Topoisomerase I Gene Mutations at F270 in the Large Subunit and N184 in the Small Subunit Contribute to the Resistance Mechanism of the Unicellular Parasite *Leishmania donovani* towards 3,3′-Diindolylmethane

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3,3′-Diindolylmethane (DIM), a novel poison targeting *Leishmania donovani* topoisomerase I (LdTOP1LS), induces programmed cell death in *Leishmania* parasites. The development of resistant parasites by adaptation with increasing concentrations of DIM generates random mutations in LdTOP1LS. Single-nucleotide mutations result in the amino acid substitutions F270L and K430N in the large subunit and N184S in the small subunit of the enzyme. DIM failed to inhibit the catalytic activity of the recombinant mutant enzyme (LdTOP1DRLS). Transfection studies of the mutant genes showed that the mutated topoisomerase I confers DIM resistance on wild-type *Leishmania* parasites. Site-directed mutagenesis studies revealed that a substantial level of resistance is conferred by the F270L mutation alone; however, all three mutations (F270L, K430N, and N184S) together are required to reach a higher-resistance phenotype. DIM fails to stabilize the topoisomerase I–DNA covalent complexes in the F270 mutant. Moreover, DIM cannot interfere with the religation step in the catalytic cycle of the recombinant F270L mutant enzyme. Taken together, these findings identify novel mutations in topoisomerase I that hinder its interaction with DNA, thereby modulating enzyme catalysis and conferring resistance to DIM. These studies advance our understanding of the mechanism of cell poisoning by DIM and suggest a specific modification of the drug that may improve its efficacy.

Eukaryotic DNA topoisomerase I is an essential enzyme that alters the topological changes of DNA that accompany DNA replication, transcription, recombination, and chromosomal segregation during mitotic cell division (3, 21). Topoisomerase I induces a transient single-stranded break of the DNA duplex and results in a reversible topoisomerase I–DNA covalent complex (2). Most eukaryotic type IB topoisomerases are monomeric enzymes, including human topoisomerase I, which comprises 765 amino acids (91 kDa). But interestingly, DNA topoisomerase I of the kinetoplast protozoan parasite *Leishmania donovani* is an unusual bisubunit enzyme, consisting of a large subunit (73 kDa) and a small subunit (29 kDa) (6). Recently, we demonstrated for the first time that in vitro reconstitution of the two recombinant proteins, the large subunit and the small subunit of *L. donovani* topoisomerase I (LdTOP1L and LdTOP1S, respectively), shows active enzyme. This active enzyme (LdTOP1LS) is located in both the nucleus and the kinetoplast of the parasite (6). Because topoisomerase I poisoning has been recognized as a promising pharmacological target for the development of therapeutic agents, a number of candidates have been identified. The potent antitumor compound 3,3′-diindolylmethane (DIM) is a well-characterized topoisomerase I inhibitor that stabilizes the topoisomerase I–DNA covalent complex. In addition to its antitumor effects, DIM has also been recognized to specifically target *L. donovani* topoisomerase I (16). Moreover, we recently demonstrated that DIM induces programmed cell death through the inhibition of mitochondrial F$_{1}$F$_{0}$ ATP synthase in the unicellular protozoan parasite *L. donovani* and reduces the parasitic loads in the livers and spleens of infected hamsters (17). Thus, DIM could be of interest in the treatment of human leishmaniasis. Therefore, precise determination of the mechanism of DIM resistance in *Leishmania* parasites could be an important strategic tool for acquiring a better understanding of its mechanism of molecular action.

Topoisomerase I is a target for anticancer as well as antiparasitic drugs, one important class of which is topoisomerase poison. These drugs act by stabilizing an intermediate stage of a reaction, where the enzyme is complexed with the cleaved gate helix. Resistance to topoisomerase I poison drugs is a clinical problem and may arise through a variety of mechanisms. Mammalian cell lines selected for resistance to camptothecin (CPT) exhibit decreased levels of topoisomerase I expression. In resistant murine P388 leukemia cells, CPT treatment is associated with a lower frequency of DNA single-strand breaks than that in wild-type (WT) cells and decreased expression of topoisomerase I at the mRNA and protein levels (8). Similar findings have been obtained with a human lung cancer cell line resistant to a CPT analog (CPT-11) (11) and
with a CPT-resistant tumor cell line (19). The levels of expression of topoisomerase I mRNA and protein in CPT-resistant human nasopharyngeal carcinoma cells (CPT30) were 30% and 40% lower, respectively, than those in the parent human nasopharyngeal carcinoma cell line (HONE-1) (4). Decreases in topoisomerase I expression have been associated with rearrangement and hypermethylation of the topoisomerase I gene (20). Thus, decreased expression and/or alteration of the topoisomerase I gene can confer resistance to CPT. Mutations affecting the way the drug interacts with the enzyme can cause resistance. Mutations that confer resistance to CPT have been identified in several regions of human topoisomerase I (Phe361Ser, Arg364His, Glu418Lys, Gly503Ser, Asp533Gly, Ala653Pro, and Asn722Ser) (5). A form of human topoisomerase I containing an Ala653Pro mutation has been shown to exhibit enhanced DNA strand religation, preventing the binding of CPT to the topoisomerase I-DNA complex (9). A point mutation resulting in the replacement of threonine with alanine at residue 729 has been found in a CPT-11-resistant human lung cancer cell line (12). Another mutation in to human nasopharyngeal carcinoma cell line (HONE-1) (4). Decreases and 40% lower, respectively, than those in the parent human nasopharyngeal carcinoma cells (CPT30) were 30% and were kept for 20 min at room temperature. The optical density was measured as the A535 on an enzyme-linked immunosorbent assay reader (Multiskan EX; Thermo Fisher Scientific, Waltham, MA).

