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A surface plasmon resonance study of the intermolecular interaction between *Escherichia coli* topoisomerase I and pBAD/ Thio supercoiled plasmid DNA

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Abstract

To date, the bacterial DNA topoisomerases are one of the major target biomolecules for the discovery of new antibacterial drugs. DNA topoisomerase regulates the topological state of DNA, which is very important for replication, transcription and recombination. The relaxation of negatively supercoiled DNA is catalyzed by bacterial DNA topoisomerase I (topoI) and this reaction requires Mg^{2+} . In this report, we first quantitatively studied the intermolecular interactions between *Escherichia coli* topoisomerase I (EctopoI) and pBAD/Thio supercoiled plasmid DNA using surface plasmon resonance (SPR) technique. The equilibrium dissociation constant (K_d) for EctopoI-pBAD/Thio interactions is determined to be about 8 nM. We then studied the effect of Mg^{2+} on the catalysis of EctopoI-pBAD/Thio reaction. A slightly higher equilibrium dissociation constant (~15 nM) was obtained for Mg^{2+} coordinated EctopoI ($Mg^{2+}EctopoI$)-pBAD/Thio interactions (~0.043 s⁻¹), compared to EctopoI-pBAD/Thio interactions (~0.043 s⁻¹), compared to EctopoI-pBAD/Thio interactions (~0.017 s⁻¹). These results suggest that enzyme turnover during plasmid DNA relaxation is enhanced due to the presence of Mg^{2+} and furthers the understanding of importance of the Mg^{2+} ion for bacterial topoisomerase I catalytic activity.

Keywords

Surface plasmon resonance; bacterial topoisomerase I; supercoiled plasmid DNA; capture covalent method; equilibrium dissociation constant

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1. Introduction

The drug resistance of bacterial pathogens to available antibacterial drugs is a serious public health issue and needs to be addressed. The bacterial topoisomerase I, a DNA topoisomerase I (topoI), is a novel target biomolecule for the discovery of new antibacterial drugs [1]. DNA topoisomerases play important roles on both the supercoiling control of DNA and the resolution of topological barriers during replication, transcription, and recombination [2,3,4,5]. The supercoiling tension caused by translocation of RNA polymerase must be relieved by topoisomerases [6,7]. Topoisomerase I cleaves and rejoins a single DNA strand during topoI-DNA reactions [8], which establishes a transient covalent linkage between these two macromolecules. These complexes can be trapped using topoisomerase inhibitors [2,8].

Topoisomerase I can catalyze interconversion of various topological isomers [9] and this type IA topoisomerase catalytic activity requires Mg^{2+} [10]. In *Escherichia coli*, EctopoI removes excess negative supercoils in order to regulate DNA supercoiling [11]. EctopoI is a single polypeptide of 865 amino acids and tyrosine 319 is the active site tyrosine in EctopoI that forms a transient covalent linkage to DNA 5' phosphoryl group during EctopoI-DNA reaction [12]. We quantitatively studied the interactions for EctopoI-pBAD/Thio supercoiled plasmid DNA (hereafter termed as pBAD/Thio) and Mg²⁺ bound EctopoI (Mg²⁺EctopoI)-pBAD/Thio using the surface plasmon resonance (SPR) technique.

SPR is a widely used label free technique to determine equilibrium dissociation constants and the kinetics of bio-molecular interactions [13]. The sensor surface modification for the SPR assay was confirmed by using electrochemical impedance spectroscopy (EIS). The equilibrium dissociation constant (K_d) for EctopoI binding to pBAD/Thio was determined to be about 8 nM. A slightly higher K_d (~15 nM) value was obtained for Mg²⁺EctopoI-pBAD/ Thio interactions. In addition, the dissociation rate constants (k_d) for the interactions between the enzymes (EctopoI or Mg²⁺EtopI) and pBAD/Thio were also derived and a larger dissociation rate constant (k_d) was obtained for Mg²⁺EctopoI-pBAD/Thio interactions. These results can help us further understand the important role of Mg²⁺ in the interactions between EctopoI and DNA substrate during catalysis [14].

