Evaluation of Two Models for Human Topoisomerase I Interaction with dsDNA and Camptothecin Derivatives

Gary S. Laco*
Laboratory of Computational and Molecular Biochemistry, Lake Erie College of Osteopathic Medicine, Bradenton, Florida, United States of America

Abstract

Human topoisomerase I (Top1) relaxes supercoiled DNA during cell division. Camptothecin stabilizes Top1/dsDNA covalent complexes which ultimately results in cell death, and this makes Top1 an anti-cancer target. There are two current models for how camptothecin and derivatives bind to Top1/dsDNA covalent complexes (Staker, et al., 2002, Proc Natl Acad Sci USA 99: 15387–15392; and Laco, et al., 2004, Bioorg Med Chem 12: 5225–5235). The interaction energies between bound camptothecin, and derivatives, and Top1/dsDNA in the two models were calculated. The published structure-activity-relationships for camptothecin and derivatives correlated with the interaction energies for camptothecin and derivatives in the Laco et al. model, however, this was not the case for several camptothecin derivatives in the Staker et al. model. By defining the binding orientation of camptothecin and derivatives in the Top1/dsDNA active-site these results allow for the rational design of potentially more efficacious camptothecin derivatives.

Citation: Laco GS (2011) Evaluation of Two Models for Human Topoisomerase I Interaction with dsDNA and Camptothecin Derivatives. PLoS ONE 6(8): e24314. doi:10.1371/journal.pone.0024314

Editor: Rakesh K. Srivastava, The University of Kansas Medical Center, United States of America

Received May 27, 2011; Accepted August 4, 2011; Published August 30, 2011

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Funding: This work was supported solely by an intramural grant from the Lake Erie College of Osteopathic Medicine. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

Competing Interests: The authors have declared that no competing interests exist.

* E-mail: glaco@lecom.edu

Introduction

Camptothecin (CPT) is a natural product that was isolated from the Chinese tree Camptotheca acuminata by Wall and Wani and was shown to inhibit the growth of cancer cells in cell culture [1]. Derivatives of CPT, including topotecan and irinotecan, have been approved by the Food and Drug Administration for the treatment of cancer. Several groups have shown that CPT specifically targets human topoisomerase type IB (Top1) [2,3]. Top1 binds dsDNA and attacks the backbone scissile bond phosphate with the active-site Tyr723 to make a covalent tyrosyl-phosphate bond resulting in religation of the scissile strand DNA backbone, the equilibrium is shifted 7-fold towards the Top1/dsDNA covalent complex [4]. As a result, Top1 remained trapped in covalent complex with the suicide-DNA in the crystals [11]. The Redinbo et al. structure allowed for the structure-based modeling of CPT and derivatives in the Top1/suicide-DNA active site and resulted in models by Redinbo et al. [11], Kerrigan et al. [14], and Laco et al. [15], the last of which is described here as the Rotated +1 Nucleoside model.

Of these structure-based Top1/dsDNA/inhibitor models, only the orientation of CPT and derivatives in the Rotated +1 Nucleoside model were experimentally tested in vitro using both a Top1 Asn352Ala mutant and a dsDNA oligonucleotide assay which used cobalt to oxidize extra helical guanines [15,16]. In the Laco et al. Rotated +1 Nucleoside model, the +1 nucleoside rotated out of the helix when Top1 was in covalent complex with supercoiled dsDNA. As a result of the rotation, the free 3'-OH of the +1 nucleoside was out of range (7.7 Å) of the tyrosyl-phosphate bond preventing religation of the DNA backbone [15]. This finding supports the processive nature of Top1 in that it stays bound to supercoiled dsDNA until the dsDNA is fully relaxed [17]. After the dsDNA is fully relaxed, the +1 nucleoside is rotated back with a suicide-DNA [11]. Since Top1 has a low affinity for relaxed dsDNA, a suicide dsDNA oligonucleotide (suicide-DNA) was used to generate the Top1/suicide-DNA covalent complexes [11]. The suicide-DNA differed from native dsDNA in that the backbone scissile bond 5'-oxygen (O5') was replaced with a sulfur. When the Top1 active-site Tyr723 attacked the suicide-DNA it generated a native tyrosyl-phosphate bond to the 3’-end of the -1 nucleotide while leaving a non-native 5’-SH on the free end of the +1 nucleoside [11,12]. And while the 5'-SH can attack the tyrosyl-phosphate bond resulting in religation of the scissile strand DNA backbone, the equilibrium is shifted ~7-fold towards the Top1/dsDNA covalent complex [13]. As a result, Top1 remained trapped in covalent complex with the suicide-DNA in the crystals [11]. The Redinbo et al. structure allowed for the structure-based modeling of CPT and derivatives in the Top1/suicide-DNA active site and resulted in models by Redinbo et al. [11], Kerrigan et al. [14], and Laco et al. [15], the last of which is described here as the Rotated +1 Nucleoside model.

