

Protection against hydrolysis by phosphate and active site electrostatics?

Transesterification events involving phosphodiester bonds in DNA and RNA have to deal with the potential problem of water acting as an intrusive alternative nucleophile. The abortive hydrolysis of the scissile MeP bond by Flp(R308A) and that of the MeP-tyrosyl intermediate by vaccinia topoisomerase highlight the threat from water during site-specific recombination and DNA relaxation. These aberrant reactions induced by charge neutralization suggest that phosphate electrostatics or active site electrostatics may help repel water during the normal reaction. MeP substitution, in conjunction with specific active site mutations, could reveal related protective mechanisms in other enzyme systems that carry out phosphoryl transfer reactions in nucleic acids. Such mechanisms are expected to be modulated among individual enzyme systems according to their active site configurations as well as DNA-protein and protein-protein interactions and conformational dynamics within their reaction complexes. These aspects are considered below for Cre, Flp and topoisomerase.

Guarding the scissile phosphodiester bond against hydrolysis during strand cleavage

Topoisomerase binds to DNA and relaxes it as a monomer. Flp and Cre also bind each of the two binding elements on their target sites as monomers. Because of their cis configuration, the topoisomerase and Cre active sites engage the scissile phosphodiester contemporaneous to DNA binding. This would leave little chance for water to channel the cleavage reaction towards hydrolysis (Figure 7A and B). Whereas the topoisomerase monomer is fully competent for strand cleavage, the Cre monomer likely requires allosteric activation by a neighboring monomer to become cleavage proficient [36].

Binding of a Flp monomer to DNA concomitantly activates the scissile phosphate [30]. However, it must await the binding of a second Flp monomer to acquire Tyr-343 and complete the assembly of its trans active site. The temporal separation between binding and active site engagement makes the scissile phosphate vulnerable to attack by water (Figure 7C). This risk appears to be minimized through the positively charged side chain of Arg-308. In Cre and topoisomerase, the role of Arg-292 and Arg-223, respectively, is purely catalytic; namely, activation of the scissile phosphate; in Flp, Arg-308 appears to play an additional non-catalytic role, namely, repelling or misorienting water from the activated phosphate.

Cre follows the topoisomerase, rather than Flp, paradigm to protect the strand cleavage reaction against hydrolysis.

Protecting the tyrosyl intermediate against attack by water during strand joining

In the case of Cre, as is true for Flp as well, the susceptibility of the strand cleavage intermediate to hydrolysis is not particularly exacerbated by the presence of MeP at the scissile DNA position. This contrasts with the vaccinia topoisomerase reaction, in which the rate of hydrolysis of the MeP-tyrosyl intermediate is accelerated by a factor of $\sim 3$ to $4 \times 10^3$ relative to the p-tyrosyl
intermediate [20,21] (Table 1). During DNA relaxation, the cleaved strand must go through a round of rotation before joining occurs. The conformational flexibility associated with the dynamics of rotation may leave the phosphotyrosyl intermediate an open target for attack by water (Figure 7D). However, the negative charge on the phosphate keeps water out of the reaction center [20,21].

Protection of the joining reaction from water during Cre and Flp recombination appears to be conferred by the compact design of their synaptic structures (see Figure 1A, B). The interface between two Flp monomers that partake in the trans donation of tyrosine and assembly of a functional active site displays extensive protein-protein interactions that bury a large surface area of, 1500 Å² [31]. The buried surface area in the Cre tetramer (1,100 Å²), though not as extensive, is nevertheless quite large [3,6,31]. Consistent with this difference perhaps, the hydrolysis product from full-site reactions appears earlier, and is more prominent, with Cre compared to Flp (this study; [19]). Crystal structures suggest that, within the tight synaptic architecture, strand cleavage and reorientation of the resulting 5'-hydroxyl groups in the strand joining orientation are strongly coupled events. The conformational dynamics associated with strand cleavage can effectively close the door to water.

Cre utilizes the Flp, rather than topoisomerase, strategy to protect the integrity of the strand joining reaction.

**Materials and Methods**

**Synthetic oligonucleotides and assembly of substrates**

A list of the oligonucleotides used for the assembly of half-site and full-site substrates is given in Table S1 (Supplementary data). All standard oligonucleotides were purchased from IDT (Coralville, IA). The methylphosphonate containing oligonucleotide (a racemic mixture of the R_P and S_P forms) was supplied by Trilink Technologies (San Diego, CA). Hybridization of oligonucleotides was performed as described previously [32].

