DNA-Binding Properties of Bis-N-substituted Tetrandrine Derivatives

Sandra Mónica González-Martínez, Drochss Petry Valencia-Ochoa, Juan Carlos Gálvez-Ruiz, Mario Alberto Leyva-Peralta, Octavio Juárez-Sánchez, María A. Islas-Osuna, Viviana Isabel Calvillo-Páez, Herbert Höpfíl, Ramón Íñiguez-Palomares, Fernando Rocha-Alonzo, and Karen Ochoa Lara*

ABSTRACT: A series of bis-N-substituted tetrandrine derivatives carrying different aromatic substituents attached to both nitrogen atoms of the natural alkaloid were studied with double-stranded model DNAs (dsDNAs) to examine the binding properties and mechanism. Variable-temperature molecular recognition studies using UV–vis and fluorescence techniques revealed the thermodynamic parameters, $\Delta H$, $\Delta S$, and $\Delta G$, showing that the tetrandrine derivatives exhibit high affinity toward dsDNA ($K \approx 10^6$–$10^7$ M$^{-1}$), particularly the bis(methyl)anthraquinone (BAqT) and bis(ethyl)indole compounds (BlnT). Viscometry experiments, ethidium displacement assays, and molecular modeling studies enabled elucidation of the possible binding mode, indicating that the compounds exhibit a synergic interaction mode involving intercalation of one of the N-aryl substituents and interaction of the molecular skeleton in the major groove of the dsDNA. Cytotoxicity tests of the derivatives with tumor and nontumor cell lines demonstrated low cytotoxicity of these compounds, with the exception of the bis(methyl)pyrene (BPyrT) derivative, which is significantly more cytotoxic than the remaining derivatives, with IC$_{50}$ values against the LS-180, A-549, and ARPE-19 cell lines that are similar to natural tetrandrine. Finally, complementary electrochemical characterization studies unveiled good electrochemical stability of the compounds.

1. INTRODUCTION

The natural alkaloid $\text{S, S-(+)-tetrandrine}$ has been extensively studied due to its multiple pharmacological activities. Tetrandrine is a macrocyclic compound that can be functionalized to yield diverse derivatives with enhanced activity. So far, the alkaloid and some of its various derivatives have been evaluated as potential treatments for cancer, viral and bacterial infections, cardiovascular afflictions, pulmonary fibrosis, diabetes, and cognitive impairment. However, one of the main limitations of using tetrandrine as a drug is its poor solubility in water and the consequent low bioavailability. On the other hand, derivatization of tetrandrine has been almost exclusively on its aromatic rings, leaving the functionalization on the nitrogen atoms largely unexplored, a strategy that can improve solubility in water and simplify the synthesis.

Our research groups have previously reported on a series of cationic mono- and bis-N-substituted tetrandrine derivatives containing up to two additional chiral centers. These compounds promote ester cleavage and bind substrates of biological importance such as nucleotides and double-stranded (ds) DNA through a combination of noncovalent interactions, including π-stacking. Recently, a bis(methyl)naphthyl tetrandrine derivative was shown to self-assemble into an organic supramolecular framework with extremely large spherical cavities, indicating that tetrandrine is a versatile scaffold molecule with several potential applications.

Selective recognition of nucleic acids remains a primary goal for diverse reasons, including the design of novel anticancer agents, antibiotics, and antiviral drugs and the intracellular detection of DNA and RNA. Specific strategies have been developed for this purpose, depending on the desired application. For instance, effective anticancer and antiviral agents contain, frequently, planar aromatic substituents that intercalate into the base pairs of dsDNA, inhibiting the replication enzyme topoisomerase II, among other effects. Other binding modes of small molecules to nucleic acids, such as groove binding or electrostatic interactions, can lead to other applications.

We hypothesize that tetrandrine fits into the major groove of dsDNA, and through the addition of substituents on the aromatic rings, we can improve its binding to DNA, thus leading to other applications.
nitrogen atoms the resulting positive charges facilitate the recognition of nucleic acids by means of the phosphate backbone. Herein, we report on dsDNA binding studies of a series of cationic bis-N-substituted tetrandrine derivatives. The substituents chosen for introduction into the alkaloid skeleton are polyaromatics commonly used for DNA intercalation and groove binding and allow evaluation of the effect of these units as well as the presence of heteroatoms. Spectrophotometric titrations, viscometry, electrophoretic mobility shift assays, and molecular modeling studies were carried out to determine the binding mode of the tetrandrine derivatives to the double helix of model dsDNA. In addition, viability MTT assays and complementary cyclic voltammetry studies were performed to examine the effect of the substituents over the cytotoxicity and gain information for further modifications of natural tetrandrine that can facilitate the development of new treatments and even fluorescent dsDNA probes for microscopic analysis.

2. RESULTS AND DISCUSSION

2.1. Synthesis and Characterization. The bis-N-substituted derivatives BBT, BNaphT, and BAqT were prepared in moderate to good yields as reported previously in the literature. In analogous manner, the novel tetrandrine salts BInT and BPyrT were synthesized by quaternization of the tertiary amine groups of the alkaloid with 3-(2-bromoethyl)indole and 1-(bromomethyl)pyrene, respectively (Figure 1).