A key result in the Top1 field was obtained when Redinbo et al. solved the X-ray crystal structure of Top1 in covalent complex +1 deoxyribonucleoside (1 deoxyribonucleoside). This results in a phosphate with the active-site Tyr723 to make a covalent tyrosyl-phosphate bond preventing religation of the DNA backbone [15]. This finding supports the processive nature of Top1 in that it stays bound to supercoiled dsDNA until the dsDNA is fully relaxed [17]. After the dsDNA is fully relaxed, the +1 nucleoside is rotated back...
into the helix allowing the +1 nucleoside 5’OH to attack the tyrosyl-phosphate bond, religate the DNA backbone, and allow Top1 to dissociate from the fully relaxed dsDNA [15]. Rotation of the +1 nucleoside out of the helix leaves a cavity in the Top1/dsDNA active-site which can be occupied by CPT and derivatives. When CPT and derivatives bind in the Rotated +1 Nucleoside model Top1/dsDNA active-site they block the +1 nucleoside from re-entering the helix thus stabilizing the Top1/dsDNA covalent complex [15].

The Rotated +1 Nucleoside model is significantly different from the Staker et al. X-ray crystallography based models of Top1 in covalent complex with suicide-DNA and either topotecan [18] or CPT [19]. In both Staker et al. structure based models the +1 nucleoside was base-paired in the helix, and the inhibitors intercalated in between the -1 and +1 base pairs, forcing the +1 base pairs ~3.4 Å further away from the -1 base pairs [18,19]. As a result, the suicide-DNA +1 nucleoside 5’S’H was 11.6 Å away from the tyrosyl-phosphate bond preventing religation of the DNA [18,19]. The Staker et al. structure-based model of Top1 in covalent complex with dsDNA and bound inhibitor is described here as the “Intercalated model”. The Rotated +1 Nucleoside model and the Intercalated model were evaluated in silico using Discovery Studio computational chemistry software (Accelrys, San Diego, CA). This approach allowed for evaluation of the binding orientations of CPT and derivatives in the Rotated +1 Nucleoside and Intercalated models, to see which binding orientation correlated best with the reported in vitro structure-activity-relationships for CPT and derivatives.

Results and Discussion

CPT and derivatives selected for this study, along with the respective in vitro inhibition results for each, are shown in Figure 1 and Table 1 [15,16,20]. There were three selection criteria for the inhibitors used in this study: 1) inhibitors had been tested in an in vitro Top1 dsDNA oligonucleotide assay in which the dsDNA oligonucleotide had a single high affinity Top1 cleavage site; 2) the assay measured the accumulation of Top1/dsDNA covalent complexes over time; and 3) inhibitors were tested at three or more concentrations to quantitate inhibition. As a result, topotecan was not included in the study due to the lack of published reports documenting the in vitro inhibition activity of topotecan using the same assay and conditions as for the inhibitors in Table 1 [21,22]. The original Laco et al. model of Top1 in covalent complex with DNA and inhibitor [15] was based on the Redinbo et al. X-ray crystal structure of Top1 in covalent complex with suicide-DNA [11]. A reconstituted Top1 was used to generate the Redinbo et al. Top1/suicide-DNA covalent-complex structure and the following Top1 residues were not resolved in the functional domain which includes the DNA binding and enzymatic domains of Top1 (aa 201–265) [17]: residues 201–214; core residues 627–634, linker residues 635-697; and C-terminal residues 698–719 which are part of the Top1 active-site [11]. A more recent structure by Staker et al. of a Top1/suicide-DNA/topotecan ternary complex resolved all main-chain Top1 atoms in the functional domain [18]. However, in that structure the intercalated topotecan, a CPT derivative with an A-ring 9-CH2N(CH3)2 and 10-OH, displaced the +1 base pairs ~3.4 Å further away from the cleavage-site -1 base pairs. In order to generate a structurally complete model of Top1 in covalent-complex with native dsDNA and no inhibitor the displaced DNA strands were replaced in the Staker et al. Top1/suicide-DNA/topotecan structure [18] with the corresponding DNA strands from the Redinbo et al. Top1/suicide DNA structure [11] using Discovery Studio and the Consistent Force Field (CFF) which has parameters for protein, DNA, and small ligands (Methods). Next, the +1 nucleoside was rotated out of the helix until it was trapped in a network of H-bonds/electrostatic interactions with Arg488, Asp533, Arg590 as previously reported [15]. CPT was then docked into the active-site of the Rotated +1 Nucleoside model and modified into the indicated CPT derivatives (Figure 1 and Table 1); with 10-OH CPT shown in the Rotated +1 Nucleoside model active-site in Figure 2. The orientations of CPT and derivatives in this structurally complete Rotated +1 Nucleoside model were similar to the Laco et al. model [15] that was based on the incomplete Top1/suicide-DNA structure by Redinbo et al. [11]. In addition, the suicide-DNA backbone sulfur, that is found as a free S’SH on the +1 nucleoside, was changed to the native oxygen [11,18,19]. Explicit waters were added to the Rotated +1 Nucleoside model presented here followed by minimization, however, individual waters did not make any bridging H-bonds/electrostatic interactions between 10-OH CPT and the Top1/dsDNA active-site (data not shown). This result is consistent with previously published results on the Rotated +1 Nucleoside model in which individual explicit waters did not make bridging H-bond interactions between ligands and the Top1/dsDNA active-site [15]. As a result, all Rotated +1 Nucleoside model Top1/dsDNA/inhibitor complexes were minimized with an implicit solvent model (Methods).

