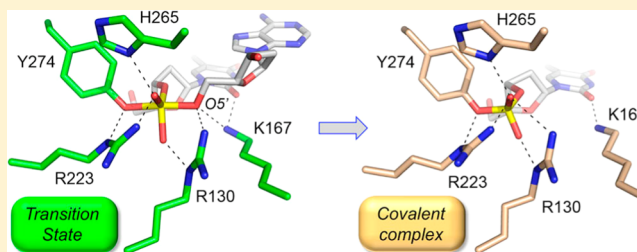


Chemical Mutagenesis of Vaccinia DNA Topoisomerase Lysine 167 Provides Insights to the Catalysis of DNA Transesterification

Lyudmila Yakovleva and Stewart Shuman*

Molecular Biology Program, Sloan-Kettering Institute, New York, New York 10065, United States

ABSTRACT: Vaccinia DNA topoisomerase IB (TopIB) relaxes supercoils by forming and resealing a covalent DNA–(3′-phosphotyrosyl)²⁷⁴–enzyme intermediate. Conserved active site side chains promote the attack of Tyr274 on the scissile phosphodiester via transition state stabilization and general acid catalysis. Two essential side chains, Lys167 and Arg130, act in concert to protonate and expel the 5′-O leaving group. Here we gained new insights to catalysis through chemical mutagenesis of Lys167. Changing Lys167 to cysteine crippled the DNA cleavage and religation transesterification steps ($k_{cl} = 4.3 \times 10^{-4} \text{ s}^{-1}$; $k_{rel} = 9 \times 10^{-4} \text{ s}^{-1}$). The transesterification activities of the K167C enzyme were revived by in vitro alkylation with 2-bromoethylamine ($k_{cl} = 0.031 \text{ s}^{-1}$; $k_{rel} \geq 0.4 \text{ s}^{-1}$) and 3-bromopropylamine ($k_{cl} = 0.013 \text{ s}^{-1}$; $k_{rel} = 0.22 \text{ s}^{-1}$), which convert the cysteines to γ -thialysine and γ -thiahomolysine, respectively. These chemically installed lysine analogues were more effective than a genetically programmed arginine 167 substitution characterized previously. The modest differences in the transesterification rates of the 2-bromoethylamine- and 3-bromopropylamine-treated enzymes highlight that TopIB is tolerant of a longer homolysine side chain for assembly of the active site and formation of the transition state.



Type IB DNA topoisomerases (TopIB) relax DNA supercoils by repeatedly breaking and rejoining one strand of the DNA duplex through a covalent DNA–(3′-phosphotyrosyl)-enzyme intermediate. Vaccinia virus TopIB, which displays stringent specificity for cleavage at the sequence 5′(C/T)CCTT↓ in the scissile strand,^{1–5} has been an instructive model system for mechanistic studies of the TopIB family. The cleavage and religation transesterification reactions are driven by four amino acid side chains (Arg130, Lys167, Arg223, and His265) that comprise the active site and catalyze the attack of Tyr274 on the scissile phosphodiester.^{6–10} The two arginines and the histidine interact directly with the scissile phosphodiester in the crystal structure of the covalent TopIB–DNA intermediate (Figure 1) and are proposed to stabilize the developing negative charge on a pentacoordinate phosphorane transition state.^{11–15} Lys167 serves together with Arg130 as a general acid to expel the 5′-OH leaving group during the cleavage reaction.^{16,17}

Van Duyne and colleagues¹⁸ exploited vanadate as a mimetic to capture a crystal structure of the pentavalent transition state of poxvirus TopIB (Figure 1), which adopts a trigonal bipyramidal geometry with the attacking tyrosine and the OS′ atom of the leaving strand at the apexes, as expected for the proposed associative mechanism.¹¹ This structure confirmed the atomic contacts to the nonbridging and 5′-bridging phosphate oxygens that were surmised from functional studies employing DNA substrates containing stereospecific methylphosphonate or bridging 5′-phosphorothioate substitutions.^{12,13,16,17} The OS′ leaving atom in the transition state is coordinated simultaneously by the Arg130 and Lys167 side

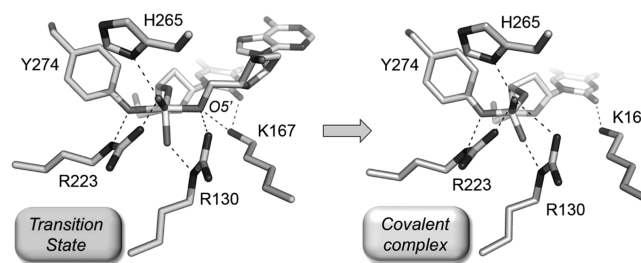


Figure 1. Structural basis of TopIB transesterification chemistry. The active sites of the pentavalent vanadate complex of poxvirus TopIB (a transition state mimetic; Protein Data Bank entry 3IGC; shown at the left) and the covalent tyrosyl-DNA intermediate of poxvirus TopIB (Protein Data Bank entry 2H7F; shown at the right) were superimposed and then offset laterally. The catalytic amino acid side chains are indicated. Atomic contacts are denoted by dashed lines.

chains implicated as general acid catalysts, suggesting a concerted mechanism of leaving group expulsion.

Comparison of the transition state¹⁸ to the covalent intermediate¹⁵ and the apoenzyme¹⁹ of poxvirus TopIB highlights the mobility of the Lys167 side chain, which is located at the tip of a β -hairpin loop. Lys167 in the apoenzyme is located far from its position near the scissile phosphodiester of the transition state and is therefore out of position to perform transesterification chemistry. The act of DNA binding elicits a major conformational switch that brings Lys167 (and

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Eliminate all cysteines in vvTopIB by mutation (C100A-C211A) and verify retention of activity. Change Lys167 to cysteine. (panel A)

Chemical aminoethylation of the cysteine by reaction with 2-bromoethylamine to form γ -thialysine. (panel B)

Install new functional groups at Lys167 by reaction of the K167C proteins with other bromoalkyl compounds. (panel C)

Test for “chemical rescue” of TopIB activity.