The products were fully characterized by elemental analysis, IR, NMR spectroscopy (Figures S1–S3), and mass spectrometry. According to the 1H NMR spectra, BInT and BPyrT constitute single stereoisomers, showing a single set of signals for the hydrogen atoms (Figures S1 and S3). UV–vis absorption and emission spectra of each derivative can also be found in the Supporting Information (Figures S4 and S5). The emission spectra show the expected bands for the fluorophore substituents of the derivatives. An excimer band was not observed for the BPyrT derivative.

Additionally, BBT crystals suitable for single-crystal X-ray diffraction analysis were grown. Crystallographic data are given in the Experimental Section, and perspective views of the molecular structure of BBT, the asymmetric unit, and the unit cell are shown in Figures 2 and S6. The incorporation of benzyl units on the nitrogen atoms of the alkaloid produced two new chiral centers. Considering the known configuration of the asymmetric carbons of natural tetrandrine enabled us to

Figure 1. Chemical structures of tetrandrine and the bis-N-substituted tetrandrine salts studied in this work. Note: The chirality of the stereogenic centers in the isolated stereoisomers is indicated as S and R.

Figure 2. Perspective view of the molecular structure of one of the four crystallographically independent molecules of BBT (without the Br− counterions) extracted from the asymmetric unit of its crystal structure (C310H237Br8N9O24). The chirality of the stereogenic centers in the isolated stereoisomer is indicated as S and R.
establish the configuration for the new chiral centers as R giving the $S_3' S_2' R_3 R_4'$-tetrandrine derivative.

2.2. DNA Binding Studies. The chemical and structural characteristics of the five tetrandrine derivatives are appropriate for binding to dsDNA. To examine the DNA binding characteristics, spectrophotometric titrations, electrophoretic mobility shift assays, and viscometry experiments were performed along with molecular modeling studies on complexes formed with model DNA to elucidate possible binding modes.

2.2.1. Recognition Studies of dsDNA. Molecular recognition studies between the derivatives BNaphT, BAqT, BInT, and BPyrT (Figure 1) and oligonucleotide DNA-1 were carried out either by UV–vis or by fluorescence emission spectroscopy. This oligonucleotide was chosen to ensure a complete turn of the double helix while maintaining a simple system that includes all nitrogenous bases.

For all spectrophotometric titrations, the samples were dissolved in phosphate buffer with 10% DMSO (v/v) containing NaCl (0.01 M), adjusted to pH = 7.2. Only BPyrT was suitable for titrations using UV–vis absorption spectroscopy; meanwhile, the remaining three derivatives showed a significant increase of the baseline and broadened bands (Figure S7), which can be attributed to aggregation of the DNA–tetrandrine derivative complexes as a consequence of the high affinity, as evidenced by the titrations followed by fluorescence emission spectroscopy. It is noteworthy that fluorescence measurements allow detection at lower concentrations; therefore, aggregation was not observed for any of the titrations reported herein. In the UV–vis spectra of compound BPyrT, a bathochromic shift was observed upon DNA binding, as seen in Figure S8, with changes in $\lambda_{max}$ from 331 and 347 nm to 342 and 354 nm, respectively. This shift can be associated with $\pi$-stacking of the pyrene units with the DNA base pairs.

In a similar manner but using fluorescence spectroscopy, titrations of BNaphT, BAqT, and BInT with dsDNA were performed (Figures S9 and S10, Supporting Information), showing strong quenching of the fluorescence intensity for BInT, as seen in Figures 3 and S10, while for the case of BAqT, an increase in the emission of fluorescence intensity was observed (Figure S9). In the case of BNaphT, the experiments were not reproducible, which might be a consequence of the self-assembly characteristics of this compound as described previously.3

To gain better insight into the supramolecular interactions involved in the binding of the tetrandrine derivatives to dsDNA, the thermodynamic parameters $\Delta H$ and $\Delta S$ were calculated from the binding constants determined at different temperatures for the complexes with DNA-1, using the van’t Hoff equation (Table S1 and Figures S8–S11, Supporting Information). BPyrT and BInT gave favorable enthalpic and entropic contributions to the Gibbs free energy (Table 1). The negative $\Delta H$ values are attributed to intercalation into the base pairs of DNA,35 particularly by means of the pyrene and indole units, which are susceptible to $\pi$-stacking interactions. In addition, the rest of the molecule can establish electrostatic ion-pair interactions with the DNA phosphate backbones and van der Waals interactions with the major groove in the double helix. The binding of tetrandrine derivative cations is associated with the displacement of cations originally surrounding the phosphate backbone and a hydrophobic effect, with a consequent increase of the entropy in the system. Haq and Chaires have established that the binding of classical intercalators to dsDNA is enthalpically driven, while complexes with groove binders are entropically favored.36,37 For BAqT, the binding constants gave a nonlinear temperature dependence of $K_w$.38 Binding constant was determined by UV–vis.