The Staker et al. structure based Intercalated model of Top1 in covalent complex with suicide-DNA and topotecan was modified

| Ligand               | Position | Inhibition |
|----------------------|----------|------------|
| CPT                  | H        | H          | OH        | 0.9*     |
| hCPT                 | H        | H          | OH        | 1.0*     |
| ± 20-deoxy CPT       | H        | H          | H         | 0.03**   |
| ± 20-CI CPT          | H        | H          | CI        | 0.29**   |
| 10-OH CPT            | OH       | H          | OH        | 3.5*     |
| 10,11-diF hCPT       | F        | F          | OH        | 12.0*    |

Table 1. Relative in vitro inhibition of Top1 by CPT, hCPT, and derivatives.

**As reported by Wang et al. for racemic CPT derivatives [20]. All Top1 inhibition assays were performed using an end-labeled dsDNA oligonucleotide.

doi:10.1371/journal.pone.0024314.g001

Figure 1. Structure of CPT and hCPT E-ring. CPT left, hCPT E-ring right; hCPT and derivatives differ from CPT in that they contain an additional E-ring carbon between the 20-OH and the adjacent carbonyl oxygen to give a seven-member E-ring.

doi:10.1371/journal.pone.0024314.g001
[18], topotecan was converted into CPT and derivatives including 10-OH CPT (CPK rendering; carbon, green; oxygen, red; nitrogen, blue; hydrogen, white). In the Staker et al. structure of topotecan in ternary complex with Top1 and suicide-DNA two crystallographic waters were found to make bridging H-bonds between the topotecan carboxylate and Top1 active-site residues [18], however, carboxylate forms of CPT derivatives have been shown to be inactive [15]. One additional water was found to be within 2.9 Å of the topotecan A-ring 10-OH, but there were no Top1/dsDNA H-bond donor/acceptor residues within 4.4 Å of that water [18]. Here the Intercalated model was initially solvated with explicit waters followed by minimization, and a water was found close to the 10-OH of 10-OH CPT, however, the only other H-bond donors/acceptors within 4.4 Å were other waters (data not shown). Since no individual waters were found to make bridging H-bond/electrostatic interactions between the inhibitor and the Top1/dsDNA active-site, all Intercalated model Top1/dsDNA/inhibitor complexes were minimized with an implicit solvent model (Methods). 10-OH CPT is shown in the Intercalated model Top1/dsDNA active-site in Figure 3. The Top1/suicide-DNA/topotecan X-ray crystallography based model was used here because Top1 in the Top1/suicide-DNA/CPT model was not structurally complete [19].

The inclusion of explicit water molecules during an interaction energy calculation between an inhibitor and Top1/dsDNA would artificially make the electrostatic component of the interaction energy score more negative (i.e., stronger) when individual waters...
do not mediate significant bridging H-bonds/electrostatic interactions between the inhibitor and the Top1/dsDNA active-site.

**Similarities and differences between Rotated +1 Nucleoside and Intercalated Models**

While the orientation of 10-OH CPT and DNA in the two models is overall strikingly different, there are similarities and they are described first. Top1 active-site residues Arg364, Lys532, and Arg533 are close to the E-ring in both models, while Asn722 faces the D-ring carbonyl oxygen in both models. In addition, the -1 scissile strand thymine base stacks under the D-ring in both models (Fig. 2 and 3).