**MATERIALS AND METHODS**

**Chemicals.** The bioactive compound DIM was synthesized chemically from indole and artrorpine by the addition of InCl3 (10 mol%) and isopropanol as described previously (16). CPT was purchased from Sigma Chemicals (St. Louis, MO). All drugs were dissolved in 100% dimethyl sulfoxide (DMSO) at 20 mM and were stored at −20°C.

**Development of a DIM-resistant parasite strain.** A highly resistant L. donovani strain (AG83) called LdDR50 was developed by stepwise exposure of the parasites to DIM at 1, 2, 5, 10, 20, 40, and 50 μM concentrations. When a 50 μM concentration was reached, the culture was diluted to ~10 cells/ml and separately cultured to generate a clonal population of parasites. The resistant parasites were maintained in drug-free medium for 72 h before use.

**Cell viability testing by an MTT assay.** The effect of the drug on the viability of the WT and LdDR50 cell lines was determined by a 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay (17). The cells were collected at the exponential phase and transferred to a 24-well plate (2 × 10⁴ cells per well). The cells were then incubated for 12 h in the presence of various concentrations of DIM (1, 5, 10, 15, and 20 μM). After incubation, the cells were centrifuged, and the cell pellet was washed twice with phosphate-buffered saline (PBS) (1×) and finally suspended in 100 μl of PBS (1×) in a 96-well plate. Ten microliters (10 μg/mL) of MTT solution was added to each sample in the 96-well plates and incubated for 4 h. After incubation, 100 μl of stop solution (stock; 4,963 μl isopropanol and 17 μl concentrated HCl) was added to each sample and kept for 20 min at room temperature. The optical density was measured as the A570 on an enzyme-linked immunosorbent assay reader.

**Measurement of DIM accumulation in cells.** WT and DIM-resistant Leishmania parasites grown to logarithmic phase were incubated at a density of 3 × 10⁴ cells/ml in culture medium containing 20 μM DIM at 22°C. At 1-, 5-, 10-, 20-, and 30-min intervals, 1-ml aliquots were taken, washed three times with ice-cold PBS (1.2 mM KH₂PO₄, 8.1 mM NaH₂PO₄, 130 mM NaCl, and 2.6 mM KCl, adjusted to pH 7.4) containing 0.1% N HCl and suspended in the same buffer. After sonication, the cell lysate was used to measure the accumulated amount of DIM by equilibrium dialysis as described previously (16). Each experiment was repeated three times with duplicate samples. Error bars were calculated from the means ± standard errors.

**Real-time RT-PCR.** Total RNA was prepared from WT and DIM-resistant promastigotes using the Total RNA isolation kit (Roche Biochemicals). cDNA was synthesized from 60 ng of total RNA using Superscript II RNaseH− reverse transcriptase (Invitrogen) and oligo(dT)₁₂₋₁₈ primers (Invitrogen) according to the manufacturer’s instructions. Semiquantitative PCR was performed in a 25-μl volume using 50 pmol each of sense and antisense primers corresponding to LdTOP1DR, LdTOP1DS, and L. donovani glyceroldehyde-3-phosphate dehydrogenase (GAPDH) by using the following profile: initial denaturation at 95°C for 5 min; 25 cycles with denaturation at 95°C for 1 min, annealing at 55°C for 40 s, and extension at 68°C for 40 s; and a final extension of 3 min. The primers were designed such that each set amplified a 300-bp fragment. Three separate reactions were carried out using three different RNA preparations in a separate reaction for each primer set in a 7300 real-time PCR system (Applied Biosystems). Reactions were carried out with the following profile: initial denaturation at 95°C for 5 min, followed by 35 cycles with denaturation at 95°C for 40 s, annealing at 60°C for 40 s, and extension at 65°C for 40 s. The PCR was followed by a melt curve analysis to ascertain that the expected products were amplified. The relative amounts of PCR products generated were obtained from the threshold cycle (Cₜ), and amplification efficiencies were normalized by dividing the values by the relative amount of the GAPDH gene, used as a control. The level of expression was calculated as 2^(-ΔΔCt) of experimental/vehicle control RT-PCR.

**Western blotting.** The DIM-resistant Leishmania strain LdDR50 and WT L. donovani cells (3 × 10⁷) were cultured for 12 h at 22°C with or without drugs. Nuclear fractions were isolated as described previously (16). Briefly, cells were suspended in hypotonic buffer (10 mM Tris-HCl [pH 7.5], 1 mM EDTA, 0.1 mM EGTA, 1 mM phenylmethylsulfonyl fluoride, 1 mM benzamidine hydrochloride, and 5 mM dithiothreitol) and homogenized. The homogenate was centrifuged for 10 min at 10,000 × g. The pellet was washed and used as the source of the nuclear fraction. The nuclear fractions, after lysis with 1% sodium dodecyl sulfate (SDS), were subjected to SDS-polyacrylamide gel electrophoresis (12%), and the proteins were electrotransfered to nitrocellulose membranes. Immobilized proteins were immunoblotted using rabbit antibodies raised against LdTOP1 (73 kDa) and LdTOP2 (29 kDa). Both polyclonal antibodies were diluted 1,000 times, and a horseradish peroxidase-conjugated secondary
antibody was used to visualize the reactive band by a diaminobenzidine color reaction.