### Table 1. Binding Constants ($K_w$) at 25 °C and Thermodynamic Parameters for the Complexes Formed between DNA-1 and the Tetrandrine Derivatives Studied Herein

| derivative | $K_w$ (M$^{-1}$) | $\Delta H$ (J/mol) | $\Delta S$ (J/(mol K)) |
|------------|-----------------|--------------------|-----------------------|
| BAqT       | $1.85 \times 10^5 \pm 2.23 \times 10^4$ | $-23.865$ | $30.7$ |
| BPyrT      | $6.16 \times 10^4 \pm 6.42 \times 10^4$ | $-7464$ | $127.7$ |
| BInT       | $9.53 \times 10^4 \pm 4.36 \times 10^4$ | $-23.865$ | $30.7$ |

DNA-1, 5'-GTA AGA TTC-3' and 5'-GAA TCA TCT TAC-3'.39,40 Binding constants were determined by fluorescence. Parameters could not be established due to a nonlinear temperature dependence of $K_w$.41 Binding constant was determined by UV–vis.
involving significant protein conformational changes.\textsuperscript{37–39} The formation of BAqT–DNA-1 aggregates is probably more likely to occur since the oligonucleotide is not large enough to present major conformational changes.

2.2.2. Ethidium Bromide Displacement Assays. To further evaluate the binding phenomena of bis-N-substituted tetrandrine derivatives with dsDNA, ethidium bromide displacement experiments were carried out, using oligonucleotides DNA-1, DNA-2, and DNA-3. Ethidium bromide displacement assays are generally accepted as an indirect method to identify intercalation in dsDNA. However, groove binders can displace ethidium bromide as well, and therefore, various experiments are needed to establish the definite mode of binding.\textsuperscript{37} The emission of the initial complex formed between dsDNA and ethidium bromide was quenched by all tetrandrine derivatives, revealing a particularly strong quenching in the case of BAqT. As an illustrative example, Figure 4 shows the results of the ethidium bromide displacement assays of BPyrT with the oligonucleotides constituted of adenine–thymine (DNA-2) or cytosine–guanine (DNA-3) nitrogenous bases; additional material is shown in Figures S12–S15.

Table 2 summarizes the Stern–Volmer constants ($K_{SV}$) obtained for the dsDNA complexes. As expected, the Stern–Volmer constants for complexes of DNA-1 are higher for all the tetrandrine derivatives compared to natural tetrandrine. Furthermore, in all derivatives, the Stern–Volmer constant was slightly larger for DNA-2 than for DNA-3, suggesting a small preference for oligonucleotides with adenine–thymine enriched sequences. In addition, ethidium bromide displacement experiments were performed with calf-thymus DNA (ctDNA) to ensure that the length of genomic DNA does not affect the magnitude of the

$\lambda_{exc} = 510$ nm, in phosphate buffer with 10% DMSO (v/v), pH = 7.2, 0.01 M NaCl, at 25 °C. $[\text{EtBr}] = 5$ μM, $[\text{dsDNA}] = 10$ μM, $[\text{BPyrT}] = 0–20$ μM.

Table 2. Stern–Volmer Constants ($K_{SV}$) for Complexes of Tetrandrine and Its Derivatives Studied Herein with DNA-1, DNA-2, and DNA-3

| compound  | DNA-1   | DNA-2   | DNA-3   |
|-----------|---------|---------|---------|
| tetrandrine | $7.34 \times 10^3$ | $b$     | $b$     |
| BNaphT    | $1.69 \times 10^4$ | $1.45 \times 10^4$ | $6.76 \times 10^3$ |
| BAqT      | $1.33 \times 10^5$ | $5.17 \times 10^4$ | $1.53 \times 10^4$ |
| BPyrT     | $4.84 \times 10^4$ | $1.34 \times 10^4$ | $8.26 \times 10^3$ |
| BInT      | $2.72 \times 10^4$ | $2.32 \times 10^4$ | $1.08 \times 10^4$ |

$^a$DNA-1, 5′-GTA AGA TGA TTC-3′ and 5′-GAA TCA TCT TAC-3′; DNA-2, 5′-poly(A)$_{18}$-3′ and 5′-poly(T)$_{18}$-3′; DNA-3, 5′-poly(G)$_{18}$-3′ and 5′-poly(C)$_{18}$-3′. Complexes were formed in phosphate buffer with 10% DMSO (v/v), pH = 7.2, 0.01 M NaCl, at 25 °C. $^b$Not determined.
occurred. Calf thymus DNA (ctDNA) was chosen for DNA decreases, bending of the double strand is likely to have do not alter the DNA length and viscosity. If the viscosity of the gel is decreased when bound to another molecule.38 Figure 5 illustrates clearly that BNaphT and natural tetrandrine reduce the mobility of the plasmid. Although some migration is still observed, it can be inferred that both compounds bind to DNA. On the other hand, BPyrT and BlnT seem to have a stronger interaction with DNA, as observed through spectrophotometric titrations, and prevent plasmid migration. Nicking of the plasmid was not observed. The BAgT-plasmid complex did not enter the pores of the agarose gel and, therefore, is not presented in Figure 5.