In terms of the differences between the two models: 1) In the Rotated +1 Nucleoside model the A-ring 10-OH makes an electrostatic interaction with Asn352 (Fig. 2B and 4A), while in the Intercalated model the A-ring 10-OH H-bonds to Glu356 (Fig. 3B and 4B); 2) In the Rotated +1 Nucleoside model 10-OH CPT only stacks with the -1 scissile strand T (Fig. 2), while in the Intercalated model 10-OH CPT stacks with the -1 and +1 base pairs (Fig. 3). And while Asn430, Pro431, Leu721, and Lys751 are within 4–7.7 Å of 10-OH CPT in the Rotated +1 Nucleoside model, those residues are 1–4 Å further away from 10-OH CPT in the Intercalated model (data not shown). By determining the orientation of CPT and derivatives in the Top1 active-site, CPT derivat could be designed to make additional H-bonds and van der Waals interactions with one or more of the above residues. The default Discovery Studio 2.1 definition for H-bonds was used with H-bonds distances reported here between 3.0–3.1 Å (Accelrys, San Diego, CA).

In order to evaluate the orientation of CPT and derivatives in the two models, and compare them to the reported in vitro activities for those same compounds (Figure 1 and Table 1), the interaction energy scores between CPT and derivatives and the active-sites of the Rotated +1 Nucleoside model (Fig. 2 and 4A) and the Intercalated model (Fig. 3 and 4B) were calculated (Fig. 5). For both models, the CPT interaction energy score (Rotated +1 Nucleoside model: vdW -48.6, E -4.67; Intercalated model: vdW -69.22, E -5.9) was subtracted from that for 10-OH CPT, 20-deoxy CPT, and 20-Cl CPT, while the hCPT interaction energy score

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**Figure 4. Flattened images of two models for Top1 interaction with dsDNA and 10-OH CPT.** A) Flat image of Rotated +1 Nucleoside model from Fig. 2. H-bonds and electrostatic interactions within 3.6 Å are indicated with dashed lines. B) Flat image of Intercalated model from Fig. 3. H-bonds and electrostatic interactions within 3.6 Å are indicated with dashed lines.

doi:10.1371/journal.pone.0024314.g004

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**Figure 5. Relative interaction energy in kcal/mol between CPT/ hCPT-derivatives and the Top1/dsDNA active-site.** The interaction energy scores for the derivatives of CPT and hCPT were subtracted from the score of the respective parent inhibitor with the resulting difference values plotted on the graph. The CPT and hCPT interaction energy scores were set to zero. A negative kcal/mol score indicates that a derivative bound tighter than the parent inhibitor, while a positive kcal/mol score indicates that it bound weaker than the parent inhibitor. A) Interaction energy values for CPT, hCPT and derivatives when bound in the Rotated +1 Nucleoside model Top1/dsDNA active-site. B) Interaction energy values for CPT, hCPT and derivatives bound in the Intercalated model Top1/dsDNA active-site.

doi:10.1371/journal.pone.0024314.g005
Ligand scores for Rotated +1 Nucleoside model

In the Rotated +1 Nucleoside model, the 10-OH CPT value was -3.23 kcal/mol (vdW -2.16, E -1.07) stronger than CPT, and the 10,11-diF hCPT value was -2.03 kcal/mol (vdW -0.16, E -1.87) (Fig. 5A) stronger than hCPT due to either the A-ring 10-OH, or 10,11-diF, interacting with Asn352 (Fig. 2B and 4A). These additional interactions explained why 10-OH CPT and 10,11-diF hCPT were more potent in vitro than CPT and hCPT, respectively (Table 1) [15,16]. The 20-deoxy CPT value was 1.81 kcal/mol (vdW 1.5, E 0.31) weaker than CPT (Fig. 5A) due to the loss of the 20-OH in vitro interaction with the Top1/dsDNA active-site (Fig. 5A). This is consistent with the racemic 20-deoxy CPT having low in vitro activity (Table 1) [20], with the 1.81 kcal/mol value (vdW 1.5, E 0.31) representing a loss of 6% of the total electrostatic interaction energy for CPT. Interestingly, the 20-Cl CPT value was only 0.49 kcal/mol (vdW 0.2, E 0.29) weaker than CPT (Fig. 5A). This result for 20-Cl CPT is consistent with the lack of a H-bond between the E-ring 20-Cl and the -1 scissile strand thymidine carbonyl oxygen (see E-ring 20-OH hydrogen in Fig. 2B and 4A). However, the 20-Cl CPT is significantly more potent in vitro than 20-deoxy CPT (Table 1), indicating that the 20-Cl is contributing to interactions between 20-Cl CPT and Top1/dsDNA through electrostatic interactions with Arg364 and Lys532. In the Rotated +1 Nucleoside model the Arg364 side-chain nitrogens make H-bonds to Asp533 and the -1 non-scissile strand cytosine and the +1 non-scissile strand adenine, and electrostatic interactions with the E-ring 20-OH (Fig. 4A). These interactions by Arg364 would contribute to the stabilization of the Top1/dsDNA/inhibitor complex (Fig. 2 and 4A). When Arg364 was mutated to His, the mutant Top1 became resistant to CPT and derivatives [23]. The shorter His side chain cannot make H-bonds to Asp533 and the -1/+1 non-scissile strand bases, nor electrostatic interactions with the E-ring 20-OH (data not shown). These in silico results support the binding orientation for CPT, hCPT, and derivatives in the Rotated +1 Nucleoside model (Fig. 2).