Cloning and construction of recombinant plasmids. The full-length large subunit gene (LdTOP1L) and small subunit gene (LdTOP1S) of L. donovani topoisomerase IB were cloned into bacterial expression vector pET16b and transformed into Escherichia coli BL21(DE3) pLysS as described previously (6). Total cellular RNA was isolated from DIM-resistant L. donovani AG83 promastigotes (strain LdDR50) by using an RNA isolation kit (Roche Applied Science) according to the manufacturer’s protocol. Reverse transcription-PCR (RT-PCR) was performed with gene-specific primers corresponding to a single open reading frame (ORF) of 1,905 bp (LdTOP1DRL) using sense primer 5′-GGAATTCC ATATGATGAGTGGAGAAATGCAGAAG-3′, containing an Ndel site created at the initiation codon of the ORF, and antisense primer 5′-CGGATCT CTCACACCCCTAAGGGCTGAAAG-3′, with a BamHI site immediately downstream from the termination codon. (Restriction endonuclease sites are underlined.) RT-PCR was performed with another set of gene-specific primers corresponding to a single ORF consisting of 786 bp (LdTOP1DRS): sense primer 5′-GGAATTCCATATGATGAGTGGAGAAATGCAGAAG-3′, containing an Ndel site created at the initiation codon of the ORF, and antisense primer 5′-CGGATCTCCTTAAATCGGAATTCGCGC-3′, with a BamHI site immediately downstream from the termination codon. The fragment amplified by RT-PCR was cloned into the Ndel/BamHI site of the pBAD/MF2 vector (Invitrogen) and used for the subsequent experiments.

Overexpression and purification of recombinant proteins. The recombinant constructs pET16b-LdTOP1L, pET16b-LdTOP1S, and pET16b-LdTOP1DRS were separately transformed into Escherichia coli BL21(DE3) pLysS cells. The recombinant proteins were overexpressed and purified using a phosphocellulose column (Qiagen). Proteins were dialyzed and purified using a phosphocellulose column (P11; Whatman) as described previously (6). Finally, the purified proteins LdTOP1L, LdTOP1S, LdTOP1DRL, and LdTOP1DRS were stored at −70°C.

In vitro reconstitution of recombinant proteins. Purified LdTOP1L was mixed with purified LdTOP1S and LdTOP1DRS, while LdTOP1S was mixed with purified LdTOP1DRS, and purified LdTOP1DRS was mixed with LdTOP1DRS, in a 1:1 molar ratio according to a standard protocol (6). The total protein concentration was 0.5 mg/ml in the standard reconstitution buffer containing 50 mM potassium phosphate (pH 7.5), 0.5 mM dithiothreitol, 1 mM EDTA, 0.1 mM phenylmethylsulfonyl fluoride, and 10% glycerol. The mixture was dialyzed at 4°C for 12 h, and the dialyzed proteins were used subsequently in all the assays.

DNA sequencing. The LdTOP1L and LdTOP1S genes were sequenced with the primers described above by using an ABI Prism DNA sequencing kit (Perkin-Elmer, Norwalk, CT).

Single-site mutagenesis. Single mutations were introduced into the Leishmania heterodimeric topoisomerase I at positions Phy270 and Lys430 of the large subunit and Asp184 of the small subunit. Mutagenesis was performed by using the QuikChange XL site-directed mutagenesis kit (Stratagene, La Jolla, CA) according to the manufacturer’s protocol. To carry out the desired mutations, pET16b-LdTOP1L and pET16b-LdTOP1S were used as templates for all mutagenesis experiments. For each mutation, the WT nucleotide was replaced using a specific pair of mutagenic primers. The following sense primers, along with their antisense counterparts (with codons in boldface and substitutions underlined), were used: for LdTOP1L720L, 5′-CCGACATGTTGGAAGTTGGA GAAGGCGGACACGC-3′ and 5′-GCTTGGCTGTCTTCTCAATCACC ATGTCGGC-3′; for LdTOP1L419N, 5′-CCGATGCGACCGAAAATGTGTTGACC GGCC-3′ and 5′-GGCGGTCGACGACCCATTTGCGG-3′; for LdTOP1S586R, 5′-GGCTGCATCTTTCTCAACCAGGATCGATCGAGGCGC-3′ and 5′-GGCTGCTGCCGTCTCCGAGGCAAAGAGC-3′. Mutations were confirmed through DNA sequencing in an ABI Prism DNA sequence (Perkin-Elmer, Norwalk, CT).

Cloning of transfection constructs and transfection into Leishmania parasites. LdTOP1L and LdTOP1S were PCR amplified using sense primers 5′-TCCCGGAGGTAGGGAGTGAATACGGGA-3′, containing a Smal site created at the initiation codon of the ORF, and 5′-ATGAAGAATGCGGCGG CATGCAGCGTGTCTTT3′, containing an Ndel site created at the termination codon of the ORF, respectively, along with antisense primers 5′-CCGG ATCCCTACATCCCTCAAGGGCTCGAAG-3′ for LdTOP1L and 5′-CCGG ATCCCTAAAAATGACATCTGCCG-3′ for LdTOP1S. Each construct contained a BamHI site immediately downstream from the termination codon. (Restriction endonuclease sites are underlined.) The amplified fragments were cloned into the SmaI/BamHI site of the pXG-HYG vector (BD Biosciences) and the NotI/BamHI site of the pXG-GFP2 vector (a kind gift from S. M. Beverley) containing the green fluorescent protein (GFP) fragment. The transfection constructs pXG-HYG-LdTOP1L and pXG-GFP2-LdTOP1S were electrotransformed according to the protocol in reference 15, and the transfected clones were obtained by selection of hygromycin B (50 μg/ml) or tetracycline (0.8 μg/ml) in 1 ml of culture medium containing 50 μg/ml of G418 (50 to 300 μg/ml) for pXG-HYG-LdTOP1L and G418 (50 to 300 μg/ml) for pXG-GFP2-LdTOP1S.