2.2.4. Viscometry Experiments. In the absence of crystallographic data, viscometry is considered one of the most sensitive and reliable techniques to determine the binding mode of small molecules to DNA. Intercalation lengthens the DNA double strands and produces an increase in viscosity relative to uncomplexed DNA,39,40 meanwhile, groove binders do not alter the DNA length and viscosity. If the viscosity of DNA decreases, bending of the double strand is likely to have occurred.41 Calf thymus DNA (ctDNA) was chosen for viscometry experiments since the solutions of the previously used oligonucleotides were not suitable for the Micro-Ubbelohde viscometer; EtBr, a classic intercalator compound, was used as a control test. BNaphT, BPyrT, and BlnT decrease the viscosity of ctDNA indicating that the double strand is bent as a consequence of the supramolecular interactions with the tetrandrine derivatives (Figure 6). This is plausible when considering that one of the N-aryl substituents can intercalate into DNA, while the rest of the molecular skeleton interacts with the major groove. Such behavior can explain the strong conformational contribution to the binding of BPyrT and BlnT (Table 1) with dsDNA and can be classified as a nonclassical intercalation. Finally, with BAgT the relative viscosity seems to remain unchanged; however, since only two concentrations of BAgT could be evaluated due to the low solubility at higher concentrations, the data points are insufficient to reach a conclusion in this case. Experiments such as dynamic light scattering could clarify whether BAgT–ctDNA complex aggregates result in the detected solubility limitations.

2.3. Molecular Modeling of the DNA–Tetrandrine Complexes. To get a better understanding of the complexes formed between DNA-1 (containing all nucleobases) and tetrandrine derivatives, molecular modeling studies were performed to propose a good representation of the binding mode of these complexes, based on the thermodynamic and viscometry binding studies. Figure S16 shows the geometry-optimized molecular gas phase structures of tetrandrine derivatives, while Figure 7 illustrates the lowest-energy structures of the complexes between DNA-1 and derivatives BlnT and BPyrT (the remaining are shown in Figure S17), determined by molecular docking that was carried out using molecular mechanics (AMBER99 force field) and the HyperChem release 8.0 software.42 In the calculated minimum-energy structures, one of the aromatic substituents of BlnT and BPyrT is tightly bound through π-stacking among the base pairs, while the macrocycle skeleton is localized in the major groove, displaying a series of noncovalent contacts including electrostatic, van der Waals, and hydrophobic interactions. Of these, the electrostatic interactions are formed among positively charged nitrogen atoms of the BlnT and BPyrT derivatives and negatively charged oxygen atoms at the phosphate backbones of the dsDNA. The short N−···O− distances of 3.87, 3.94, 3.64, and 4.69 Å for the case of DNA-1−BPyrT, as well as 3.78 and 4.24 Å for DNA-1−BlnT, are typical for strong ion-pair interactions.25,26 Similar results were obtained for the calculated minimum-energy structure of DNA-1−BNaphT (Figure S17b), while the structure of DNA-1−BBT (Figure S17a) shows that BBT, unable to intercalate, is positioned in the major groove with short N−···O− distances.

Finally, a very interesting result was obtained for the minimum-energy structure of the complex formed between DNA-1 and BAgT. As shown in Figure S17c, the derivative has a very good fit within the major DNA groove without intercalation of any of the anthraquinones; other calculated structures showed at least partial intercalation of one anthraquinone although with higher energies.

The arrangements illustrated in Figures 7 and S17 are in good agreement with the data established from the experimental binding and viscometry studies, as well as with
the depth, width, electrostatic potential, and steric effects of the DNA major groove. Generally, small molecules are localized in minor grooves, while larger molecules bind to the major groove even though it involves at least partial intercalation.

### 2.4. Cytotoxicity Assays

Since DNA binding is usually linked to cytotoxicity, we evaluated the biological activity of the bis-N-substituted tetrandrine derivatives studied herein, including BBT, and unmodified tetrandrine for comparison purposes, against the human cancer cell lines HeLa (cervical cancer), LS-180 (colon cancer), and A-549 (lung cancer), and the nontumor ARPE-19 (retinal epithelium) cell line.

As shown in Table 3, three of the derivatives revealed low cytotoxicity after 48 h of incubation. BPyrT, which carries the largest aromatic residues, is significantly more cytotoxic than the remaining derivatives, with IC<sub>50</sub> values against the LS-180, A-549, and ARPE-19 cell lines that are comparable to those of natural tetrandrine. This effect is probably related to better intercalation as seen formerly for MAnT, a tetrandrine derivative substituted with one anthracene unit, previously reported by our group.

Complementary electrochemical studies were performed to characterize the redox-active tetrandrine derivatives (Figure S18). The cyclic voltammograms of the tetrandrine derivatives with aryl substituents that do not possess heteroatoms, BNaphT and BPyrT, revealed each a single reduction and oxidation peak, with more negative <i>E<sub>1/2</sub></i> than BInT and BAqT. The peak current ratio (i<sub>pa</sub>/i<sub>pc</sub>) of the former approaches unity, indicating that the redox behavior is chemically reversible (Table S3). These parameters suggest that the derivatives described here do not possess any particular electrochemical properties indicating that they are oxidizing agents that could explain differences in cytotoxicity. Interestingly, and in contrast to tetrandrine, only low cytotoxicity was observed for BPyrT against HeLa, which deepens our drive to elucidate the