However, it is important to note that while the 10,11-diF hCPT in vitro inhibition (Table 1, 12-fold) follows the respective Rotated +1 Nucleoside model difference value of -2.03 kcal/mol (Fig. 5A), the two values do not correlate in magnitude with the corresponding values for 10-OH CPT (Table 1, 3.5-fold; and Fig. 5A, -3.23). Fluorinated compounds have increased hydrophobicity [24], which in vitro likely results in 10,11-diF hCPT partitioning more efficiently out of the solvent phase and into the Top1/dsDNA binding cavity than does 10-OH CPT which is more hydrophilic due to the A-ring 10-OH. However, the interaction energy calculations in this study did not take that into account. Consistent with the reported 12-fold inhibition enhancement for 10,11-diF hCPT (Table 1), an inhibitor of glycogen phosphorylase, in which two hydroxyls were replaced with fluorines, was 10-fold more potent than the non-fluorinated inhibitor [25]. And while the addition of a hydroxyl to an inhibitor can be used to probe the active-site of an enzyme for a H-bond donor/acceptor, the substitution of a hydroxyl on an inhibitor with fluorine can be used to probe the active-site of an enzyme for the presence of just a H-bond donor [24,25]. The in vitro data in Table 1 indicates that 10-OH CPT, 10,11-diF hCPT, and 20-Cl CPT interact with H-bond donors in the Top1 active-site. The Rotated +1 Nucleoside model supports this conclusion since Asn352 can donate H-bond/electrostatic interactions to 10-OH CPT and 10,11-diF hCPT (Fig. 2B and 4A), and Arg364 and Lys532 which are H-bond donors can make electrostatic interactions with 20-Cl CPT at ~4 Å (data not shown).

Ligand scores for Intercalated model

In the Intercalated model, the 10-OH CPT value of -3.31 kcal/mol (vdW -2.06, E -1.25) (Fig. 5B) was stronger than CPT due to the 10-OH CPT A-ring 10-OH making a H-bond to Glu356 (Fig. 3B and 4B). In the Staker et al. structure based Top1/suicide-DNA/TopoI model, the Glu356 side-chain oxygen was 3.8 Å from the topocean A-ring 10-OH with a reported B-factor of 48 [18]. The topocean A-ring 9-CH2-N-(CH3)2 may have disrupted the interaction between Glu356 and the adjacent A-ring 10-OH. In contrast, the 10,11-diF hCPT value of -0.2 kcal/mol (vdW -0.34, E 0.13) was only slightly stronger than hCPT (Fig. 5B), due to an electrostatic clash between the A-ring 10,11-diF and the Glu356 side-chain oxygens (note positive E value in the above score). These in silico results for 10,11-diF hCPT appear inconsistent with the reported in vitro 12-fold increase in inhibition (Table 1) [15]. As pointed out above, fluorine has been used to probe the active-site of enzymes for H-bond donors, however, the Glu356 side-chain oxygens can not donate a H-bond to fluorine. In the Intercalated model, the 20-deoxy CPT value was 6.03 kcal/mol (vdW 1.91, E 4.12) weaker than CPT with the electrostatic component representing a loss of 70% of the total electrostatic interaction energy for CPT (see above, Fig. 5B). The decrease in electrostatics for 20-deoxy CPT can be explained by the lack of a H-bond between the 20-deoxy CPT E-ring and Asp533; the racemic 20-deoxy CPT had low in vitro activity (Table 1) [20]. The 20-Cl CPT value of 5.64 kcal/mol (vdW 0.74, E 4.9) was also significantly weaker than CPT due again to the lack of a H-bond between the 20-Cl CPT E-ring and the Asp533 side-chain oxygen (see 20-OH group of 10-OH CPT in Fig. 3B and 4B). In the Intercalated model, the 20-OH H-bond to Asp533 is the dominant electrostatic interaction between the inhibitor and the Top1/dsDNA active-site. However, in the Intercalated model the 20-Cl makes an electrostatic clash with Asp533 (see 20-OH of 10-OH CPT in Fig. 3B and 4B); The in silico results for 10,11-diF hCPT and 20-Cl CPT in the Intercalated model (Fig. 5B) do not follow the trend for the known in vitro inhibition by these CPT/hCPT derivatives (Table 1). It is important to note that these in silico results allow for further testing of both models. For the Intercalated model to be biologically relevant (Fig. 3 and 4B) mutation of Top1 Glu356 to Ala should result in a Top1 mutant that could not distinguish between CPT and 10-OH CPT due to the lack of a H-bond between the Ala356 and the A-ring 10-OH. In contrast, for the Rotated +1 Nucleoside model, the same Top1 Glu356Ala mutant should still be inhibited ~3.5-fold more by 10-OH CPT, than by CPT, because in the Rotated +1 Nucleoside model the 10-OH CPT A-ring does not interact with Glu356 (Fig. 2B).