Plasmid relaxation assay. The type I DNA topoisomerase was assayed by the decreased mobility of the relaxed isomers of supercoiled pBluescript SK (+) DNA in an agarose gel as described previously (16). The fluorescence of the supercoiled monomer DNA band after staining with ethidium bromide (0.5 μg/ml) was quantitated by integration using Gel Doc 2000 under UV illumination (Quantity One software; Bio-Rad).

Equilibrium cleavage assay. The 25-mer oligonucleotide 1′(5′-GAAAAAGACCTT 3′AGAGAAAATTTTTAAG-3′) (with the downward-pointing arrow marking a cleavage site) was 5′-end labeled with [γ-32P]ATP and annealed with oligonucleotide 2′(5′-TAAATTTTTCTAGTTTCTTCTTCT-3′) to make a duplex oligonucleotide containing a topoisomerase I binding motif as described previously (16). Cleavage was carried out using a 20-fold molar excess of the WT (LdTOP1L) or mutant (LdTOP1L720L, LdTOP1L419N) enzyme over the duplex 25-mer DNA (enzymes, 0.2 μM; DNA, 10 nM). The reactions were carried out in the presence or absence of DIM at 37°C for 30 min. The samples were analyzed on a 12% denaturing polyacrylamide gel, followed by autoradiography as described previously (16).

Single-site mutation assay. A 25-mer oligonucleotide containing a topoisomerase IB-specific cleavage site (5′-GAAAAAGACCTT 3′AGAGAAAATTTTTAAG-3′) was 5′-end labeled with [32P] and annealed to a 25-mer oligonucleotide 3′(CTTTTTTCTAAGCTTTTTTTTTTTTTAAT-P′) as described previously (16). The suicidal cleavage reaction was carried out with 5 nM DNA substrate and 0.15 μM WT (LdTOP1L) or mutant (LdTOP1L720L, LdTOP1L419N) enzyme in 20-μl reaction mixtures under standard assay conditions at 25°C for 4 h in the absence or presence of DIM as described previously (16). For religation experiments, covalent complexes generated by incubating the suicidal DNA substrate with the WT or mutant enzyme in the absence or presence of DIM (20 μM) were transferred to 30°C and preincubated for 2 min. The religation reaction was initiated by the addition of a 300-fold molar excess of the 11-mer religation acceptor oligonucleotide (5′-OH-AGAGAAAATTTTT-3′) to the same reaction mixture and incubation for the indicated times. Finally, all the reactions were stopped by the addition of SDS, and DNAs were subsequently precipitated by ethanol. Samples were digested with 5 μl of 1-mg/ml trypsin, electrophoresed in 12% denaturing polyacrylamide gels, and autoradiographed.

Equilibrium dialysis. The equilibrium dialysis experiment was performed as described previously (16). The binding solution, as well as the buffer, used for equilibrium dialysis was 20 mM Tris-HCl (pH 7.5)–30 mM NaCl–10 mM MgCl2. Four sets of reactions were performed in 500 μl of binding solution containing 200 μM enzyme (LdTOP1LDRS, LdTOP1L586R, LdTOP1S419N) to the same suicidal DNA substrate in the absence or presence of DIM as described previously (16). For religation experiments, covalent complexes generated by incubating the suicidal DNA substrate with the WT or mutant enzyme in the absence or presence of DIM (20 μM) were transferred to 30°C and preincubated for 2 min. The religation reaction was initiated by the addition of a 300-fold molar excess of the 11-mer religation acceptor oligonucleotide (5′-OH-AGAGAAAATTTTT-3′) to the same reaction mixture and incubation for the indicated times. Finally, all the reactions were stopped by the addition of SDS, and DNAs were subsequently precipitated by ethanol. Samples were digested with 5 μl of 1-mg/ml trypsin, electrophoresed in 12% denaturing polyacrylamide gels, and autoradiographed.

RESULTS
Development of DIM-resistant parasites and accumulation of DIM in the cells of WT and resistant parasites. In order to determine the DIM resistance mechanisms of Leishmania parasites, we selected the parental L. donovani parasites (AG83) for resistance to DIM in a step-by-step manner until they reached a resistance level of 42-fold over their WT counterparts. More specifically, a resistant L. donovani strain was selected by stepwise exposure to increasing concentrations
One of the possible potential mechanisms of drug resistance in parasites is increased efflux or decreased influx. So for *Leishmania* parasites, one possible way to develop resistance to DIM could be to block the uptake or increase the efflux of DIM. Since DIM has an absorbance maximum at 280 nm, we tested these possibilities by measuring its accumulation inside the parasites. Moreover, spectrophotometric analysis of the drug kinetics was performed to determine whether the WT strain was more permeable to DIM than resistant parasites. As reported in Fig. 1B, no significant differences in permeability were detected between the WT strain and LdDR50, even at a high concentration of DIM (20 μM), suggesting no alteration in the accumulation of the drug in the resistant strain. Since DIM accumulation appears to remain same in the WT and resistant parasites, we further investigated the molecular aspects of topoisomerase I and its interaction with DIM.