Table 3. Cytotoxicity of the Bis-substituted Tetrandrine Derivatives Studied Herein against Human Cancer Cell Lines and Retinal Epithelium Cells<sup>a</sup>

| compound | HeLa  | LS-180          | A-549        | ARPE-19          |
|----------|-------|-----------------|--------------|------------------|
| tetrandrine | 16.64 ± 0.02 (26.72 ± 0.03) | 6.43 ± 0.34 (10.33 ± 0.55) | 12.07 ± 1.51 (19.38 ± 2.42) | 8.45 ± 0.19 (13.57 ± 0.30) |
| BBT      | >30.00* (>31.10) | >30.00* (>31.10) | >30.00* (>31.10) | >30.00* (>31.10) |
| BNaphT   | >30.00* (>28.18) | >30.00* (>28.18) | 24.23 ± 0.29 (22.76 ± 0.27) | 21.10 ± 1.42 (19.82 ± 1.33) |
| BPyrT    | >30.00* (>24.66) | 5.84 ± 0.72 (4.80 ± 0.59) | 17.47 ± 1.05 (14.36 ± 0.86) | 9.23 ± 1.97 (7.59 ± 1.62) |
| BInT     | >30.00* (>28.02) | >30.00* (>28.02) | 28.25 ± 3.00 (26.39 ± 2.80) | 28.25 ± 1.49 (26.39 ± 1.39) |
| BAqT     | >30.00* (>24.50) | >30.00* (>24.50) | >30.00* (>24.50) | >30.00* (>24.50) |

<sup>a</sup>Data are shown as the mean IC<sub>50</sub> ± SD (standard deviation) in μg/mL (μM ± SD in parentheses) from three independent repetitions after 48 h exposure to the test compounds. The asterisk (*) represents the maximum concentration tested, which did not reach an IC<sub>50</sub> value.
underlying mechanisms to achieve specificity through proper selection of the substituents attached to the periphery of tetranda. Chemotherapeutic potential is generally valued for small molecules that bind to nucleic acids; however, the lack of relevant cytotoxicity of these tetrandrine derivatives could be an advantage if used as DNA probes.

3. CONCLUSIONS

The binding properties of a series of bis-N-substituted tetrandrine derivatives were examined using various double-stranded DNA models. The molecular recognition studies and electrophoretic mobility shift assays showed that cationic tetrandrine derivatives functionalized with polyatomic units have high affinity to dsDNA. Variations of the type of aryl substituents on the nitrogen atoms of the macrocyclic alkaloid allow the modulation of the physicochemical properties by varying the supramolecular interaction pattern and binding mode toward dsDNAs. These derivatives appear to bind to dsDNA through a dual binding mode involving partial intercalation of one substituent unit and accommodation of the remaining macrocyclic skeleton in the major groove of the double helix.

While the presence of heteroatoms on the substituent increases the binding constant to dsDNA, it does not necessarily result in higher cytotoxicity. As a matter of fact, viability MTT assays against tumor and nontumor cell lines did not necessarily result in higher cytotoxicity. As a matter of fact, tetrandrine macrocycle and the ten aromatic hydrogen atoms 

4. EXPERIMENTAL METHODS

4.1. General Procedures and Materials. 4.1.1. Reagents.

All reagents (including natural tetrandrine) were purchased from Sigma-Aldrich and used without further purification. Desalted oligonucleotides were purchased from Integrated DNA Technologies. All solutions were prepared in deionized water. Desalted oligonucleotides were purchased from Integrated DNA Technologies. All reagents (including natural tetrandrine) were purchased from Sigma-Aldrich and used without further purification. Desalted oligonucleotides were purchased from Integrated DNA Technologies. All solutions were prepared in deionized water.

4.1.2. Instruments. NMR studies were carried out with a Bruker Avance III 400 spectrometer, using TMS as reference (δ H = 0). Mass spectra were recorded on an Agilent 6100 LC/MS instrument using the ES^+ mode. UV-vis and fluorescence spectra were performed on Agilent 8435 diode array and PerkinElmer LS50B equipment using quartz cells of 1 cm optical path. The cuvettes were thermostatted at 25 °C through circulating water with a Thermo Scientific SC100 Circulating Bath. Melting points were recorded on a Büchi melting point B-545 instrument.

4.2. Preparation of Tetrandrine Derivatives. Bis-N-substituted tetrandrine derivatives BBT, BNNaphT, and BAqT were prepared following the synthetic procedures reported previously. The syntheses of the novel compounds BInT and BPyrT are described below.

4.2.1. Synthesis of BInT (C_{68}H_{62}Br_{2}N_{4}O_{4}). S,S-(−)-Tetrandrine (0.180 g, 0.289 mmol) and 3-(2′-bromomethyl)naphthalene (0.136 g, 0.606 mmol) were refluxed and stirred during 48 h in acetone (60 mL). The reaction was monitored by TLC until all tetrandrine was consumed. The resulting reaction mixture in the form of a suspension was rota-evaporated until dryness, whereupon ethyl acetate was added. A precipitate formed and was washed with the same solvent, and dried under vacuum to yield the product as a colorless solid (0.29 g, yield 93.5%). Decomposition point: 219-222 °C. UV absorption (DMSO/water, 10:90 v/v), λ (nm), ε (dm^3 mol^-1 cm^-1): λ_{240} e = 18435; λ_{280} e = 15925.