Stereochemically pure R_P and S_P methylphosphonate containing oligonucleotides

The oligonucleotide substrates containing R_P or S_P form of methylphosphonate were obtained as described previously [19]. Details of the individual steps of stereoisomer separation of MeP containing dinucleotides and oligonucleotide synthesis can be provided upon request.

**Purification of Cre and Cre mutants**

Wild type Cre and its mutant derivatives, harboring a His-6 tag at their N-termini, were obtained as described previously [33]. Proteins were expressed in *Escherichia coli* BL21(DE3)pLysS strain by induction with 1 mM IPTG. They were purified from cell extracts by affinity chromatography on Ni Sepharose 6 Fast Flow columns (GE Healthcare, Piscataway, NJ). After buffer exchange through Econo-Pac 10DG columns (Bio-Rad, Hercules, CA), they were stored in 100 μl aliquots (containing 20% glycerol) at −70°C. The final purity of the protein preparations ranged from 65−80%, as judged by SDS-PAGE analysis.

**In vitro Cre reactions**

In vitro Cre reactions were carried out using 5'-end labeled substrates at 30°C under conditions described by Ghosh et al. [26]. Each 30 μl reaction mixture contained 0.2 pmole end-labeled DNA substrate in 50 mM Tris-HCl, pH 7.5, 50 mM NaCl, 10 mM MgCl₂ and 1 mM DTT. After addition of approximately 6−8 monomers of Cre or Cre mutant per Cre

![Figure 7](image-url)
binding element, reaction mixtures were incubated at 30°C for various times. Reactions were terminated by addition of SDS to a final concentration of 0.1%. For assaying the Cre-DNA covalent adduct, samples were analyzed directly by SDS-PAGE. For assaying Cre endonuclease activity, the proteinase K treatment that normally precedes DNA extraction was omitted. By doing so, following electrophoresis, the hydrolysis product could be visualized without interference from the strand cleavage product carrying short Cre peptides at the 3’ end.

Characterization of hydrolysis products by the sequenase extension reaction

Cre reactions were stopped by addition of SDS (0.1%), and samples were switched to the sequenase reaction buffer by passage over QiAquick PCR purification columns (QiAGEN). Extension reactions were done for 5 min at 37°C in the presence of all four deoxynucleoside triphosphates (0.5 mM final concentration), and analyzed by denaturing PAGE. The marker oligonucleotide or DNA purified (by phenol-chloroform extraction and ethanol precipitation) from the Maxam-Gilbert sequencing reactions was hybridized to the bottom stand of the half-site prior to treatment with sequenase.

Dephosphorylation by T4 polynucleotide kinase

The T4 polynucleotide kinase (New England BioLabs) reaction was performed using 10 units of the enzyme without addition of ATP. The incubation was done in the T4 polynucleotide kinase reaction buffer provided by the supplier (70 mM Tris-HCl, 10 mM MgCl₂, 5 mM DTT, pH 7.6 at 25°C) for 30 min at 37°C.

Curve fitting to the kinetic data

Curves were fit to kinetic data by non-linear regression according to ‘one-phase-association’ using the software GraphPad Prism (version 5.02). For reactions that displayed early rapid kinetics followed by later slower kinetics, separation of the individual kinetic components by biphasic curve fitting was not attempted. First order rate constants were derived from the fitted curves based on the assumption that strand cleavage in the half-site substrates was irreversible (see also legend to Table 1).

Quantification of reaction products

Bands corresponding to substrate and products were quantified using a phosphorimager. For low efficiency reactions, Conversion of substrate to product was quantified as follows. Multiple serial dilutions of a reaction were fractionated alongside the undiluted reaction, and the substrate band was quantified from those that fell within the linear range of estimation. Appropriate correction factors were then applied in the calculation of product yields.