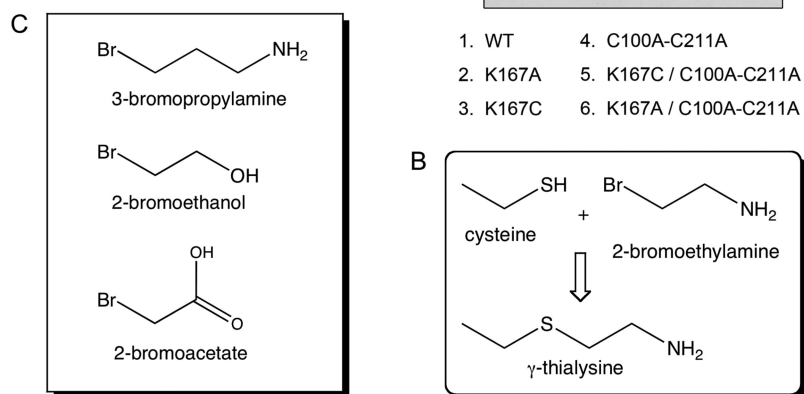


Figure 2. Chemical mutagenesis of vaccinia virus TopIB Lys167. The mutagenesis strategy is outlined. (A) Aliquots (5 μ g) of the indicated phosphocellulose preparations of vaccinia TopIB were analyzed by SDS–PAGE. The Coomassie blue-stained gel is shown. The positions and sizes (kilodaltons) of marker polypeptides are indicated at the left. (B) Chemical reaction of 2-bromoethylamine with cysteine to form γ -thialysine. (C) Chemical structures of other bromoalkyl compounds tested.

also Arg130 and Tyr274) into the active site. After transesterification, Lys167 moves away from the leaving OS' atom and becomes “parked” in the DNA minor groove via its contact with the O2 carbonyl atom of the +1T base (Figure 1).

Traditional mutagenesis methods have illuminated structure–activity relations for the components of the vaccinia TopIB active site.^{6–10,13,16,17,20–22} In the case of Lys167, an alanine substitution causes a 10^{-4} -fold decrement in the rate of transesterification, which is substantially revived when Lys167 is replaced with arginine^{9,16} or when the 5'-bridging oxygen of the scissile phosphodiester is replaced with sulfur.^{16,17} The residual $\sim 10^2$ -fold difference in transesterification rates between wild-type TopIB and the K167R mutant could reflect the predicted 2 log unit difference in the pK_a values for the Lys and Arg side chains.²³ Alternatively, there might be steric constraints on the larger arginine side chain that preclude full restoration of the catalytic power. The genetically programmable protein “tool kit” offers no additional options to instructively modify this functional group at the TopIB active site.

A recent innovation in the analysis of TopIB mechanism is the use of “chemical mutagenesis” to introduce unnatural analogues in lieu of a catalytic amino acid. As implemented by Hecht and colleagues^{24–27} for vaccinia TopIB and human TopIB, this entails in vitro translation of the TopIB protein, from an mRNA with a stop codon in lieu of the codon for the catalytic tyrosine nucleophile, in the presence of a chemically misacylated suppressor tRNA that incorporates a tyrosine analogue. Studies of the effects of *meta*-substituted analogues of Tyr274 on vaccinia TopIB revealed that a *meta*-OH reduced the rate of single-turnover DNA cleavage 130-fold without affecting the rate of single-turnover religation. By contrast,

meta-OCH₃ and NO₂ groups elicited an only 6-fold decrement in cleavage rate. It was proposed that the *meta*-OH uniquely antagonizes deprotonation of the *para*-OH nucleophile during the cleavage transesterification step.²⁶

In this study, we take a different approach to chemical mutagenesis, by producing recombinant vaccinia TopIB with cysteine in lieu of Lys167 and then reacting the K167C protein in vitro with haloalkyl compounds to install lysine-like analogues. The successful chemical reactivation of the K167C mutant provides insights to TopIB general acid catalysis.

EXPERIMENTAL PROCEDURES

Recombinant TopIB. Vaccinia virus TopIB was produced in *Escherichia coli* with a C-terminal His₆ tag. Expression plasmid pET21-TopoHis was constructed as follows. The TopIB open reading frame was amplified by polymerase chain reaction (PCR) with an antisense primer designed to introduce an XhoI restriction site at the translation stop codon while converting it to a serine codon. The PCR product was digested with NdeI (at the translation initiation codon) and XhoI and then inserted into T7-based expression vector pET21b that had been digested with NdeI and XhoI. This maneuver fused the 314-amino acid TopIB polypeptide in frame to nine-amino acid His tag, SLEHHHHHH. Single and compound missense mutations K167A, K167C, C100A/C211A, C100A/C211A/K167A, and C100A/C211A/K167C were introduced into the TopIB gene by two-stage PCR overlap extension. The inserts of the wild-type and mutant pET21-TopoHis plasmids were sequenced to exclude the acquisition of unwanted changes during amplification and cloning. The pET21-TopoHis plasmids were transformed into *E. coli* BL21(DE3). IPTG