Despite the large number of hydrogen atoms, the ^1H NMR spectra showed characteristic signals that could be assigned using two-dimensional experiments. The signals for the two N-CH₃ and four O-CH₃ groups were found at δ_{H} = 3.86 (s, 3H), 3.68 (s, 3H), 3.57 (s, 3H), 3.10 (s, 3H), 2.63 (s, 3H), and 2.56 (s, 3H) ppm. The signals at δ_{CH₃} = 5.13 (m, 2H), 4.74 (m, 2H), and 4.27 (m, 4H) were assigned to the N-Ch₃ and N-CHR hydrogen atoms. The ten aromatic hydrogen atoms of the tetrandrine macrocycle and the ten aromatic hydrogen atoms of the two indole units are found in the region of δ_{H} = 7.72-6.17 (m, 20H). Finally, the indole NH-group gave a characteristic single signal at δ_{H} = 10.97 (s, 2H). MS-ESI(+): m/z: 455.3 ([BInT - Br^-]^+), 100%. Elem. Anal. Calcdd for C_{68}H_{62}Br_{2}N_{4}O_{4}: C, 61.43; H, 6.13; N, 4.94. Found: C, 61.44; H, 6.13; N, 4.35.

4.2.2. Synthesis of BPyrT (C_{72}H_{64}Br_{2}N_{2}O_{6}). S,S-(−)-Tetrandrine (0.180 g, 0.289 mmol) and 1-(bromomethyl)pyrene (0.136 g, 0.606 mmol) were refluxed and stirred for 24 h in 60 mL of acetone/chloroform (1:1 v/v). The reaction mixture was filtered and rota-evaporated until dryness, whereupon acetone was added. The precipitate was washed with acetone, separated by filtration, and dried under vacuum to yield the product as colorless solid (0.213 g, yield 43%). Decomposition point: 203-207 °C. UV absorption (DMSO/water, 10:90 v/v), λ (nm), ε (dm^3 mol^-1 cm^-1): λ_{240} e = 80808; λ_{279} e = 43642; λ_{347} e = 34980; λ_{374} e = 48244.

Despite the large number of hydrogen atoms, the ^1H NMR spectra showed characteristic signals that could be assigned using two-dimensional experiments. The signals for the N-CH₃ and O-CH₃ groups were found at δ_{H} = 3.86 (s, 3H), 3.63 (s, 3H), 3.59 (s, 3H), 3.52 (s, 3H), 2.87 (s, 3H), and 2.45 (s, 3H) ppm. The signals at δ_{CH₃} = 5.59 (m, 1H), 5.64-5.53 (m, 2H), 5.43-5.27 (m, 2H), 5.09-5.00 (m, 2H), 4.68-4.32 (m, 3H) were assigned to the N-Ch₃ and N-CHR hydrogen atoms. The ten hydrogen atoms of the aromatic skeleton of tetrandrine were found at δ_{H} = 7.51 (d, J = 8.3 Hz, 1H), 7.35 (s, 1H), 7.00 (d, J = 8.3 Hz, 1H), 6.96 (s, 1H), 6.88 (m, 2H), 6.81 (m, 1H), 6.55 (d, J = 10.2 Hz, 1H), 6.36 (s, 1H), and 6.24 (s, 1H) ppm. Finally, the hydrogen atoms of the pyrene units gave a series of signals in the range from 8.0 to 9.0 ppm that integrate for 18 hydrogen atoms. MS-ESI(+): m/z: 526.5 ([BPyrT - 2Br^-]^+) 100%. Elem. Anal. Calcdd for C_{72}H_{64}Br_{2}N_{2}O_{6}: C, 61.43; H, 6.13; N, 4.94. Found: C, 61.44; H, 6.13; N, 4.35.
4.3. Single-Crystal X-ray Diffraction Analysis. Single crystals of BBT (Figure 1) were grown by slow solvent evaporation from solution in acetonitrile at room temperature. Single-crystal X-ray diffraction data were collected at T = 100 K on an Agilent Technologies SuperNova diffractometer equipped with a CCD area detector (EosS2) using Cu Kα radiation (λ = 1.54184 Å) from a microfocus X-ray source and an Oxford Instruments Cryojet cooler. The measured radiation (equipped with a CCD area detector (EosS2) using Cu Kα evaporation from solution in acetonitrile at room temperature. Crystals of BBT (Figure 1) were grown by slow solvent evaporation from solution in acetonitrile at room temperature. Several of which indicated the presence of acetonitrile. Of BBT, a large number of additional peaks were detected, Fourier maps and assigned to the bromine counterions.

Prior to the refinement the SOLVENT MASK methodology 47 was used to achieve the final data set. Analysis of the solvent accessible volume gave a total of 6693.2 Å³ corresponding to 28.6% of the unit cell volume (electron count = 806). Figures have been created with Diamond. 48 Selected crystallographic data for BBT are shown in Table 4.

**Table 4. Selected Crystallographic Data for BBT**

| Formula         | C72H64Br2N2O6·4H2O |
|-----------------|---------------------|
| MW (g mol⁻¹)    | 16387               |
| Space group     | P2₁2₁2₁             |
| T (K)           | 100(3)              |
| a (Å)           | 24.27507(18)        |
| b (Å)           | 28.6715(2)          |
| c (Å)           | 33.1633(3)          |
| V (Å³)          | 23081.8(3)          |
| Z               | 4                   |
| μ (mm⁻¹)        | 2.119               |
| R Å             | 0.0681              |
| R Å (Å)         | 0.1868              |
| GOF             | 1.03                |

Crystalllographic data for the crystal structure have been deposited with the Cambridge Crystallographic Data Centre as supplementary publication no. 2123405. Copies of the data can be obtained free of charge on application to CCDC, 12 Union Road, Cambridge CB2 1EZ, UK (fax (+44)1223-336-033; e-mail deposit@ccdc.cam.ac.uk; http://www.ccdc.cam.ac.uk).