**ABC transporters are not involved in the resistance of parasites to DIM.** In CPT-resistant parasites, CPT accumulation decreased primarily through the ABC transporters present in the cell membrane. The ABCG6 gene is threefold overexpressed in that phenotype (1). Therefore, we investigated the involvement of ABC transporters in DIM resistance in parasites. Real-time RT-PCR quantitation revealed that the expression levels of ABCC3, ABCG4, and ABCG6 in DIM-resistant parasites were similar to those in WT parasites. Parasites resistant to CPT, sodium antimony gluconate (SAG), and miltefosine were used as positive controls. The *ABCG6*, *ABCC3*, and *ABCG4* transporters were approximately threefold, fourfold, and threefold overexpressed in CPT-, SAG-, and miltefosine-resistant parasites, respectively (Fig. 2; see also Fig. S1A in the supplemental material), results consistent with our previous findings (1). The expression patterns of ABC genes were also compared for DIM-resistant cell types. No differences in the expression of the *ABC* genes at the level of mRNA were observed between DIM-resistant cells and WT cells. For all cell types, the expression of the *GAPDH* gene was observed to be equivalent, and it served as the loading control.

**Reduced expression of topoisomerase I in DIM-resistant *Leishmania* parasites.** The basis for DIM resistance in *Leishmania* parasites was studied by examining topoisomerase I gene expression at the mRNA and protein levels. Real-time RT-PCR analysis of total RNA extracted from exponentially
growing WT *Leishmania* parasites and parasites resistant to DIM (LdDR50), CPT (Ld20C), SAG (LdGE1), or miltefosine (Ld30M) revealed 4.6-fold and 3-fold decreased expression of the large subunit and 4.3-fold and 3.7-fold decreased expression of the small subunit of mRNA in DIM-resistant and CPT-resistant parasites, respectively, compared to WT cells (Fig. 2; see also Fig. S1A in the supplemental material). There were no such changes in the expression of either subunit at the level of mRNA in SAG- or miltefosine-resistant parasites. For all cell types, the expression of the GAPDH gene was observed to be equivalent, and it served as the loading control. Western blot analysis was performed to assess the levels of topoisomerase I protein expression in both WT and LdDR50 cells (see Fig. S1B in the supplemental material). The expression of the large and small subunits of LdTOP1LS in LdDR50 cells was 4.0-fold and 3.8-fold less than that in WT cells. The mRNA expression results are consistent with the protein expression results, suggesting that the downregulation of topoisomerase I in DIM-resistant parasites occurs at the transcriptional level.

**Topoisomerase I gene sequencing.** Two point mutations in the large subunit of *L. donovani* topoisomerase I confer CPT resistance (13). To address whether point mutations confer resistance, we cloned the genes from WT and mutant parasites by RT-PCR and sequenced them. By sequence alignment of the WT and DIM-resistant genes for topoisomerase I, two amino acid substitutions resulting from two single-nucleotide mutations were observed in LdTOP1DRL and one amino acid substitution resulting from one single-nucleotide mutation was observed in LdTOP1DRS (see Table S1 and Fig. S2 in the supplemental material). Two amino acids found in WT parasites, phenylalanine 270 (F270) and lysine 430 (K430), were mutated by site-directed mutagenesis to leucine (L) and asparagine (N) by C-to-G and A-to-T transitions in the LdTOP1DRL of LdDR50 cells, respectively. Based on the crystal structure of *Leishmania* topoisomerase I (7), we observed that F270 in the large subunit is located near the active-site tyrosine 222 of the small subunit (Fig. 3) and that the site tyrosine 222 of the small subunit (Fig. 3) and that the K430N mutation occurs far from the active-site tyrosine. This observation suggests that amino acid F270 of the large subunit mediates the interaction between DIM and the enzyme. In addition, a mutation of asparagine 184 (N184) to serine (S) was observed for the small subunit in DIM-resistant parasites due to an A-to-G transition (see Table S1 in the supplemental material). None of the three mutations F270L, K430N, and N184S are conserved in human topoisomerase I.

**Transfection of WT *L. donovani* with mutated LdTOP1DRLS.** To investigate the potential resistance mechanism and to evaluate the contribution of the three mutations observed to DIM resistance, we cloned the mutated LdTOP1DRL and LdTOP1DRS genes into a *Leishmania* expression vector. We also performed site-directed mutagenesis on the LdTOP1L and LdTOP1S genes to obtain amino acid substitutions resulting in the formation of mutated LdTOP1L and LdTOP1S. As a control, we also transfected *Leishmania* parasites with overexpressed WT LdTOP1L and LdTOP1S in order to eliminate the possible effect of the endogenous enzyme in conferring resistance on the parasites. When WT parasites were transfected with LdTOP1DRLS containing three amino acid substitutions (F270L and K430N in the large subunit and N184S in the small subunit), they showed a higher level of DIM resistance, similar to that shown by LdDR50 parasites (Fig. 4). Furthermore, we found that transfection of WT parasites with an F270L (large subunit with Phe270Leu reconstituted with the WT small subunit), F270L K430N (large subunit with Phe270Leu and Lys430Asp reconstituted with the WT small subunit), or F270L N184S (large subunit with Phe270Leu reconstituted with the small subunit with Asp184Ser) mutant individually resulted in the formation of cells that also conferred strong levels of resistance to DIM. The growth of the resulting cells is inhibited by 34%, 34%, and 31%, respectively, at 20 μM DIM.

Transfection of WT cells individually with (i) a K430N mutation in the large subunit and a WT small subunit (LdTOP1S), (ii) a K430N mutation in the large subunit and an N184S mutation in the small subunit, or (iii) a WT large subunit (LdTOP1L) and an N184S mutation in the small subunit did not induce significant resistance to DIM. The growth of these mutants was inhibited 94%, 86%, and 91%, respectively, in the presence of 20 μM DIM. These results, taken together, suggest that the F270L mutation in the large subunit is responsible for the acquisition of resistance to DIM by the parasites.