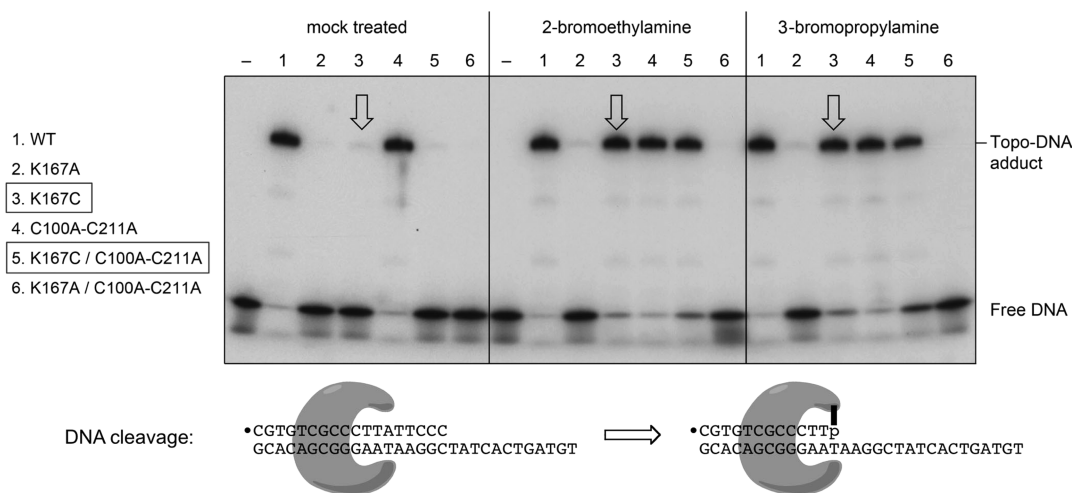


Figure 3. Aminoalkylation of Cys167 restores DNA cleavage activity. The single-turnover cleavage reaction is depicted at the bottom. The 18-mer/30-mer suicide cleavage substrate is shown with the 5' ^{32}P label depicted as a filled circle. The poxvirus TopIB forms a C-shaped protein clamp around the CCCTTpJA recognition site. Transesterification results in the formation of a covalent ^{32}P -labeled pCGTGTGCGCCCTTp-(TopIB) adduct and dissociation of the unlabeled HOATTCCC leaving strand. Cleavage reaction mixtures (20 μL) containing 50 mM Tris-HCl (pH 7.5), 0.3 pmol of 5'- ^{32}P -labeled DNA substrate, and 150 ng (4 pmol) of TopIB as specified (by numbers at the left and above the gel lanes) were incubated at 37 $^{\circ}\text{C}$ for 10 min. The reactions were quenched with 1% SDS. The products were analyzed by SDS-PAGE. An autoradiogram of the dried gel is shown. Free DNA migrated near the dye front. Covalent complex formation was revealed by the transfer of radiolabeled DNA to the TopIB protein. The arrows overlying lanes 3 highlight the successful chemical rescue of K167C DNA cleavage activity by 2-bromoethylamine and 3-bromopropylamine.

induction of TopIB expression and purification of the TopIB proteins from soluble bacterial extracts by sequential Ni-agarose and phosphocellulose chromatography were performed as described previously.²⁶ Protein concentrations were determined by using the Bio-Rad dye reagent with BSA as the standard.

Chemical Modification of Vaccinia TopIB at Cys167.

Reaction mixtures containing (per 20 μL) 100 mM Tris-HCl (pH 8.0), 3 μg of vaccinia TopIB, and 100 mM freshly prepared alkylating agent as specified were incubated at room temperature in the dark for 2 h. The cysteine alkylation reactions were terminated by adding 2-mercaptoethanol to a final concentration of 200 mM. Mock-treated control reactions lacking the alkylating agent were performed in parallel. The treated and mock-treated TopIB proteins were prepared freshly for each experiment shown. The alkylating agents 2-bromoethylamine, 3-bromopropylamine, 2-bromoethanol, and 2-bromoacetate were purchased from Sigma-Aldrich.

Single-Turnover DNA Cleavage. An 18-mer CCCTT-containing DNA oligonucleotide was 5'-end-labeled by enzymatic phosphorylation in the presence of [γ - ^{32}P]ATP and T4 polynucleotide kinase, gel-purified, and hybridized to a complementary 30-mer strand. Cleavage reaction mixtures containing (per 20 μL) 50 mM Tris-HCl (pH 7.5), 0.3 pmol of ^{32}P -labeled 18-mer/30-mer DNA, and 150 ng (4 pmol) of TopIB (mock-treated or treated with alkylating agent) were incubated at 37 $^{\circ}\text{C}$. The cleavage reactions were initiated by adding TopIB to prewarmed reaction mixtures. The reactions were quenched at the times specified by adding SDS to a final concentration of 1%. The samples were analyzed by electrophoresis through a 10% polyacrylamide gel containing 0.1% SDS. Covalent complex formation was revealed by transfer of radiolabeled DNA to the TopIB protein. The extent of covalent complex formation was quantified by scanning the dried gel with a Fujifilm BAS-2500 imager.

RESULTS

Chemical Mutagenesis of Lys167 by Postsynthetic Installation of a Lysine Analogue.

The chemical approach to introducing a lysine-like functional group at a specific site in a recombinant protein exploits the spontaneous reactivity of a cysteine side chain with 2-bromoethylamine to form the lysine analogue γ -thialysine^{28–42} (Figure 2B). The analogue is structurally and chemically similar to lysine; the replacement of a methylene group with a sulfur increases side chain length by ~ 0.3 Å and lowers the pK_a of $\text{N}\zeta$ by ~ 1.1 compared to that of lysine.^{32,39} The chemical modification is quite specific for cysteine at mildly alkaline pH.³⁹ Optimal use of this method to illuminate structure–activity relations at an essential lysine in an enzyme active site entails the following steps (Figure 2): (i) elimination of other cysteine residues in the protein by mutation to alanine, (ii) mutation of the lysine of interest to cysteine, which is now the only cysteine in the polypeptide, (iii) verification that the K-to-C change ablates enzyme activity, (iv) chemical aminoethylation of the cysteine to form γ -thialysine, and (v) interrogation of whether enzymatic activity is restored as a consequence of the cysteine aminoethylation maneuver. An advantage of this approach is that it can be applied to the purified recombinant cysteine-containing enzyme and thereby yield quantities of chemically modified protein for functional studies greater than the amount that can be obtained by in vitro translation.