Mycobacterial tuberculosis topoisomerase I (MttopoI), has a different C-terminal DNA binding domain (CTD) that lacks the three Zn^{2+} binding motifs in the CTD of EctopoI [15]. Binding to pBAD/Thio plasmid DNA by the two enzymes were compared. Under our experimental conditions, weak SPR signals were observed for interactions between MttopoI and pBAD/Thio. Therefore, the K_d value could not be recovered for MttopI-pBAD/Thio interactions.

2. Materials and methods

2.1 Materials

Triethylene glycol mono-11-mercaptoundecyl ether (PEG-thiol), nickel (II) sulfate hexahydrate, sodium, chloride, potassium hexacyanoferrate (III) and ethanolamine HCL were purchased from Sigma-Aldrich, ethanol (200 proof) from Decon Laboratories LLC, 2- $\{2-[2-(1-mercaptoundec-11-yloxy)-ethoxy]-ethoxy)-ethoxy nitrilotriacetic acid (NTA-thiol)$ from ProChimia Surfaces, Poland and potassium ferrocyanide trihydrate from AcrosOrganics. N-hydroxysuccinimide (NHS), N-(3-dimethylaminopropyl)-N'-ethylcarbodiimidehydrochloride (EDC) and GeneJET Plasmid Maxiprep Kit were received from Thermo-Scientific. All other reagents were purchased from VWR international, Randor, PA, USA. $Solutions were prepared using deionized (DI) water (~18 M<math>\Omega$) (Ultra Purelab system, ELGA/Siemens or Milli-Q Direct 8 water system). The polycrystalline gold chips (50 nm

Au over 2.5 nm titanium adhesion layers, coated on 18mm × 18mm cover slip glass slides) were purchased from Platypus Technologies, LLC, Madison, WI and each chip was cut into two halves before further processing.

2.2 Methods

2.2.1 Isolation and purification of Ectopol, Mttopol and pBAD/Thio-

Recombinant EctopoI and MttopoI were expressed and purified as N-terminal His-tagged proteins according to previously reported procedures [16,17]. For the SPR experiments, the EctopoI was dialyzed against HEPES buffer (10 mM Hepes, pH 7.4 containing 100 mM NaCl and 0.005% (V/V) tween 20 in DI water) overnight at 4°C. The MttopoI was dialyzed against HEPES buffer with 2.5% (V/V) glycerol overnight at 4°C. The glycerol was needed to maintain solubility of MttopoI. The pBAD/Thio supercoiled plasmid DNA was purified using Genejet maxiprep kit (Thermo Scientific) according to the manufacturer's protocol.

2.2.2 Sensor preparation and characterization—Gold chips were used to prepare SPR sensors. After rinsing in ethanol for 2–3 minutes, the chips were cleaned by oxygen plasma (Plasma Cleaner PDC-001, Harric Plasma, Ithaca, NY) for 40 s at an oxygen pressure of 500–600 mTorr and RF power of 10.2 W. The chips were then annealed with hydrogen flame for 20 s to reduce the surface roughness [18]. To form a self-assembled monolayer (SAM), the hydrogen flamed chip was immediately immersed in mixed thiol solution (1:9 V/V mixture of 1mM NTA-thiol and 1mM PEG-thiol in ethanol respectively) and incubated overnight. The chip was then copiously rinsed with ethanol and DI water to remove physically adsorbed thiol molecules and dried with argon. The formation of SAM on the cleaned gold surface was confirmed by electrochemical impedance spectroscopy (EIS). A detailed explanation of EIS experiments can be found in supplementary information (section S1). Figure 1a shows the scheme of SAM formation and His-tagged EctopoI/ MttopoI immobilization. The modified chip was then mounted inside SPR flow cell. A detail of the SPR system similar to the system used here can be found in a previous report [19].