Supporting Information

**Figure S1** Strand cleavage and endonucleolytic activities of Cre on a full-site substrate. In a schematic representation of the full-site substrate, the 32P-label at the 5’-end of the top strand and the scissile phosphates are indicated by the asterisk and ‘P’, respectively. Reactions were split into two equal portions, and analyzed for the covalent Cre-DNA adduct (line ending a circular knob) and the hydrolysis product (HP; 24-mer). SDS-PAGE and denaturing PAGE profiles are shown in A and B, respectively. In B, the mid-section of the gel, bereft of radioactive bands, was trimmed out. The end-labeled full-site (A) or its labeled strand (B) is denoted by ‘S’. The labeled strand was 54 nucleotides long. The results from two independent experiments are plotted below with error bars. The nearly complete hydrolysis of the covalent intermediate under the reaction conditions employed was rather surprising. Nevertheless, the recombination reaction carried out under similar conditions goes to completion well before hydrolysis gains ground (data not shown). Even in a half-site reaction, a 5’-hydroxyl group present on the bottom strand strongly competes out water as the nucleophile in the strand joining reaction (see Figure S3). Assays using the full-site 5’ end-labeled on the bottom strand yielded qualitatively similar results except that the hydrolysis reaction was not as strong as with the top strand labeled substrate. The maximal yield of the 24-mer product was approximately 20-30% of the input substrate (data not shown). This difference is perhaps due to the asymmetry in the cleavage of the scissile phosphodiester bonds on the two strands by Cre [1–7]. The direction of this asymmetry, towards the top or the bottom strand, is apparently influenced by whether cleavage occurs prior to or following synopsis of the recombination sites [2]. The cleavage reaction in full-sites is readily reversible by reformation of the parental strands or by strand exchange in the recombinant mode to form the Holliday junction. At early time points in our reactions (before its diminution by hydrolysis), the amount of covalent Cre-DNA complex formed on the top strand was greater than that on the bottom strand (data not shown). The longer-lived intermediate is, the more susceptible it may become to attack by water. 1. Ennifar E, Meyer JE, Bachholz F, Stewart AF, Suck D (2003) Crystal structure of a wild-type Cre recombinase-loxP synapase reveals a novel spacer conformation suggesting an alternative mechanism for DNA cleavage activation. Nucleic Acids Res 31: 5449–5460. 2. Ghosh K, Lau CK, Gupta K, Van Dunye GD (2005) Preferential synthesis of loxP sites drives ordered strand exchange in Cre-loxP site-specific recombination. Nat Chem Biol 1: 275–282. 3. Hoess R, Wierzbiick A, Ahrens K (1987) Isolation and characterization of intermediates in site-specific recombination. Proc Natl Acad Sci USA 84: 6840–6844. 4. Lee G, Saito I (1998) Role of nucleotide sequences of loxP spacer region in Cre-mediated recombination. Gene 216: 55–65. 5. Lee L, Sadowski PD (2003) Sequence of the loxP site determines the order of strand exchange by the Cre recombinase. J Mol Biol 326: 397–412. 6. Martin SS, Pulido E, Chu VC, Lechner TS, Baldwin EP (2002) The order of strand exchanges in Cre-LoxP recombination and its basis suggested by the crystal structure of a Cre-LoxP Holliday junction complex. J Mol Biol 319: 107–127. 7. Tribble G, Ahn YT, Lee J, Dandekar T, Jayaram M (2000) DNA recognition, strand selectivity, and cleavage mode during integrase family site-specific recombination. J Biol Chem 275: 22255–22267.

**Figure S2** Reactions of Cre(Y324G) on MeP-DNA

- **Figure S3** Competition between the 5’-OH and water nucleophiles during strand joining in a P-half-site. The half-site designed to follow strand joining contained a longer bottom strand than the standard half-sites shown in **Figure S2**. The 5′-hydroxyl group of this strand was left unblocked. Looping back of the single stranded region would place the 5′-hydroxyl in position to attack the tyrosyl intermediate formed by strand cleavage. This intra-half-site reaction would produce a hairpin (63-mer). Strand joining could...
potentially occur between two half-sites to yield a ‘pseudo-full-site’. The individual strands of the pseudo-full-site and the hairpin would be identical in sequence and display the same mobility during denaturing PAGE. JP refers to this joined product, which was formed early and in greater abundance than the hydrolysis product (HP). The labeled strand of the half-site is denoted by ‘S’.

The gel profile shown here was compressed by excluding its bare mic section.

Found at: doi:10.1371/journal.pone.0007248.s006 (0.46 MB TIF)

**Table S1** The synthetic oligonucleotides used for the assembly of the substrates used in this study are listed. *The bottom strand with a longer than normal 3'-extension was used to assemble the half-site substrate used in the strand joining assay (see Figure S5).*

Found at: doi:10.1371/journal.pone.0007248.s005 (0.03 MB DOC)

**Figure S4** Incubation of the 24-mer hydrolysis product with Cre does not cause 3' end dephosphorylation. The 5' end-labeled P-half-site was treated with Cre for 144 hr to convert it nearly quantitatively into the hydrolysis product via the cleaved intermediate. After phenol-chloroform extraction and ethanol precipitation, the isolated DNA was reincubated with Cre in the same buffer as that employed in the hydrolysis reaction. Incubation for up to 6 hr did not result in the removal of the 3'-phosphate (the lower 24-mer band). By contrast, incubation for 30 min with T4 polynucleotide kinase without addition of ATP led to complete removal of the phosphate group from the 3' end.