To execute this agenda, we first generated a cysteine-less C100A-C211A mutant of vaccinia TopIB. Then we replaced Lys167 with alanine or cysteine in the context of the wild-type and C100A-C211A proteins. The wild-type and mutated topoisomerases were produced in *E. coli* with His₆ tags and purified from soluble bacterial extracts by Ni-agarose and phosphocellulose chromatography steps. Sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS–PAGE) analysis verified that the recombinant proteins were of similar high purity with respect to the TopIB polypeptide (Figure 2A).

The purified proteins were reacted for 2 h at 22 °C with 100 mM 2-bromoethylamine (2-BEA) in 100 mM Tris-HCl (pH 8.0). Control “mock-treated” protein samples were incubated in the same buffer without 2-BEA. The 2-BEA and mock reactions were quenched by adjustment to 200 mM 2-mercaptoethanol. The mock-treated and 2-BEA-modified proteins were then tested for DNA transesterification under single-turnover conditions using a “suicide” substrate containing a single CCCTT site. The suicide substrate (Figure 3) consisted of a 5′-³²P-labeled 18-mer scissile strand (5′-pCGTGTGCGCCCT-TATTCCC) annealed to an unlabeled 30-mer strand. The cleavage transesterification reaction of mock-treated wild-type vaccinia TopIB and the cysteine-less C100A/C211A mutant resulted in covalent attachment of the ³²P-labeled 12-mer (5′-pCGTGTGCGCCCTp) to the enzyme and spontaneous dissociation of the unlabeled 6-mer 5′-OH leaving strand from the TopIB–DNA complex. The reaction yielded a discrete SDS-stable protein–DNA adduct that was resolved via SDS–PAGE from the free DNA that migrated near the electrophoretic front (Figure 3, left panel, lanes 1 and 4). Quantification of the protein–DNA adduct and free DNA showed that 90% of the labeled DNA strand was transferred to the mock-treated wild-type and C100A/C211A topoisomerases during a 10 min reaction (Figure 4). The extents of

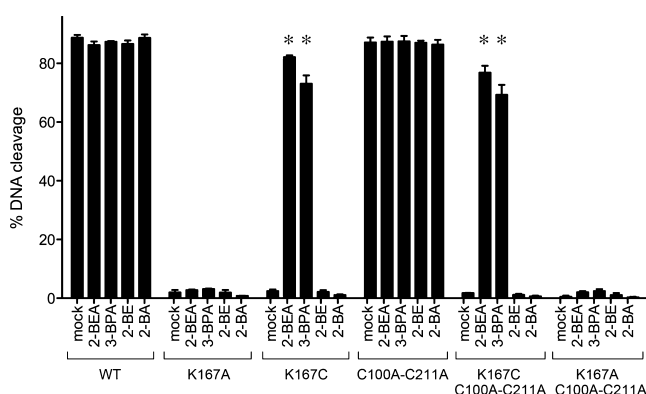


Figure 4. Tests of chemical rescue of K167C by haloalkyl compounds. Single-turnover cleavage by mock-treated and haloalkyl-treated TopIB proteins was performed as described in the legend of Figure 3. The extents of covalent complex formation were quantified by scanning the dried SDS gels with a Fujifilm BAS-2500 imager. Each datum in the bar graph is the average of three independent experiments \pm the standard error of the mean (SEM). The alkylating agents were 2-bromoethylamine (2-BEA), 3-bromopropylamine (3-BPA), 2-bromoethanol (2-BE), and 2-bromoacetate (2-BA).

transesterification by the 2-BEA-treated wild-type and C100A/C211A proteins were the same as the mock-treated controls (Figure 3, middle panel, lanes 1 and 4) (Figure 4).

Changing Lys167 to either alanine or cysteine suppressed DNA cleavage, whether in the context of the otherwise wild-type TopIB (Figure 3, left panel, lanes 2 and 3) or the C100A/C211A mutant (Figure 3, left panel, lanes 5 and 6) (Figure 4, mock). To gauge and compare the impact of the K167A and K167C mutations, we employed a longer duplex DNA substrate containing a single CCCTT site to assay transesterification under equilibrium conditions⁴ (Figure 5). We refer to this DNA as an equilibrium substrate because the 22-mer leaving strand generated upon cleavage at CCCTT remains associated with the topoisomerase DNA complex via base pairing to the nonscissile strand. The K167A mutant displayed

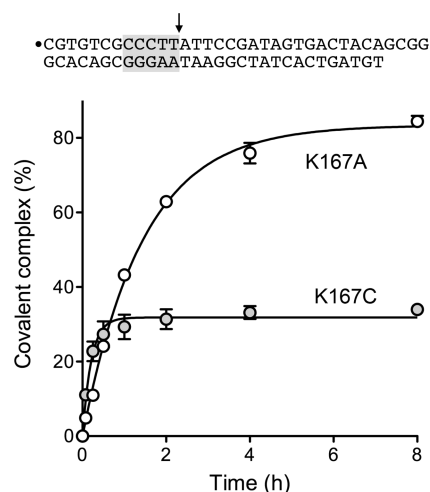


Figure 5. Kinetics of equilibrium DNA cleavage by K167A and K167C. The 34-mer/30-mer equilibrium cleavage substrate is depicted at the top with the 5′ ³²P label depicted as a filled circle, the CCCTT sequence shaded gray, and the cleavage site denoted with the arrow. Reaction mixtures containing (per 20 μ L) 50 mM Tris-HCl (pH 7.5), 0.3 pmol of ³²P-labeled DNA, and 150 ng (4 pmol) of purified K167A or K167C TopIB were incubated at 37 °C. The reactions were initiated by adding TopIB to prewarmed reaction mixtures. Aliquots (20 μ L) were withdrawn at the times specified, and the reactions were quenched immediately with SDS. The quenched reaction mixtures were treated with 10 μ g of proteinase K for 60 min at 37 °C, then mixed with an equal volume of a 95% formamide/20 mM EDTA mixture, and heat-denatured prior to electrophoresis through a 17% polyacrylamide gel containing 7 M urea in TBE (90 mM Tris-borate and 2.5 mM EDTA). The radiolabeled DNAs and the DNA–peptide adducts derived by proteolysis of the covalent TopIB–DNA complex were quantified by scanning the gel. Covalent adduct formation (expressed as the percent of the total labeled DNA) is plotted as a function of reaction time. Each datum is the average of three separate experiments \pm SEM. Nonlinear regression curve fits to a pseudo-first-order exponential function (executed in Prism) are shown.