For this study, interactions between Top1 residues Glu356 and Asp533 and CPT derivatives were critical in showing that the Rotated +1 Nucleoside model is biologically relevant (Fig. 3 and 4B). The A-ring 10-OH making a H-bond to Glu356 (Fig. 3B and 4B). In the Intercalated model, the 20-deoxy CPT value was 6.03 kcal/mol (vdW 1.91, E 4.12) weaker than CPT with the electrostatic component representing a loss of 70% of the total electrostatic interaction energy for CPT (see above, Fig. 5B). The decrease in electrostatics for 20-deoxy CPT can be explained by the lack of a H-bond between the 20-deoxy CPT E-ring and Asp533; the racemic 20-deoxy CPT had low in vitro activity (Table 1) [20]. The 20-Cl CPT value of 5.64 kcal/mol (vdW 0.74, E 4.9) was also significantly weaker than CPT due again to the lack of a H-bond between the 20-Cl CPT E-ring and the Asp533 side-chain oxygen (see 20-OH group of 10-OH CPT in Fig. 3B and 4B). In the Intercalated model, the 20-OH H-bond to Asp533 is the dominant electrostatic interaction between the inhibitor and the Top1/dsDNA active-site. However, in the Intercalated model the 20-Cl makes an electrostatic clash with Asp533 (see 20-OH of 10-OH CPT in Fig. 3B and 4B); The in silico results for 10,11-diF hCPT and 20-Cl CPT in the Intercalated model (Fig. 5B) do not follow the trend for the known in vitro inhibition by these CPT/hCPT derivatives (Table 1). It is important to note that these in silico results allow for further testing of both models. For the Intercalated model to be biologically relevant (Fig. 3 and 4B) mutation of Top1 Glu356 to Ala should result in a Top1 mutant that could not distinguish between CPT and 10-OH CPT due to the lack of a H-bond between the Ala356 and the A-ring 10-OH. In contrast, for the Rotated +1 Nucleoside model, the same Top1 Glu356Ala mutant should still be inhibited ~3.5-fold more by 10-OH CPT, than by CPT, because in the Rotated +1 Nucleoside model the 10-OH CPT A-ring does not interact with Glu356 (Fig. 2B).
hCPT, respectively (Table 1 and Fig. 5B). The results for the Intercalated model suggest that the use of suicide-DNA, and high inhibitor-to-enzyme ratios (~34:1) to generate Top1/suicide-DNA/inhibitor ternary complexes resulted in non-biologically relevant binding modes for CPT and derivatives [10,19]. This is supported by the fact that the Top1/suicide-DNA/topotecan active-site was found to contain both the lactone (63%) and carboxylate (37%) forms of topotecan (the same ratio was also found for topotecan in mother liquor alone) [18]. The Top1/suicide-DNA/CPT active-site was also found to contain both the lactone and carboxylate forms of CPT, for which the percentages were so close they could not be differentiated [19]. In contrast, Top1 and dsDNA oligonucleotides were reported in vitro to preferentially stabilize the active lactone form of CPT (97.1%) versus the inactive carboxylate form (2.9%) [15]. The carboxylate form of hCPT, which can not reverse to the lactone form, has been shown to have no detectable inhibitory activity in vitro [15]. It can be argued that the Staker et al. Top1/suicide-DNA active-site could not distinguish between active and inactive forms of CPT and derivatives because the +1 nucleoside was not rotated out of the helix to open a biologically important inhibitor binding cavity (Fig. 2) [15]. The Rotated +1 Nucleoside model H-bond/electrostatic interactions between Arg364 and -1 non-scissile strand A and +1 non-scissile strand C, Arg488 and +1 scissile strand G, Lys532 and -1 scissile strand T, and Arg590 and +1 scissile strand G (Fig. 2B and 4A) are consistent with CPT stabilized Top1/dsDNA covalent-complexes occurring preferentially at sequences containing a -1 scissile strand T/+1 scissile strand G and followed by other combinations of -1 pyrimidines and +1 purines/pyrimidines [26].