2.2.3 Enzyme immobilization and DNA binding—The sensor surface was activated using a 40 mM nickel (II) sulfate solution prepared in DI water for 2 minutes at a flow rate of 50 μ L/min followed by DI water flushing for two minutes. The surface was then equilibrated with the HEPES buffer for 5–10 minutes. His-tagged Ectopol (2 μ M) or Mttopol (2.5 μ M) was immobilized on the activated SAM surface at a flow rate of 50 μ L/min via the capture covalent method [20]. Sequential treatment of the Ectopol or Mttopol immobilized surface was accomplished with a pulsed injection of NHS-EDC solution (25 mM NHS and 100 mM EDC) in DI water followed by 1 M ethanolamine (pH 8.2) in DI water. This treatment allowed us to achieve the capture covalent immobilization. The experiments with various pBAD/Thio concentrations were accomplished by regeneration of the sensor surface using 1M NaCl. For the experiments involving Mg²⁺, 0.5 mM MgCl₂ (in Tris buffer, pH 8.0) was passed over EctopoI immobilized surface for 5 minutes. All the experiments were carried out at 22°C.

2.2.4 Data analysis—Complex non-linear least square (CNLS) fitting algorithm was used for EIS data using an equivalent circuit model (supplementary information, section S1). Equilibrium data analysis methods were used to analyze the SPR data [21]. The simple nonlinear hyperbolic (SNLH) equation (equation 1 below) was used to fit the equilibrium SPR response plotted vs DNA concentrations,

$$R = \frac{R_{max}[DNA]}{K_d + [DNA]} \quad (1)$$

where, R is the equilibrium response, [DNA] is the analyte (pBAD/Thio) concentration, K_d is the equilibrium dissociation constant, and R_{max} is the fitting parameter representing maximum response. The dissociation rate constants were also derived by fitting the SPR dissociation profiles. A detailed explanation of the fitting procedure is given the supplementary information (section S2).

3. Results

3.1 Sensor surface characterization

As shown in Figures 1b&c, the EIS experimental data (symbols) are fitted (continuous lines) using the equivalent circuit (inset, Figure 1c). The parameters obtained from fitting are listed in Table 1. The charge transfer resistance (R_{ct}) for SAM modified gold is significantly higher (~1060 k Ω) compared to bare gold (~0.5 k Ω). This clearly confirms the successful modification of the gold surface. Furthermore, due to the presence of SAM on the gold surface, the interfacial capacitance for modified surface is obviously reduced (313nF) compared to clean gold (1227 nF).

3.2 Enzyme immobilization and pBAD/Thio binding

As explained in the methods section, both EctopoI (Figure 2a) and MttopoI (Figure S3, supplementary information) were immobilized onto the Ni²⁺ chelated NTA surface via the capture covalent method. Figure 2b depicts the sensor surface stability after chemical treatment of the enzyme immobilized surfaces with NHS-EDC followed by ethanolamine. Figure 2c is the representation of pBAD/Thio binding to the enzyme immobilized sensor surfaces, including association, saturation and dissociation of SPR signal. The sensor surface was regenerated using 1 M NaCl solution.

3.3 pBAD/Thio molecular interaction with Ectopol, Mg²⁺Ectopol and Mttopol

The concentration-dependent pBAD/Thio binding to EctopoI and Mg^{2+} EctopoI are shown in Figure 3a and 3b, respectively. Under our experimental conditions, the pBAD/Thio did not show a considerable response change upon its binding to MttopoI (Figure 3c). The SNLH fitting, using equation 1, to the plot of equilibrium response vs concentration of pBAD/Thio (Figure 3d) provides quantitative information of the interaction; more precisely, the K_d can be derived. The K_d value obtained from the fit is about 8 nM for EctopoI-pBAD/Thio interactions and a slightly higher K_d value (~15 nM) for Mg²⁺EctopoI-pBAD/Thio interactions. We could not obtain quantitative information for MttopoI-pBAD/Thio interactions.

3.4 Effect of Mg²⁺on Ectopol-pBAD/Thio molecular interaction

The SPR dissociation signals were fitted using the exponential dissociation rate constant equation (supplementary information, section S2). As shown in Figure 4, stronger dissociation signals were observed for Mg²⁺EctopoI-pBAD/Thio interactions compared to EctopoI-pBAD/Thio interactions. The derived dissociation rate constant (k_d) value for EctopoI-pBAD/Thio interactions is ~0.017 s⁻¹ and for Mg²⁺EctopoI-pBAD/Thio interactions is ~0.043 s⁻¹. Compared to EctopoI-pBAD/Thio interactions, a larger k_d value for Mg²⁺EctopoI-pBAD/Thio interactions suggests that the rate of enzyme turnover following DNA religation during catalysis of EctopoI-pBAD/Thio relaxation reaction is enhanced in the presence of Mg²⁺.