a slow approach to equilibrium over 4 h (Figure 5) and achieved a much higher extent of covalent adduct formation at equilibrium (84%) than what one sees for wild-type TopIB.⁴ The increase in the cleavage equilibrium constant (K_{eq}) attests that the K167A mutation has relatively less impact on the rate of cleavage (k_{cl}) than on the rate of religation (k_{rel}). A nonlinear regression fit of the data to a pseudo-first-order one-phase association in Prism (Figure 5) yielded a rate constant (k_{obs}) of $1.9 \times 10^{-4} \text{ s}^{-1}$ for the approach to equilibrium. Solving the equations $k_{obs} = k_{cl} + k_{rel}$ and $K_{cl} = k_{cl}/k_{rel}$ yielded the component rate constants ($k_{cl} = 1.6 \times 10^{-4} \text{ s}^{-1}$, and $k_{rel} = 3.2 \times 10^{-5} \text{ s}^{-1}$) for K167A, which are in the same range as those derived previously for K167A with a 60 bp equilibrium cleavage substrate.⁹ The K167C mutant displayed a faster approach to equilibrium ($k_{obs} = 1.3 \times 10^{-3} \text{ s}^{-1}$) and a lower end point cleavage value than K167A (Figure 5). From these data, we derived component rate constants ($k_{cl} = 4.3 \times 10^{-4} \text{ s}^{-1}$, and $k_{rel} = 9 \times 10^{-4} \text{ s}^{-1}$) for K167C. Whereas cysteine in lieu of alanine had a minor restorative effect (~ 2.5 -fold) on forward cleavage, it elicited an ~ 30 -fold enhancement of religation (though K167C remained 1000-fold slower at religation than wild-type vaccinia TopIB).

The instructive finding was that treatment with 2-BEA restored single-turnover DNA cleavage activity to the K167C and C100A/C211A/K167C proteins (Figure 3, middle panel,

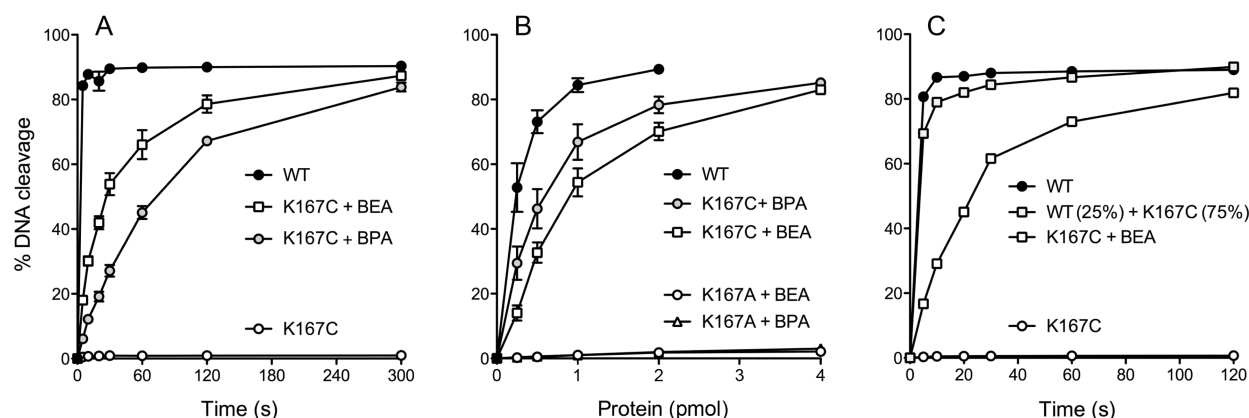


Figure 6. Kinetics of single-turnover cleavage by 2-BEA- and 3-BPA-modified TopIB. (A) Kinetic analysis of single-turnover cleavage of the 18-mer/30-mer substrate was performed as described in Experimental Procedures. The percentage of input DNA cleaved was plotted as a function of reaction time for the indicated TopIB preparations. Each datum in the graph is the average of three independent time course experiments \pm SEM. The cleavage rate constants were derived by nonlinear regression fitting of the data to an exponential one-phase association in Prism. (B) Cleavage reaction mixtures containing 0.3 pmol of 5'- 32 P-labeled 18-mer/30-mer DNA substrate and increasing amounts of TopIB as specified were incubated for 15 min at 37 °C. The percentage of input DNA cleaved was plotted as a function of input TopIB. Each datum in the graph is the average of three independent protein titration experiments \pm SEM. (C) Kinetics of single-turnover cleavage of 0.3 pmol of 5'- 32 P-labeled 18-mer/30-mer DNA substrate by 150 ng (4 pmol) of either pure wild-type TopIB (WT), pure K167C, 2-BEA-treated K167C, or a 25/75 mixture of WT and K167C.

lanes 3 and 5) (Figure 4). By contrast, 2-BEA treatment had no salutary effect on the K167A and C100A/C211A/K167A proteins (Figure 3, middle panel, lanes 2 and 6) (Figure 4). Thus, the chemical rescue maneuver works for TopIB. This experiment told us also that (i) we did not need to convert the two native cysteines of vaccinia TopIB (C100 and C211) to alanine to conduct a successful chemical mutagenesis study of Lys167 and (ii) we did not need to denature the protein with urea^{30–32,37} to chemically modify Cys167, a finding consistent with the location of amino acid 167 on a surface-exposed β -hairpin loop of the TopIB apoenzyme.¹⁹