Here it was also found that the lactone forms of CPT and derivatives made direct H-bonds to the Top1/dsDNA active-site in both the Rotated +1 Nucleoside and Intercalated models which is consistent with: 1) a previous report for a solvated Rotated +1 Nucleoside model [15]; 2) for the X-ray crystallography based Intercalated model [18]; and 3) molecular dynamic simulations of the topotecan lactone in the solvated Top1/dsDNA active-site by Mancini et al. [27].

The probing of the Top1 active-site with CPT derivatives containing chlorine and fluoride substitutions for hydroxyls revealed that 20-Cl CPT was able to make an electrostatic interaction with the Top1/dsDNA active-site, and 10,11-dF hCPT interacted with active-site H-bond donors near the A-ring. Only the binding of these inhibitors in the Rotated +1 Nucleoside model allows for 20-Cl CPT to maintain electrostatic interactions with Top1/dsDNA via Arg364 and Lys532, and for Asn352 to interact with fluorines on 10,11-dF hCPT (see 10-OH CPT in Fig. 2B and 4A). Together these findings further define the orientation of CPT and derivatives in the Top1 active-site, and allow for the rational design of CPT derivatives that make additional interactions with the Top1 active-site residues, including Asn430, Pro431, Lecu721, and Lys751. This approach, in combination with the rational design of CPT derivatives which avoid cancer cell drug resistance mechanisms [9,10], has the potential to result in more efficacious CPT derivatives for the treatment of drug resistant cancers and cancers that are currently not treated with CPT derivatives.

**Methods**

**Structure based Top1/dsDNA Rotated +1 Nucleoside model**

All Top1/dsDNA and inhibitor modifications, minimizations, and calculations, were carried out using Discovery Studio 2.0 with conditions and parameters that were optimized to mimic general in vitro conditions for direct interactions between ligands and an enzyme active site (Accelrys, San Diego, CA). 1) In the Redelino et al. IA31.pdb file [11], all waters were deleted, and all 5-iodo-2'-uridines in the DNA were replaced with thymidines. In the Staker et al. 1K4T.pdb file [10], topotecan (lactone and carboxylate forms), mercury, PEG, and all waters were deleted. 2) All incomplete Top1 side-chains were built out in the IA31.pdb and 1K4T.pdb structures. 3) The IA31.pdb Top1/suicide-DNA structure was superimposed with the 1K4T.pdb Top1/suicide-DNA/topotecan structure using selected atoms in the respective enzymes and bound dsDNAs. The aligned non-scissile strand nucleotides 115–122, and scissile strand nucleotides 1–10 in the IA31.pdb and 1K4T.pdb structures had an RMSD of 0.598 Å. The IA31.pdb Top1 protein was then deleted. 4) The intercalated topotecan in the 1K4T.pdb structure displaced the non-scissile strand nucleotides (101–114) and scissile strand nucleotides 11 and nucleotides 12–22, and so these portions of the DNA in the 1K4T.pdb structure were deleted leaving the corresponding DNA from the IA31.pdb structure. The IA31.pdb non-scissile strand bases 115–122 and scissile strand bases 1–10 in the DNA, that were aligned to the corresponding 1K4T.pdb bases (RMSD, 0.598 Å), were also deleted. The IA31.pdb scissile strand uridine 11 free 5’ sulfur (S’) was replaced with the native oxygen and the thymine base was replaced with a guanine (the 1K4T.pdb scissile strand base 11 is a guanine), and the base pairing non-scissile strand adenine in the 1A31.pdb DNA was replaced with a cytosine to allow for proper base pairing to the scissile strand guanine 11. Note: in the RCSB Protein Data Bank current “remediated” version of the IA31.pdb entry, the scissile strand nucleotide 11 S’ is incorrectly labeled as an O5’ 5). The backbone atoms of nucleotides 114/115 in the remaining non-scissile strand DNA (nucleotides 101–114 from IA31.pdb and 115–122 from 1K4T.pdb) were bonded and the entire Top1/dsDNA covalent-complex was typed with CFF, the bond orders for the atoms in the tyrosyl-phosphate bond were corrected as needed, and then minimized with an Implicit Solvent Model using Implicit Distance-Dependent Dielectrics and no restraints using first Steepest Descent and then Conjugate Gradient algorithms to a Final Convergence of 0.1. 6) The +1 nucleoside was rotated, around the backbone oxygen-phosphorus bond, out of the helix until the guanine base and deoxyribose 5’OH made a network of H-bonds/electrostatic interactions with Arg488, Asp533, and Arg590 as previously reported [15]. 7) CPT was docked into the Top1/dsDNA active-site such that it made the same network of H-bond/electrostatic interactions with Top1 and dsDNA as previously reported [15]. CPT was then modified into the following derivatives: 20-doxo CPT, 20-Cl CPT, hCPT and 10,11 diF-hCPT. Inhibitors were first typed with CFF and then the C-ring C15 and C16 atom types in hCPT and derivatives, and C10 and C11 atoms in CPT and derivatives, were manually corrected from cpb (bridge carbon biphenyl functional group) to cp (sp2 aromatic carbon in either five- or six-member ring). Inhibitors were then minimized with an Implicit Solvent Model using Implicit Distance-Dependent Dielectrics and no restraints using Steepest Descent and then Conjugate Gradient algorithms to a Final Convergence of 0.1. 8) Next the entire Top1/dsDNA/inhibitor covalent-complex was typed with CFF (again, inhibitor C-ring C15 and C16 atom types in hCPT and derivatives, and C10 and C11 atoms in CPT and derivatives, were corrected from cpb to cp), and then minimized with an Implicit Solvent Model (implicit distance-dependent dielectrics, dielectric constant of 3), Electrostatics spherical cutoff, Nonbond List Radius of 14 (nonbond higher cutoff distance 12, nonbond lower cutoff distance 10), and no restraints using Steepest Descent and then Conjugate
Gradient algorithms until either a RMS Gradient of 0.1, or Energy Change of 0.0, was reached.