4 Discussion

Recognizing EctopoI as an important target biomolecule for the discovery of new antibacterial drugs [1], EctopoI-plasmid DNA interactions were quantitatively studied for the first time (to the best of our knowledge) using SPR, a surface based affinity technique. In the SPR technique, the sensor surface modification plays a crucial role for the minimization of non-specific binding and reproducibility of SPR signals. We carried out EIS measurements to confirm the gold surface modification via the mixed thiols (supplementary information, section S1). The efficient charge (electron) transfer between the redox couple and the working electrode in solution is basically evaluated by charge transfer resistance (R_{ct}) determination. The R_{ct}, obtained either from fitting or crudely from the semicircular diameter (Figures 1b&c) [22], for Au surface is much smaller compared to Au/SAM surface (Table 1). This clearly confirms the successful sensor surface modification. Our data also show that the interfacial capacitance (C) decreases as the surface thickness increases because of the surface modification (Table 1). The successful immobilization of EctopoI or MttopoI on the modified sensor surface was achieved via capture covalent method. Previously reported result shows that the single His-tag has a weak affinity (~1 μ M) to the Ni²⁺-NTA surface [23]. Even immobilized hexahistidine tagged proteins would dissociate upon continuous buffer flushing, and this phenomenon is not suitable for protein-small molecule interaction studies [20]. The immobilized enzyme surface must therefore be stable. This can be achieved by chemical treatment of the enzyme immobilized surface with sequential flow of NHS-EDC and ethanolamine solutions [20] (explained in methods). 30 µL of each solution was injected as a pulse for 20 seconds. Figure 2a and Figure S3 (supplementary information) show the Ectopol and Mttopol immobilization, respectively, onto the sensor surface via capture covalent immobilization. As shown in Figure 2b, the SPR response was found to be stable for 10 minutes before and after the treatment of Ectopol/Mttopol immobilized surfaces and thus guarantees the sensor surface stability and the successful capture covalent immobilization of the enzymes.

Figures 3a shows the interaction of EctopoI-pBAD/Thio in a concentration dependent manner. The EctopoI-pBAD/Thio binding responses saturated quickly; therefore an equilibrium analysis method could easily produce the K_d value via SNLH fitting (Figure 3d). The K_d value for EctopoI-pBAD/Thio interactions was found to be about 8 nM. A slightly higher K_d value (~15 nM) was obtained for Mg²⁺EctopoI-pBAD/Thio interactions. For the experiments shown in Figures 3a&b, except for 40 nM data, each SPR trace was repeated three times. Due to sample limitation, the 40 nM data was not repeated. After saturated SPR responses were achieved, dissociation SPR profiles were recorded for some experiments. Buffer flushing could dissociate the EctopoI-pBAD/Thio complex considerably. The covalent topoisomerase-DNA complex is transient and reversible in nature [3]. Treatment of the topoisomerase-DNA complex with high salt would promote dissociation of the DNA from enzyme following relegation [24]. After desired SPR signals were recorded, the surface was treated using 1M NaCl (in DI water) via pulsed injection for 2-3 minutes. This could successfully regenerate the surface to the initial (buffer) level and the surface was ready for next pBAD/Thio concentration. Figure 2c shows a typical SPR sensorgram showing association, saturation and dissociation of enzyme-pBAD/Thio interaction followed by regeneration. It should be noted that 10 mM Tris (pH 8.0) buffer was used as running buffer for enzymes-pBAD/Thio interaction experiments. The immobilized enzyme surfaces were equilibrated with the Tris buffer, for approximately 5 minutes, before pBAD/Thio injection. The experiments for each enzyme were performed either on the same chip or on different sensor chips. The variation of chip surfaces resulted in a variation of equilibrium response (lowest to highest) of ~0.6 pixels, on an average.