4.4. Spectrophotometric Titrations. Prior to the titration experiments, the molar absorptivities were obtained at the absorption maximum of the spectrum of each tetrandrine derivative (Figure S4), ensuring a linear concentration dependence of the absorbance. The fluorescence spectra of the derivatives (Figure S5) are in agreement with the known spectral characteristics of each of the fluorophore units.

Spectrophotometric titrations were performed with 10⁻⁶ to 10⁻³ M solutions of the corresponding tetrandrine derivative dissolved in 0.0064 M phosphate buffer with 10% DMSO (v/v), pH = 7.2, 0.01 M NaCl. There was no visible aggregation under such conditions, and the chosen concentration for the titration of any derivative falls in the linear response of the observed property. For the recognition experiments with the double-stranded DNA models, the respective oligomer was mixed in the same medium with its complementary sequence and heated for 5 min to 98 °C in a Thermo Scientific dry bath/block heater, followed by slow cooling to room temperature. The following dsDNA sequences were used: DNA-1, 5’-GTA TCA TCT TAC-3’ and 5’-GAA TCA TCT TAC-3’; DNA-2, 5’-poly(A)₈-3’ and 5’-poly(T)₈-3’; DNA-3, 5’-poly(G)₈-3’ and 5’-poly(C)₈-3’.

Binding constants for the complexes formed between the tetrandrine derivatives and the model dsDNA were obtained by recording changes of the absorption or emission spectrum of a fixed amount of the tetrandrine derivative dependent on increasing concentrations of added dsDNA. All UV–vis and fluorescence spectrometric titration curves were fitted with the Microlab OriginPro8 software package using the following equation, which here is expressed in terms of absorbance and is analogous for emission intensity:

\[
A_{\text{obs}} = \frac{A_0 + A_{\infty} K_b [L]}{1 + K_b [L]}
\]

where \(A_{\text{obs}}\) is the observed absorbance, \(A_0\) is the initial absorbance, \(A_{\infty}\) is the maximum absorbance change, \(K_b\) is the binding constant, and \([L]\) is the total concentration of ligand, in this case dsDNA. The nonlinear curve fitting of the data was compared with the classical Scatchard linear plot, which is commonly used in dsDNA binding studies in combination with the McGhee and von Hippel equation. 26 As expected, the same results were obtained without dependence on the fitting model employed.

4.5. Ethidium Bromide Displacement Assays. For the ethidium bromide (EtBr) displacement assays, increasing amounts of a tetrandrine derivative stock solution were added to a solution of the dsDNA–EtBr complex ([DNA] = 10 μM and [EtBr] = 5 μM) in 0.0064 M phosphate buffer with 10% DMSO (v/v), pH = 7.2, 0.01 M NaCl. The fluorescence spectra of EtBr bound to dsDNA were measured using an excitation wavelength of 510 nm. The resulting titration data were plotted according to the Stern–Volmer equation to obtain the quenching constant, \(K_{SV}\).

\[
\frac{I_0}{I} = K_{SV}[Q] + 1
\]

where \(I_0\) and \(I\) are the emission intensities of the dsDNA–EtBr complex in the absence and in the presence of quencher (Q), respectively. \(K_{SV}\) corresponds to the slope obtained from the plot of \(I/I_0\) versus [Q].

4.6. Electrophoretic Mobility Shift Assays. Samples of plasmid DNA pCR 2.1 (100 ng, Thermo Scientific) containing a cloned DNA sequence of a total of 4.5 kb were combined with the absorption maximum of the spectrum of each tetrandrine derivative (Figure S4), ensuring a linear concentration dependence of the absorbance. The fluorescence spectra of the derivatives (Figure S5) are in agreement with the known spectral characteristics of each of the fluorophore units.
with 0.1 nmol of the respective tetrandrine derivative in a solvent mixture of DMSO and deionized water 10:90 (v/v) and incubated for 2 h at 37 °C. The samples were loaded into 1% agarose gel, and their electrophoretic mobility was examined after electrophoresis for 30 min at 60 V. After staining the gel under gentle agitation for 5 min with ethidium bromide (5 μg/mL), the samples were washed with water for 2 h, and then revealed with UV-light using a Gel Doc EZ documentation system from Bio-Rad. The solvent mixture and a solution of plasmid only were used as controls.