Structure based Top1/dsDNA Intercalated model
All modifications and minimizations were carried out using Discovery Studio 2.0 (Accelrys, San Diego, CA). 1) In the Staker et al. 1K4T.pdb [18] the topotecan carboxylate, mercury, PEG, and all waters were deleted. 2) All incomplete Top1 side-chains were built out and the scissile strand +1 nucleoside 5′ sulfur (S5′) was changed to the native oxygen. 3) The topotecan A-ring 9-N(CH₃)₂ was replaced with a hydrogen to give 10-OH CPT, and then the A-ring 10-OH was replaced with a hydroxyl to give CPT. CPT was then modified into the following derivatives: 20-deoxy CPT, 20-CI CPT, hCPT and 9,10 diF-hCPT. Inhibitors were first typed with CFF and then the C-ring C15 and C16 atom types in hCPT and derivatives, and C10 and C11 atoms in CPT and derivatives, were manually corrected from cpb (bridge carbon biphenyl functional group) to cp (Sp2 aromatic carbon in either five- or six-member ring) and minimized with an Implicit Solvent Model using Implicit Distance-Dependent Dielectrics and no restraints using Steepest Descent and then Conjugate Gradient algorithms to a Final Convergence of 0.1. 4) Next the entire Top1/dsDNA/ inhibitor covalent complex was typed with CFF (inhibitor C-ring atom types were corrected from cpb to cp), and then minimized with an Implicit Solvent Model (implicit distance-dependent dielectrics, dielectric constant of 3), Electrostatics spherical cutoff, Nonbond List Radius of 14 (nonbond higher cutoff distance 12, deoxyguanosine, via in vitro experiments and molecular modeling studies. Bioorg Med Chem 12: 5225–5235.

Interaction energy calculations
All calculations were carried out using Discovery Studio 2.0 (Accelrys, San Diego, CA).

The interaction energy scores between inhibitors and Top1/dsDNA were calculated using an Implicit Solvent Model (implicit distance-dependent dielectrics, dielectric constant of 3) and a Nonbond List Radius of 14 (nonbond higher cutoff distance 12, nonbond lower cutoff distance 10), and no restraints using the Steepest Descent and then Conjugate Gradient algorithms until either a RMS Gradient of 0.1, or Energy Change of 0.0, was reached.

Acknowledgments
The author thanks both Dr. Sidney Hecht and Dr. Venkat Mathura for thoughtful discussions and review of the manuscript.

Author Contributions
Conceived and designed the experiments: GSL. Performed the experiments: GSL. Analyzed the data: GSL. Contributed reagents/materials/analysis tools: GSL. Wrote the paper: GSL.

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