**Mutant *L. donovani* topoisomerase I is insensitive to DIM.** To investigate the effect of DIM on resistant *L. donovani* topoisomerase I (LdTOP1DRLS), we cloned, overexpressed, and purified DIM-resistant mutated topoisomerase I and per-
formed plasmid relaxation assays (Fig. 5) as described in Materials and Methods. WT recombinant LdTOP1LS was purified as described previously (6). The relaxation experiments were performed under standard assay conditions where the plasmid DNA and the enzyme (LdTOP1DRLS or LdTOP1LS) were mixed at a molar ratio of 3:1. Under this condition (ratio of DNA to LdTOP1DRLS, 3:1), DIM was used at increasing concentrations from 1 μM to 50 μM. There was no catalytic inhibition of LdTOP1DRLS in the presence of DIM (Fig. 5, lanes 4 to 8), whereas the WT enzyme, LdTOP1LS, was completely inhibited at 10 μM DIM (Fig. 5, lane 12). It can be inferred from these results that the DIM-resistant mutant enzyme LdTOP1DRLS is insensitive to DIM.

To investigate the interacting residues of the enzyme that bind with DIM, we purified the subunits generated by site-directed mutagenesis, reconstituted the enzymes, and performed a standard relaxation assay (Fig. 6) in the presence of DIM. The enzymes showed no sensitivity to DIM when the mutated subunits were reconstituted as follows: the large subunit with F270L plus K430N and the small subunit with N184S (Fig. 6, lane 6), the large subunit with F270L and the WT small subunit (lane 8), the large subunit with F270L plus K430N and the WT small subunit (lane 12), and the large subunit with F270L and the small subunit with N184S (lane 14). Moreover, DIM sensitivity was restored when the large subunit with K430N was reconstituted with the WT small subunit (Fig. 6, lane 10) or with the small subunit with N184S (lane 16) and when the WT large subunit was reconstituted with the small subunit with N184S (lane 18). These results suggest that F270 in the large subunit is responsible for the sensitivity of the enzyme to DIM.

F270 is responsible for the DIM-induced stabilization of the cleavable complex. To investigate the mechanism of DIM resistance in the mutant enzymes, the stability of the topoisomerase I–DNA covalent complex was analyzed using the 25-mer full duplex oligonucleotide substrate in the presence of increasing concentrations of DIM (5 to 20 μM). After 30 min of incubation, the reactions were stopped with SDS, the samples were treated with trypsin, and the products were analyzed by polyacrylamide-urea gel electrophoresis (Fig. 7). In the absence of DIM, the cleavage-religation equilibrium was shifted toward religation for the WT enzyme; little trapped cleavable complex was found, and most of the cleaved product was religated and migrated as an uncleaved 25-mer (Fig. 7A, lane 2). When the WT enzyme was exposed to increasing concentrations of DIM, the cleavage-religation equilibrium was shifted toward cleavage, as reported previously (16). The band corre-
sponding to the 12-mer cleaved product is clearly detectable, and the band intensity increases with increasing concentrations of DIM (Fig. 7A, lanes 3 to 6). For the F270L mutant, the equilibrium shifted toward religation in the absence of DIM, as for the WT enzyme (Fig. 7B, lane 2). But in the presence of increasing concentrations of DIM, the equilibrium was not shifted toward cleavage; there are no bands of 12-mer cleaved products in the gel (Fig. 7B, lanes 3 to 6). This observation revealed that the F270L mutant is insensitive to DIM and that DIM is unable to stabilize its cleavable complex. Moreover, in the presence of increasing concentrations of DIM, the cleavage-religation equilibrium shifted toward cleavage for the K430N mutant, as for the WT enzyme (Fig. 7C). However, for the N184S mutant, the cleavage equilibrium also shifted to religation at lower concentrations of DIM, and stabilization of cleavage occurred only at higher concentrations (Fig. 7D, lane 6). These results indicate that the K430N mutation in the large subunit is not responsible for the stabilization of the cleavable complex but might play some role in DIM binding. Moreover, the N284S mutant enzyme has some role in stabilizing the formation of the cleavable complex only at the highest concentration (20 μM) of DIM.

DIM cannot interfere with the religation step in the F270L mutant topoisomerase I reaction. To investigate whether DIM interferes with the religation reaction carried out by DIM-resistant mutant topoisomerase I, we performed time-dependent single-turnover religation experiments in the absence and in the presence of DIM (20 μM) with mutant enzymes (Fig. 8). In these experiments, religation was studied by assaying the ability of the covalent intermediate to attach a 5'-OH-terminated 11-mer to the cleaved oligonucleotide, forming a 23-mer product. Since we have observed that the F270 residue is solely responsible for the DIM-induced stabilization of the cleavable complex, the religation reactions were performed with WT LdTOP1LS and mutant LdTOP1LF270/LdTOP1LS as described in Materials and Methods. Reactions were stopped at various time points by the addition of an equal volume of 1% SDS. The samples were treated with trypsin to remove all but a short trypsin-resistant peptide from the topoisomerase I–DNA co-
valent complex before analysis in a sequencing gel. The WT and mutant enzymes were incubated for 60 min with a suicide substrate in order to generate the cleavable complexes. Once cleavage had occurred, 11-mer religation oligonucleotides were added for different time periods in the absence or presence of 20 μM DIM to start the religation process. As shown in Fig. 8, in the absence of DIM, the religation rate of the F270L mutant enzyme (Fig. 8B) was higher than that of the WT enzyme (Fig. 8A). Moreover, the presence of DIM did not affect the religation rate of the F270L mutant (Fig. 8B, lanes 5 to 7), but it inhibited the religation of the WT enzyme (LdTOP1LS) (Fig. 8A, lanes 5 to 7), as reported previously (16). We also performed the religation experiment with the N184S mutant (Fig. 8C) and found that the religation reaction was inhibited in the presence of DIM (Fig. 8C, lanes 5 to 7), as with the WT enzyme. These results indicate that DIM is able to bind the WT enzyme and inhibit its religation process but is unable to stabilize the enzyme-DNA covalent complex, and fails to inhibit the religation process, for the DIM-resistant LdTOP1F270L/S mutant enzyme.