Test of Other Cysteine Alkylating Agents for the Chemical Rescue of DNA Cleavage Activity. Having achieved a successful chemical rescue with 2-BEA, we sought to modify Cys167 with other haloalkyl compounds (Figure 2B) to create a series of non-native side chains and then evaluate which (if any) restore activity and the extent to which they do so. We applied this strategy with 3-bromopropylamine (3-BPA) and found that (i) 3-BPA treatment had no effect on the DNA cleavage activity of wild-type TopIB or C100A/C211A (Figure 3, right panel, lanes 1 and 4), (ii) 3-BPA treatment rescued the DNA cleavage activity of the K167C and C100A/C211A/K167C topoisomerases (Figure 3, right panel, lanes 3 and 5), and (iii) 3-BPA did not rescue the transesterification defects of the K167A and C100A/C211A/K167A proteins (Figure 3, right panel, lanes 2 and 6) (Figure 4). We conclude that TopIB catalysis can accommodate an unnatural γ -thiahomolysine side chain with a longer alkyl spacer between the main chain and the amino functional group that serves as the general acid.

We also treated the Cys167-containing TopIB proteins with 2-bromoethanol (2-BE) and 2-bromoacetate (2-BA) (Figure 2C) to install hydroxyl and carboxylate functional groups with lysine-like spacers. Whereas treatment with 2-BE or 2-BA did not adversely affect the extent of DNA cleavage by wild-type TopIB and C100A/C211A, it also failed to revive the activity of the K167A and C100A/C211A/K167A proteins (Figure 4). These results attest to the critical role of the alkylamine moiety at position 167 in TopIB catalysis.

Effects of γ -Thialysine and γ -Thiahomolysine on the Rate of DNA Cleavage. We conducted a kinetic analysis of

the suicide cleavage reactions of the 2-BEA- and 3-BPA-modified K167C proteins to gauge the impact of the lysine analogues on TopIB catalysis. Mock-treated K167C and wild-type TopIB served as controls. The TopIB proteins were added in 13-fold molar excess over the 32 P-labeled DNA substrate. The pertinent feature of the suicide cleavage reaction is that the unlabeled 6-mer 5'-OH leaving strand ATTCCC dissociates spontaneously from the covalent TopIB–DNA complex. Loss of the leaving strand drives the reaction toward the covalent state, so that the reaction can be treated kinetically as a first-order unidirectional process. As expected, the K167C protein was virtually unreactive within the 5 min time frame of the experiment (Figure 6A; <1% DNA cleaved). The wild-type TopIB cleavage reaction attained 93% of the end point value at 5 s, the earliest time tested, consistent with a k_d of ~ 0.4 – 0.5 s^{–1} as reported previously.⁴ The 2-BEA- and 3-BPA-modified K167C proteins displayed a steady increase in their levels of covalent adduct formation and attained extents of DNA cleavage at 5 min (87 and 84%, respectively) similar to that of wild-type TopIB (90%) (Figure 6A). Fitting the data in Figure 6A to a single exponential yielded k_d values of 0.031 and 0.013 s^{–1} for the 2-BEA- and 3-BPA-modified K167C proteins, respectively.

Having shown that the yield of the covalent adduct by the chemically modified K167C proteins under single-turnover conditions was comparable to that of wild-type TopIB, we could gauge the extent of Cys167 chemical modification by performing an enzyme titration in the suicide cleavage assay. Figure 6B shows a plot of the yields of the covalent TopIB–DNA adduct in a 15 min DNA cleavage reaction with 0.3 pmol of 32 P-labeled DNA substrate versus the amounts of input wild-type and 2-BEA- and 3-BPA-treated K167C and K167A proteins. As expected, the 2-BEA- and 3-BPA-treated K167A proteins were inactive. By contrast, wild-type TopIB generated near-stoichiometric yields of the covalent adduct, i.e., 0.25 pmol of input TopIB, the lowest level tested, formed 0.15 pmol of covalent protein–DNA complex (Figure 6B). The titration profiles of the 2-BEA- and 3-BPA-treated K167C proteins were shifted to the right compared to that of the wild-type enzyme. By comparing the slopes of titration curves, we estimated that

half of the 3-BPA-treated enzyme was catalytically active (i.e., contained γ -thiahomolysine in lieu of Cys167) and one-fourth of the 2-BEA-treated enzyme was catalytically active (i.e., contained γ -thialysine at position 167).

With this information in hand, we queried whether the presence of significant residual K167C might have affected the kinetics of single-turnover cleavage by the chemically reactivated 2-BEA-modified TopIB containing γ -thialysine. To model this scenario, we premixed wild-type TopIB and the K167C mutant (to achieve 25% wild-type and 75% K167C, thus mimicking the composition of the 2-BEA-treated TopIB). The kinetics of single-turnover DNA cleavage by the WT/K167C mixture are shown in Figure 6C. The presence of the K167C protein had no effect on the cleavage reaction end point but resulted in a slight reduction of the extent of cleavage at the 5 s time point vis-à-vis pure wild-type TopIB. The apparent k_{cl} values of the wild-type TopIB and WT/K167C mixtures were 0.44 and 0.32 s⁻¹, respectively (Figure 6C). [The apparent k_{cl} of a 2-BEA-treated K167C preparation assayed in parallel was 0.043 s⁻¹ (Figure 6C), which is close to the value of 0.031 s⁻¹ derived from the experiments in Figure 6A.] We regard the apparent k_{cl} of the WT/K167C mixture as the upper bound for the full restoration of activity and the k_{cl} of the pure K167C mutant (derived from the experiment in Figure 5) as the baseline activity against which the activities of the treated K167C preparations (comprising mixtures of chemically modified K167C and residual unmodified K167C) can be compared. Accordingly (using the k_{cl} data from Figure 6A), the conversion of Cys167 to γ -thialysine elicited an ~70-fold enhancement in the rate of DNA cleavage and thereby restored the cleavage rate to within a factor of 10 of that of the Lys167-containing enzyme. Modification of the Cys167 to γ -thiahomolysine increased k_{cl} by ~30-fold. The finding that the rate of DNA cleavage by the 2-BEA-modified K167C protein was only 2.4-fold faster than that of the 3-BPA-modified K167C signifies that TopIB is relatively tolerant of the extra methylene substituent of the homolysine side chain during the cleavage transesterification step. A salient point is that γ -thialysine (k_{cl} = 0.031 s⁻¹) and γ -thiahomolysine (k_{cl} = 0.013 s⁻¹) at position 167 promote faster catalysis of DNA cleavage than arginine does, the reported⁹ K167R k_{cl} value being 0.0067 s⁻¹.