In order to understand the influence of the topol C-terminal domain (CTD) sequence on the interactions between topoI and supercoiled plasmid DNA, we also studied the interaction between MttopoI and pBAD/Thio using SPR. There are tetracysteine Zn^{2+} binding motifs [25] that follow the N-terminal 67kDa transesterification domain [26]. The three tetracysteine motifs are part of a DNA-binding domain at the C-terminus of Ectopol [27]. Unlike Ectopol [28], Mttopol lacks Zn²⁺ coordination and has evolved to have a different CTD sequence [15]. Under our experimental conditions, we could not see the concentration dependent interaction for MttopoI-pBAD/Thio interactions. As shown in figure 3c, weak SPR signals were observed instead. This result suggests that the tetracysteine Zn^{2+} binding motifs are required for observing the interaction with plasmid DNA with the SPR protocol described here. Mycobacterium smegmatis topoisomerase I (MstopoI) has a CTD similar to Mttopol that also does not coordinate Zn²⁺ [15]. Previous reports indicate that Mstopol CTD interacts with DNA during catalysis [29] and is responsible for sequence specific recognition of duplex DNA by Mstopol [30]. Because of its specific interaction with duplex DNA sequence, the *Mycobacterium topoisomerase* CTD may not be as efficient in promoting high affinity binding to the single-stranded DNA region in supercoiled plasmid DNA as the CTD in Ectopol. Alternatively, the result observed here might be due to the loss of Mttopol activity during protein immobilization. The dialyzed Mttopol sample used for protein immobilization was assayed for relaxation activity and found to be active (Figure S4, supplementary information).

It has been revealed from crystal structure of the 67 kDa N-terminal fragment of Ectopol that there is presence of acidic and basic amino acid residues nearby the active site region [31]. It has also been proposed that Lys-13 and Arg-321 (both basic residues) participate in DNA cleavage [32,33] and three acidic residues Asp-111, ASP-113, and Glu-115 coordinate with Mg^{2+} [34]. We could not detect the concentration dependent Mg^{2+} binding to Ectopol, which might be the limiting case of our SPR systems to resolve the detection of very small molecular weight ions/molecules-protein interactions. We also did not analyze our SPR sensorgrams for association rate constant measurements due to rapid association and fast saturation of the SPR signals. Therefore, to better understand the effect of Mg²⁺ coordination, the SPR dissociation profiles were recorded for the SPR sensorgrams shown in Figures 3a&b. These SPR profiles were fitted to the exponential dissociation rate constant equation. The fitting procedure is explained in detail in the supplementary information (section S2). A larger dissociation rate constant (k_d) (~0.043 s⁻¹) was obtained for Mg²⁺EctopoI-pBAD/Thio compared to EctopoI-pBAD/Thio interactions (~0.017 s⁻¹). This increase in k_d value confirms that Ectopol catalytic activity is enhanced with Mg²⁺. This is consistent with the role of Mg^{2+} to increase the dissociation rate constants for type IB topoisomerase I-DNA interactions, as reported previously [14]. Such analysis for type IA topoisomerase-DNA interactions has not been previously reported.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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Research Highlights

- Study of *E. coli* topoisomerase I interactions with supercoiled DNA quantitatively
- Quantitative analysis for the effect of Mg²⁺ ions on the enzyme-DNA interactions
- Higher dissociation rate constant obtained for Mg²⁺ bound *E. coli* topoisomerase I
- Enzyme turnover during the DNA relaxation would be enhanced in the presence of Mg^{2+}

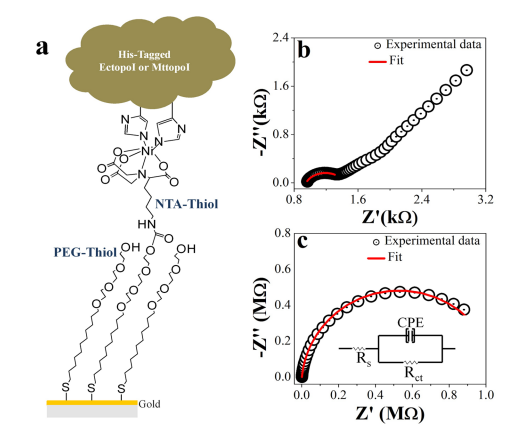


Figure 1.

a) Scheme showing the sensor surface modification with mixed thiols followed by the Histagged EctopoI/MttopoI immobilization. b) Electrochemical Impedance Spectroscopy (EIS) for the bare gold. c) EIS for SAM modified gold surface, inset: equivalent circuit for EIS data fitting and analysis. A frequency range from 10^{-1} Hz to 10^{4} Hz was used during EIS measurements. In both Figures b and c, the symbols are experimental data and continuous lines are the CNLS fit (see supplementary information, section S1, for details).