![FIG. 8](image)

**FIG. 8.** Effect of DIM on single-turnover cleavage and religation by WT and mutant enzymes. Shown are results of time course religation experiments with the 32P-5′-end-labeled suicide DNA substrate (14-mer/25-mer) as indicated and the WT enzyme (LdTOP1LS) (A) or the F270L (B) or N184S (C) mutant enzyme (C). The DNA substrate was incubated with WT or mutant topoisomerase I in the absence or presence of DIM (20 μM) at 23°C as described in Materials and Methods. Lanes 1, suicide DNA substrate with the enzyme. Lanes 2 to 4, same as lanes 1, but the enzyme was incubated with 11-mer religation oligonucleotides for 30, 60, and 120 s, respectively, in the absence of DIM. Lanes 5 to 7, same as lanes 2 to 4, respectively, but in the presence of DIM. All the reactions were stopped by the addition of 2% (wt/vol) SDS. The samples were precipitated with ethanol, digested with trypsin, and analyzed by denaturing polyacrylamide sequencing gel electrophoresis. The uncleaved suicidal oligonucleotide, the covalent complex, and the religation products are indicated.

The binding affinity of DIM for the WT enzyme is higher than that for the DIM-resistant mutant enzyme. From the crystal structure of the large subunit of *L. donovani* topoisomerase I, it is found that F270 is close to the active-site tyrosine residue of the small subunit and is a DNA binding residue, present in the DNA binding core domain. To investigate the binding affinity of DIM for the mutant enzyme, we performed equilibrium dialysis experiments as described in Materials and Methods. The number of DIM molecules bound to the proteins was plotted against increasing concentrations of DIM (Fig. 9). The dissociation constant ($K_D$) was calculated from the equilibrium dialysis experiments. The $K_D$ of DIM-LdTOP1DRLS is $2.4 \times 10^{-4}$ M. Previously, we found that the $K_D$ of the WT enzyme, LdTOP1LS, bound to DIM was $9.7 \times 10^{-9}$ M (16). From these results it was calculated that the affinity of DIM for LdTOP1DRLS is 2.5 $\times$ 10^3-fold less than that for the WT enzyme. This finding reveals that DIM interacts very weakly with the DIM-resistant mutant enzyme, per-
haps explaining the insensitivity of the enzyme to DIM in the relaxation assays.

To find out the differential affinity of DIM for the mutated subunits of LdTOP1LS, we also investigated the binding of both the mutated subunits to DIM by equilibrium dialysis experiments. The numbers of DIM molecules bound by LdTOP1L,F270L,K430N/1S,N184S, LdTOP1L,F270L,K430N, and LdTOP1S,N184S were plotted against increasing concentrations of DIM (Fig. 9). The binding constants of DIM-LdTOP1L,F270L,K430N/1S,N184S (Fig. 9A), DIM-LdTOP1L,F270L,K430N (Fig. 9B), DIM-LdTOP1L,F270L (Fig. 9C), and DIM-LdTOP1SL,N184S (Fig. 9D) were 2.4 × 10⁻⁵ M, 7.6 × 10⁻⁴ M, 5.3 × 10⁻⁴ M, and 3.8 × 10⁻⁴ M, respectively. It was reported previously that the binding constants of DIM-LdTOP1L and DIM-LdTOP1S were 4.28 × 10⁻⁷ M and 5 × 10⁻⁸ M, respectively (16). The results obtained here suggest that the affinity of DIM for the large subunit is reduced 1.2 × 10⁻⁵-fold by the F270L mutation and 7.6 × 10⁻⁴-fold by the N184S mutation. Therefore, the results indicate that the low affinity of DIM for the mutated holoenzyme is due to the F270L mutation of the large subunit and the N184S mutation of the small subunit.

**DISCUSSION**

The structure-function analysis of the DIM–topoisomerase I interaction might be exploited in the development of rational approaches to chemotherapy for human leishmaniasis and also in the study of drug resistance. Resistance to traditional first-line drugs underlines the urgent need for new treatments. The unusual bisubunit topoisomerase I of *Leishmania* parasites is an attractive chemotherapeutic target. Various topoisomerase I inhibitors have been tested for their efficacies as antileishmanial agents. Recently, we have reported that DIM is a potent noncompetitive topoisomerase I inhibitor (16).

The possible potential mechanisms of drug resistance in a population of parasites include the following: (i) differential selection of resistant individuals from a mixed population, (ii) physiological adaptations, such as increased efflux or decreased influx, (iii) conversion of the drug to an inactive form by an enzyme, (iv) changes in gene expression, and (v) spontaneous mutations followed by selection. Hence, by developing DIM-resistant *Leishmania* parasites, a better understanding of the various mechanisms of development of chemoresistance can be obtained. Because the *Leishmania* plasma membrane represents the first cellular barrier that DIM encounters before reaching its specific target, we determined the efficiency of the drug at accumulating inside WT and resistant parasites in order to verify differences in the kinetics of DIM transport. The experiment on the transport of DIM in LdDR50 and WT parasites revealed no alteration in drug accumulation for the resistant parasites. Eukaryotic ABC transporters are involved in the translocation of various substances across the cell membrane. Drug resistance arising out of altered influx or efflux of drugs involves different classes of ABC transporters. Resistance to SAG is manifested by overexpression of the ABCB5 transporter, while overexpression of the ABCG4 transporter is involved in miltefosine resistance and overexpression of the ABCG6 transporter is involved in CPT resistance, in *Leishmania* parasites (1).