Effects of γ -Thialysine and γ -Thiahomolysine on the Rate of DNA Religation. The effects of the lysine 167 analogues on the religation transesterification reaction were studied under single-turnover conditions by assaying the ability of preformed suicide intermediates to transfer the covalently held 5'-³²P-labeled 12-mer strand to a 5'-OH-terminated 18-mer strand to generate a 30-mer product (Figure 7). After the suicide intermediate on the 18-mer/30-mer DNA substrate had been formed, the religation reaction was initiated by adding a 50-fold molar excess of the 18-mer DNA acceptor strand. The sequence of the added 18-mer is fully complementary to the 5'-single-stranded tail of the suicide intermediate. The ionic strength was adjusted simultaneously to 0.5 M NaCl to promote dissociation of the topoisomerase after strand ligation and prevent recleavage of the ³²P-labeled 30-mer strand transfer product. Aliquots were withdrawn immediately prior to the addition of 18-mer and NaCl (defined as time zero) and at various times afterward. The extent of religation at each time point was quantified as the fraction of the ³²P-labeled DNA present as the covalent adduct at time zero that was converted to 30-mer strand transfer product (Figure 7). The religation

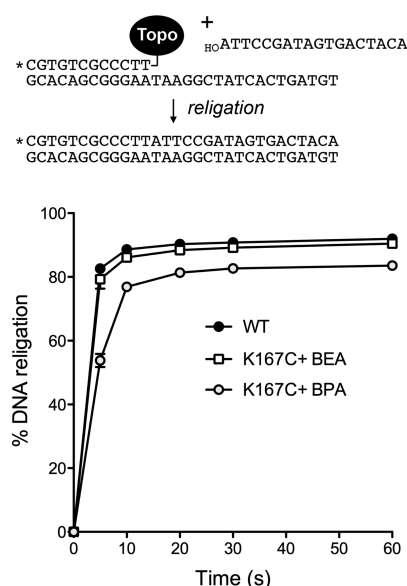


Figure 7. Effects of lysine analogues on DNA religation. Cleavage reaction mixtures containing (per 20 μ L) 50 mM Tris-HCl (pH 7.5), 0.3 pmol of 5'-³²P-labeled 18-mer/30-mer DNA substrate, and 150 ng (4 pmol) of TopIB as specified were incubated at 37 °C for 5 min to form the suicide intermediate depicted at the top. Religation was initiated by the simultaneous addition of NaCl to a concentration of 0.5 M and a 5'-OH 18-mer acceptor strand d(ATTCCGATAGTGACTACA) to a concentration of 15 pmol/22 μ L (a 50-fold molar excess over the input DNA substrate). Aliquots (20 μ L) were withdrawn at the times specified and reactions quenched immediately with 1% SDS. A time zero sample was withdrawn before the addition of the acceptor strand. The samples were digested for 1 h at 37 °C with 10 μ g of proteinase K, then mixed with an equal volume of a 95% formamide/20 mM EDTA mixture, heat-denatured, and analyzed by electrophoresis through a 17% polyacrylamide gel containing 7 M urea in TBE. The radiolabeled DNAs and DNA-peptide adducts (derived by proteolysis of the covalent TopIB-DNA complex) were quantified by scanning the gel. Religation of the covalently bound 12-mer strand to the 18-mer acceptor DNA yielded a 5'-³²P-labeled 30-mer strand transfer product. The extents of religation (expressed as the percent of the covalent intermediate converted into 30-mer) are plotted as a function of reaction time for each TopIB protein. Each datum for the alkylated K167C proteins is the average of three separate experiments \pm SEM. Each datum for wild-type TopIB is the average of five experiments \pm SEM.

reaction of wild-type TopIB was effectively complete by 5 s, the earliest time analyzed, in keeping with prior measurements of k_{rel} of 1.2–1.4 s⁻¹ for vaccinia TopIB.^{9,43} The religation profile of the 2-BEA-modified TopIB(K167C)-DNA intermediate was indistinguishable from that of wild-type TopIB, within the limits of the assay (i.e., $k_{rel} \geq 0.4$ s⁻¹), signifying that the installation of γ -thialysine in lieu of Lys167 had a <3–4-fold effect on the rate of religation. Taking the k_{rel} value of untreated K167C (from the experiment in Figure 5) as a baseline, we found the conversion of Cys167 to γ -thialysine increased the rate of religation by at least 400-fold. The religation profile of the 2-BPA-modified TopIB(K167C)-DNA intermediate was detectably slowed (k_{rel} = 0.22 s⁻¹), indicating that the longer γ -thiahomolysine side chain exerted an only modest (at least 2-fold, but likely <6-fold) effect on the rate of the religation transesterification reaction. Note that γ -thialysine and γ -thiahomolysine at position 167 are at least 1 order of magnitude faster in catalysis of religation than is arginine, the K167R k_{rel} value⁹ being 0.012–0.022 s⁻¹.