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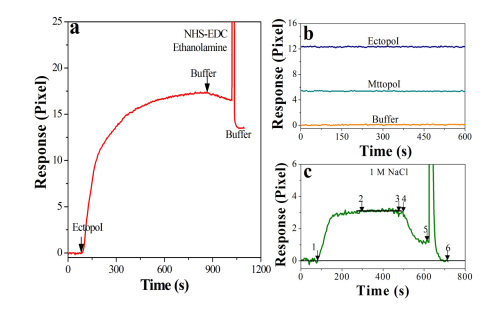


Figure 2.

a) SPR sensorgram for the immobilization of Ectopol via capture covalent method. The arrows show the start of immobilization and buffer wash. b) Representative SPR signals showing stability of sensor surface; with buffer before enzyme immobilization and after capture covalent immobilization of the enzymes. c) Representative SPR profile for enzyme-pBAD/Thio interaction: 1 to 2 association, 2 to 3 saturation, 3 to 4 minor signal shift due to manual changing of valve, 4 to 5 dissociation, and 5 to 6 regeneration.

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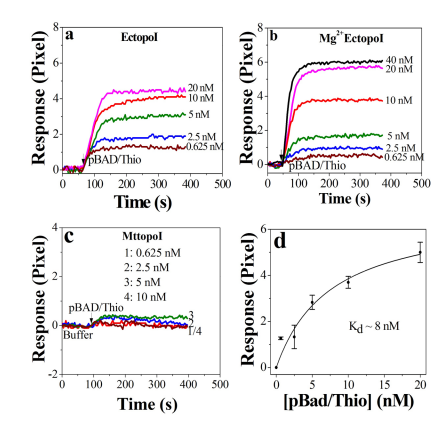


Figure 3.

SPR sensorgrams for a) pBAD/Thio binding to EctopoI, b) for pBAD/Thio binding to Mg^{2+} bound EctopoI and c) for pBAD/Thio binding to MttopoI. d) Plot of equilibrium SPR response vs pBAD/Thio concentration. The symbols are average experimental data of three different measurements with error bars as standard deviation and the continuous lines are the simple hyperbolic fit (explained in text).

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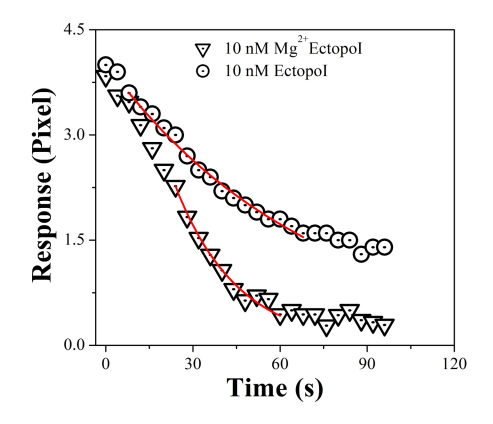


Figure 4.

Representative SPR dissociation profiles fitted to exponential dissociation rate equation (equation S2a, supplementary information). The symbols are experimental data and the continuous lines are the best fit to the rate equation, S2a.

Table 1

The parameters obtained by SNLH fitting of EIS experimental data to an equivalent circuit (Inset, Figure 2c). The errors are the standard errors of the fitting.

Surface	C (nF)	$R_{ct}\left(k\Omega\right)$	$R_{s}\left(k\Omega\right)$	a
Au	1227±122	0.48 ± 0.08	0.95 ± 0.04	0.75±0.01
Au/SAM	313±3	1060±10	0.93±0.08	0.94