4.7. Viscosity Measurements. Viscosity studies of complexes formed between the tetrandrine derivatives and calf thymus DNA (ctDNA) were carried out with a Micro-Ubbelohde viscometer from SI Analytics at a temperature of 20 °C, which was maintained with a PolyScience temperature controller. Ethidium bromide (EtBr), which is a classic intercalator compound, was used as a control test. For the experiment, the flow time of calf thymus DNA (ctDNA) was measured upon addition of the tetrandrine derivative. Each data point was established from ten measurements using a digital stopwatch. The relative viscosity of the ctDNA solution (ηr) and the tetrandrine–ctDNA complexes (η) were calculated according to the equation:

\[
\eta = \frac{(t - t_0)}{t_0}
\]

where \( t \) is the average flow time of the respective solution and \( t_0 \) is the average flow time of the medium (phosphate buffer with 5% DMSO v/v). The results were plotted as \( \eta/\eta_r \) against the [tetrandrine derivative]/[ctDNA] ratio.

4.8. Molecular Modeling. For the molecular modeling studies of the tetrandrine derivatives, the initial molecular structures were constructed by using the X-ray diffraction coordinates of BBT and those previously reported for BNaphT for the macrocycle skeleton, attaching when necessary the respective N-CH2-aryl substituents using the Avogadro software tool. The tetrandrine derivatives were then geometry-optimized in the gas phase by using the PM6 semiempirical method implemented in the Gaussian 09 program package. The structure of double-stranded B DNA-1 was generated from the nucleic acid database of HyperChem release 8.0 software and optimized in the gas phase by applying the AMBER99 force field and a Polak–Ribiere conjugate algorithm with a convergence criterion of 0.001 kcal/(Å mol). For the molecular modeling of the DNA–tetrandrine derivative complexes, the optimized conformer of the tetrandrine derivative was manually docked with DNA-1 based on the findings obtained from the titration and viscosity studies. Geometry optimizations of the complexes with DNA were carried out using molecular mechanics (AMBER99 force field) and the HyperChem release 8.0 software.

4.9. Cytotoxicity Assays. Cytotoxicity study tests of the bis-N-substituted tetrandrine derivatives studied herein were performed through MTT assays on the human cancer cell lines HeLa (cervical cancer), LS-180 (colon cancer), and A-549 (lung cancer). For comparison, the nontumoral cell line ARPE-19 (retinal epithelium) was also included. The cells were initially cultured in a 96-well sterile plate with DMEM medium supplemented with 5% FBS at a density of 10000 cells per well. Different concentrations of the tetrandrine derivatives were added, followed by 48 h incubation. After that time, 10 μM MTT [3-(4,5-dimethyl-2-thiazolyl)-2,5-diphenyltetrazolium bromide] dissolved in DMSO was added to each well followed by incubation for 4 h at 37 °C. Acidic isopropanol was added under stirring to dissolve formazan crystals, and after 10 min, the plate was analyzed in a Bio-Rad ELISA plate reader, with 750 nm being the test wavelength and 630 nm the reference wavelength. Control experiments were performed in parallel. Cytotoxicity was reported as the 50% inhibitory concentration, IC50. All experiments were carried out in triplicate.

The data are reported as mean IC50 ± SD (standard deviation) and were subjected to statistical analysis of variance (ANOVA), comparing the mean values with the Tukey test (p < 0.05). For the statistical analyses, the IBM® SPSS® 20 Program was used.

4.10. Cyclic Voltammetry Studies. Cyclic voltammograms (CV) were recorded at room temperature in a conventional three-electrode cell, using a platinum wire as the auxiliary electrode and Ag/AgCl (KBr, 3 M) as reference electrode. Measurements were performed in DMSO solutions containing 0.20 M tetrabutylammonium hexafluorophosphate (NBu4PF6) as supporting electrolyte, using a Metrohm PGSTAT 128N equipment. Previously, the working electrodes were prepared by evaporating a drop (5.0 μL) of a suspension of the corresponding tetrandrine derivative in acetonitrile (1 mg/mL) located on a glassy carbon electrode (GCE, BAS MF 4012, exposure area 0.071 cm²). CVs were collected with scan rates of 100 mV/s with iR compensation.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsomega.2c00225.

NMR spectra of BlnT and BPyrT, UV–visible absorption and emission spectra of BNaphT, BPyrT, BAqT, and BlnT, perspective views of the asymmetric unit and the unit cell of BBT, spectroscopic titrations and fitting curves for all derivatives, as well as Stern–Volmer constants, and binding constants for BPyrT, BAqT, and BlnT at different temperatures with corresponding van’t Hoff plots, geometry-optimized molecular gas phase structures of BBT, BNaphT, BPyrT, BAqT, and BlnT, lowest-energy structures of the complexes between DNA-1 and derivatives BBT, BNaphT, and BAqT, and cyclic voltammograms and electrochemical parameters of BNaphT, BPyrT, BAqT, and BlnT (PDF)

**AUTHOR INFORMATION**

Corresponding Author

Karen Ochoa Lara – Departamento de Investigación en Polímeros y Materiales, Universidad de Sonora, 83000 Hermosillo, Sonora, México; orcid.org/0000-0002-5271-2718; Phone: +52 (662) 259 2161; Email: karenol@polimeros.unison.mx, karen.ochoa@unison.mx

Authors

Sandra Mónica González-Martínez – Departamento de Investigación en Polímeros y Materiales, Universidad de Sonora, 83000 Hermosillo, Sonora, México

Drochss Pettry Valencia-Ochoa – Departamento de Ciencias Naturales y Matemáticas, Facultad de Ingeniería y Ciencias, Pontificia Universidad Javeriana, 760031 Cali, Colombia
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