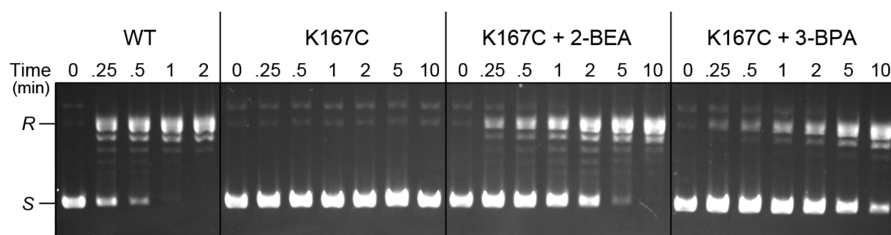


Figure 8. Aminoalkylation of Cys167 restores supercoil relaxation activity. Relaxation reaction mixtures containing (per 20 μ L) 50 mM Tris-HCl (pH 7.5), 100 mM NaCl, 2.5 mM EDTA, 300 ng of supercoiled pUC19 DNA, and 2.25 ng of WT, K167C, 2-BEA-treated K167C, or 3-BPA-treated K167C TopIB as specified were incubated at 37 $^{\circ}$ C. The reactions were initiated by the addition of enzyme. Aliquots (20 μ L) were withdrawn at the times specified and reactions quenched immediately with SDS. The time zero samples were taken prior to the addition of enzyme. The reaction products were analyzed by 1% agarose horizontal gel electrophoresis. The gels were stained with ethidium bromide, and DNA was visualized under UV illumination.

Effects of γ -Thialysine and γ -Thiahomolysine on Supercoil Relaxation. The wild-type, K167C, and amino-alkylated K167C proteins were also assayed for their rates of supercoil relaxation (Figure 8). The K167C protein failed to relax the supercoiled DNA within the 10 min time frame of the experiment, as expected from its feeble DNA cleavage and religation kinetics. Wild-type TopIB relaxed all of the input DNA within 1–2 min. 2-BEA-treated K167C relaxed the supercoils at $\sim 1/10$ the rate of wild-type TopIB, and 2-BPA-treated K167C relaxed them at $\sim 1/20$ of the wild-type rate. After correcting for the fractions of alkylated enzymes in the 2-BEA- and 3-BPA-treated K167C preparations, we estimate that γ -thialysine and γ -thiahomolysine promote supercoil release $1/3$ and $1/10$ as well as the native lysine. The hierarchy of the effects of alkylation on relaxation was concordant with the effects on single-turnover DNA cleavage.

DISCUSSION

Here we successfully applied postsynthetic chemical modification of a defective K167C mutant TopIB to show that lysine analogues γ -thialysine and γ -thiahomolysine rescue the DNA cleavage, DNA religation, and DNA supercoil relaxation activities of vaccinia TopIB. The chemically installed lysine analogues are more effective than a genetically programmed conservative arginine substitution⁹ in promoting the DNA transesterification steps. The modest rate differential between γ -thialysine and γ -thiahomolysine attests to the ability of TopIB to accommodate a longer alkylamino side chain during assembly of the active site and adoption of a transition state in which the lysine or homolysine side chain coordinates the OS' leaving group. The functional evidence for Lys167 catalysis of leaving group expulsion is well established.^{16,17} The incomplete restoration of function by arginine,⁹ and the failure of 2-bromoethanol to revive the cleavage activity of K167C, are consistent with the model of general acid catalysis, whereby the arginine guanidinium ($pK_a = 12.5$) is less amenable to proton donation and an ϵ -OH analogue of lysine (the product of cysteine alkylation by 2-bromoethanol, with a pK_a of ~ 15) is much worse in this respect.

It is not immediately obvious why γ -thialysine, with a pK_a 1.1 units lower than that of lysine,³⁹ would slow the single-turnover cleavage rate by 1 order of magnitude. Assuming that Lys167 has a "consensus" pK_a of 10.5, γ -thialysine 167 would have a pK_a of 9.4 and $N\zeta$ of thialysine would still be fully protonated in the ground state at pH 7.5. However, it is possible that the pK_a of Lys167 is perturbed downward, either in the apoenzyme or in the noncovalent TopIB-DNA complex, so that a further

reduction of 1.1 units for γ -thialysine 167 could result in a significant fraction of the amino moiety being unprotonated and hence unavailable for general acid catalysis of DNA cleavage. Alternatively, the reduced cleavage activity of the γ -thialysine 167 of TopIB could be caused by unfavorable contacts or conformational effects of the larger $S\gamma$ atom, independent of a pK_a effect.

Lys167 in its unprotonated state is presumed, by microscopic reversibility, to function (along with Arg130) in abstracting a proton from the attacking 5'-OH DNA strand during the religation transesterification reaction. This scenario is consistent with the apparently benign effect of γ -thialysine on the rate of religation. The K167A mutation slows the rate religation by more than 4 orders of magnitude; our observation that the K167C enzyme is 3-fold faster at religation than K167A suggests that the cysteine sulfur can function, albeit quite poorly, in lieu of the lysine or γ -thialysine nitrogen as a proton acceptor during religation. The pK_a of cysteine in proteins can vary widely around an average pK_a value of 6.8.²³ If the average value pertains to Cys167 in TopIB, then it would be largely unprotonated under our assay conditions and potentially suitable for function as a general base. It is conceivable that the shorter distance of the cysteine sulfur atom from the main chain (compared to the nitrogen of lysine or homolysine) limits its access to the OS' atom in the TopIB active site (Figure 1) and thereby accounts for the feeble activity of Cys167 in DNA religation compared to Lys167.

AUTHOR INFORMATION

Corresponding Author

*Phone: (212) 639-7145. E-mail: s-shuman@ski.mskcc.org.

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