Found at: doi:10.1371/journal.pone.0007248.s004 (0.46 MB TIF)

**References**

1. Jayaram M, Mehta S, Uzri D, Vozyanov Y, Velmurugan S (2004) Site-specific recombination and partitioning systems in the stable high copy propagation of the 2-micron yeast plasmid. In: Moldave K. ed. Prog Nucl Acid Res Mol Biol Academic Press. Vol 77: pp 127–172.

2. Jayaram M, Tribble G, Grainge I, editors (2002) Site-specific recombination by the Flp protein of Saccharomyces cerevisiae. In: Craig NL, Craigo R, Gelfett M, Lambowitz AM, eds. Mobile DNA II. Washington DC: ASM Press. pp 192–210.

3. Van Dyne GD (2002) A structural view of tyrosine recombinase site-specific recombination. In: Craig NL, Craigo R, Gelfett M, Lambowitz AM, eds. Mobile DNA II. Washington DC: ASM Press. pp 93–117.

4. Rice PA (2002) Theme and variation in tyrosine recombinases: structure of a Flp-DNA complex. In: Craig NL, Craigo R, Gelfett M, Lambowitz AM, eds. Mobile DNA II. Washington DC: ASM Press. pp 219–229.

5. Chen Y, Narendra U, Iype LE, Cox MM, Rice PA (2000) Crystal structure of a Flp recombinase-Holliday junction complex: assembly of an active oligomer by helix swapping. Mol Cell 6: 885–897.

6. Guo F, Gopal DN, Van Dyne GD (1997) Structure of Cre recombinase complexed with DNA in a site-specific recombination synapse. Nature 389: 49–46.

7. Krogh BO, Shuman S (2000) Catalytic mechanism of DNA topoisomerase IB. Mol Cell 5: 1033–1041.

8. Redinbo MR, Stewart I, Kuhn F, Champoux JJ, Hol WG (1998) Crystal structures of human topoisomerase I in covalent and noncovalent complexes with DNA. Science 279: 1504–1513.

9. Chen Y, Rice PA (2003) The role of the conserved Trp330 in Flp-mediated end dephosphorylation. The 5'-30 min with T4 polynucleotide kinase without addition of ATP led to complete removal of the phosphate group from the 3' end.

Found at: doi:10.1371/journal.pone.0007248.s006 (0.46 MB TIF)

**Figure S4** Incubation of the 24-mer hydrolysis product with Cre does not cause 3' end dephosphorylation. The 5' end-labeled P-half-site was treated with Cre for 144 hr to convert it nearly quantitatively into the hydrolysis product via the cleaved intermediate. After phenol-chloroform extraction and ethanol precipitation, the isolated DNA was reincubated with Cre in the same buffer as that employed in the hydrolysis reaction. Incubation for up to 6 hr did not result in the removal of the 3'-phosphate (the lower 24-mer band). By contrast, incubation for 30 min with T4 polynucleotide kinase without addition of ATP led to complete removal of the phosphate group from the 3' end.

Found at: doi:10.1371/journal.pone.0007248.s006 (0.46 MB TIF)

**Figure S4** Incubation of the 24-mer hydrolysis product with Cre does not cause 3' end dephosphorylation. The 5' end-labeled P-half-site was treated with Cre for 144 hr to convert it nearly quantitatively into the hydrolysis product via the cleaved intermediate. After phenol-chloroform extraction and ethanol precipitation, the isolated DNA was reincubated with Cre in the same buffer as that employed in the hydrolysis reaction. Incubation for up to 6 hr did not result in the removal of the 3'-phosphate (the lower 24-mer band). By contrast, incubation for 30 min with T4 polynucleotide kinase without addition of ATP led to complete removal of the phosphate group from the 3' end.

Found at: doi:10.1371/journal.pone.0007248.s006 (0.46 MB TIF)

**Table S1** The synthetic oligonucleotides used for the assembly of the substrates used in this study are listed. *The bottom strand with a longer than normal 3'-extension was used to assemble the half-site substrate used in the strand joining assay (see Figure S5).*

Found at: doi:10.1371/journal.pone.0007248.s005 (0.03 MB DOC)

**Acknowledgments**

We thank Milena Sobczak for excellent technical assistance during synthesis of methylphosphonate containing oligonucleotides. We are grateful to Malgorzata Bukowiecka-Matusiak for useful suggestions for synthesis of methylphosphonate dimers. We acknowledge the many useful comments from E. P. Baldwin for improving the interpretation and presentation of results.

**Author Contributions**

Conceived and designed the experiments: CHM AHK PG MJ. Performed the experiments: CHM AHK. Analyzed the data: CHM AHK. Contributed reagents/materials/analysis tools: CHM AM TYC PG. Wrote the paper: CHM MJ.