In the present study, we have characterized the mechanism of resistance of *Leishmania* parasites to DIM. LdDR50 parasites were about 42-fold more resistant to DIM than the WT strain. As with other drugs, resistance to DIM might involve alterations in cellular drug accumulation, the drug target, or the response to the drug-target interaction. The present study indicates that ABC transporters were not overexpressed in DIM-resistant *Leishmania* parasites. However, this study shows that without changes in the drug-trafficking pathway, these parasites develop a chemoresistance phenotype by harboring random mutations in the target genes.

To further study the mechanisms for the development of resistance in LdDR50, a series of quantitative and qualitative assays for topoisomerase I was performed. Real-time RT-PCR and Western blot analysis showed lower expression of the large and small subunits at both the mRNA and protein levels in LdDR50 parasites than in their WT counterparts. This result was consistent with that of a previous report where expression of topoisomerase I was reported to be decreased in CPT-resistant tumor cell lines (19). Moreover, the study with CPT-resistant *Leishmania* parasites identified two point mutations in the large-subunit gene of LdTOP1LS (13). To address this possibility, we sequenced both the large- and small-subunit genes (LdTOP1L and LdTOP1S) of DIM-resistant parasites. LdDR50 parasites had two amino acid substitutions (F270L and K430N) in the core domain of LdTOP1L and one amino acid substitution (N184S) in LdTOP1S. Based on mutations that we have identified in both the large and the small subunit, we hypothesized that these amino acid residues of the enzyme are crucial for the cytotoxic action of DIM against *Leishmania* parasites. Despite substantial resistance conferred by each amino acid substitution individually, site-directed mutagenesis studies indicated that all three substitutions were required to reach the highest level of the DIM-resistant phenotype. From a homology modeling study of *L. donovani* topoisomerase I and human topoisomerase I, we determined that F270 and K430 of LdTOP1L and N184 of LdTOP1S are equivalent to Y448, D614, and K642 of human topoisomerase I, respectively, residues that are predicted to interact with the essential indole moieties of DIM. The crystal structure of *L. donovani* topoisomerase I (7) revealed that K430 is quite far from the active-site tyrosine (Tyr222) of the small subunit, which is involved in cleavage activity, but that F270 is very close to the active-site Tyr222. The distance between the carbon of F270 and the closest atom of Tyr222, the phenolic oxygen atom, is 3.28 Å. F270 plays a role in stabilizing the phenolic anion during the attack on the scissile phosphate group. The distance between the carbon of F270 and the phosphate of bound DNA is 7.41 Å. Moreover, in a previous study of a CPT-11-resistant human lung cancer cell line, it was demonstrated that a mutation in residue 729, near the active site of the enzyme, is associated with decreased sensitivity (12). The equilibrium cleavage/religation experiment revealed that K430 has no role in the DIM-induced stabilization of the covalent complex, whereas this stabilization is inhibited for the F270L mutant enzyme compared to the WT enzyme. The results presented here provide a molecular explanation for the DIM resistance of the mutants of topoisomerase I.

The religation experiment indicates that DIM strongly inhibits the religation process for the native enzyme but it is inefficient at inhibiting religation for a mutant enzyme.
(F270L). Moreover, in the absence of DIM, the religation process of the mutant enzyme is faster than that of WT. Previously it was reported that the Ala653Pro mutation of human topoisomerase I exhibits enhanced DNA strand religation and inhibits the CPT-induced formation of the topoisomerase I–DNA cleavable complex (5). Based on these findings, we hypothesize that the increased religation rate of the F270L mutant enzyme results from conformational changes within the topoisomerase I catalytic pocket that enhance the binding of DNA to the covalent complex. In particular, mutation of the highly flexible phenylalanine might extend the DNA to the covalent complex. In particular, mutation of the mutant enzyme results from conformational changes within the hypothesize that the increased religation rate of the F270L in inhibits the CPT-induced formation of the topoisomerase I cleavage complexes in the structure shown in Fig. 3. The insensitivity of the F270L mutant to DIM might be due to the high affinity of DIM for the phenyl ring of the aromatic residue F270 in the WT enzyme. Our studies also suggest that mutation-induced alterations in the enzyme active-site architecture coincide with decreased DIM sensitivity.

In conclusion, we have shown for the first time the mechanism of DIM resistance in Leishmania parasites and the occurrence of a novel point mutation, F270L, in the large subunit. F270 is very close to the active-site tyrosine and is responsible for DIM-induced cleavage. The inability to induce topoisomerase I–DNA cleavable complexes with the DIM-resistant topoisomerase I enzyme demonstrates that DIM is a promising non-CPT topoisomerase I inhibitor for therapeutic development and a useful reagent for mapping topoisomerase I cleavage complexes in the Leishmania genome. Interestingly, this is the first report concerning the mechanism of resistance to DIM in any living organism. Moreover, we have identified three topoisomerase I mutation sites conferring resistance to DIM. These mutations provide new insight into the molecular interactions between DIM and the enzyme. This study advances our understanding of the mechanism of cell poisoning by the dietary phytochemical compound DIM and suggests specific modifications to the drug that may improve the efficacy of the molecule.

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