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Novel poly(ADP-ribose) polymerase inhibitor veliparib: biophysical studies on its binding to calf thymus DNA

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Veliparib (ABT-888), which can inhibit cancer growth by blocking DNA base excision repair, is one of several recently developed oral inhibitors of poly(ADP-ribose) polymerases, which are currently used in clinical trials. In this work, interaction of calf thymus DNA (ctDNA) with ABT-888 was first investigated following UV-visible absorption, nuclear magnetic resonance (NMR) spectroscopy, steady-state and time-resolved fluorescence, viscosity measurements, circular dichroism (CD), and Fourier transform infrared (FT-IR) spectroscopy coupled with molecular docking methods. UV-visible absorption indicated that ABT-888 was indeed bound to ctDNA. Broadening and upfield shift of the proton peaks of ABT-888 in the proton NMR spectrum indicated that ABT-888 interacted with ctDNA primarily by partial intercalation. Fluorescence quenching and time-resolved fluorescence spectroscopy studies showed that binding of ABT-888 with ctDNA occurred through a static quenching mechanism, resulting in the formation of a ctDNA-ABT-888 complex. Thermodynamic calculations revealed that interaction was an enthalpy-driven process caused by hydrogen bonds and van der Waals forces. Competitive fluorescence experiments coupled with viscosity, CD, and FT-IR studies revealed that ABT-888 intercalates partially and binds to the groove, phosphate group, and deoxyribose sugar of ctDNA and also induces conformational changes. Molecular docking showed that ABT-888 preferably binds to the DNA groove. However, other types of binding, including classic intercalation and partial intercalation, cannot be ruled out.

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1. Introduction

Currently, cancer remains a leading cause of death in countries around the world. A majority of anti-cancer drugs cause cell death through DNA damage, and in this situation, functioning DNA repair pathways act as mechanisms of drug resistance.^{1,2} Cancer cells can activate their own DNA damage repair to restore themselves, thus causing drug resistance against anti-cancer drugs and radiotherapy.^{3,4} Further research indicates that there is over-expression of DNA repair pathways in cancer cells. Thus, blocking DNA damage repair pathways is an important and new method of cancer therapy. Polyadenylation diphosphate ribose transferase (poly ADP-ribose polymerase, PARP) is a DNA repair enzyme, which plays a key role in DNA repair pathways. The superfamily of PARPs consists of six different enzymes: PARP-1, PARP-2, PARP-3, PARP-4/VPARP, tankyrase-1 and tankyrase-2.⁵ These enzymes engage in various cellular functions, including cell cycle regulation, transcription, and DNA damage repair.⁶ Thus, PARP inhibitors could be developed as chemosensitizers of DNA-damaging agents. PARP-1 is currently the best characterized enzyme in

this group and is critical to repair of breaks in single-strand DNA through base excision repair pathway.^{5,7}

Veliparib (also called ABT-888) is a novel oral benzimidazole derivative and is the most represented PARP-1 inhibitors.⁸ Fig. 1 shows the complete chemical name of the compound (2-[(R)-2-ethylpyrrolidin-2-yl]-1*H*-benzimidazole-4-carboxamide) and its chemical structure. *In vivo* and *in vitro* experiments confirmed that ABT-888 increases cellular apoptosis and consequently inhibits cancer growth *via* blocking DNA base excision repair.⁹⁻¹¹ Efficacy and safety data from recent phase I and II trials demonstrated that ABT-888, in combination of cytotoxic agents or as monotherapy, was well-tolerated and showed anticancer activity.^{12,13} In treatment of metastatic breast cancer, colon cancer, metastatic melanoma, and brain tumor, the compound has significant effects, and its combination with temozolomide is going to enter phase III clinical trials for treatment of breast cancer.¹⁴ Recent studies reported that ABT-888 also inhibits vessel formation in radiated cells.¹³ Many reports focus on functional targets of ABT-888, but none centered on detailed aspects of binding mode and binding region of ABT-888 and DNA and DNA structural change. Interestingly, investigation of interactive nature of ABT-888 with DNA can provide insight into drug anticancer activities involved



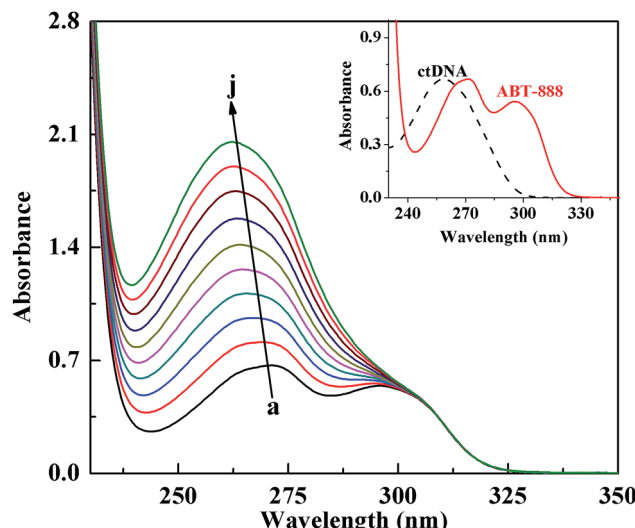


Fig. 1 UV-visible absorption spectra of ct-DNA-ABT-888 system in PBS buffer (pH 7.4). $C[ABT-888] = 8.0 \times 10^{-5} \text{ M}$; $C[\text{ctDNA}]/(a-j) = 0.0, 2.6, 5.1, 7.6, 10.1, 12.5, 15.0, 17.4, 20.1, 22.8 \times 10^{-5} \text{ M}$; inset: UV-visible absorption spectra of ABT-888 and ctDNA in PBS buffer. $C[ABT-888] = 8.0 \times 10^{-5} \text{ M}$; $C[\text{ctDNA}] = 7.6 \times 10^{-5} \text{ M}$; $T = 298 \text{ K}$.

in DNA damage repair pathways and provide support for continued clinical investigation of the drug.

Given the information above, this work aimed to investigate the mechanism of interaction (including binding mode, binding region, and structural feature) between ABT-888 and DNA using calf thymus DNA (ctDNA) *in vitro* under simulated physiological conditions. ctDNA has a relatively low protein content with a highly polymerized DNA skeleton which contains alternating sugar phosphate sequence.^{15,16} It is a natural DNA widely used in studies of binding between DNA and anticancer agents, which modulate DNA structure and function.^{17–19} It is also used in physicochemical studies of DNA behavior in solution. In the present study, UV-visible absorption and proton nuclear magnetic resonance (¹H NMR) investigations were carried out to explore whether ABT-888 binds to ctDNA in solution. The quenching mechanism, binding constants, number of binding sites, and thermodynamic parameters were evaluated by steady-state and time-resolved fluorescence spectroscopy. Ethidium bromide (EB) and Hoechst 33258 were used as DNA probes for comparative study of the interaction of ABT-888 with ctDNA helix, and comparison was further supported by viscosity measurements, circular dichroism (CD), and Fourier transform infrared (FT-IR) spectroscopy. Furthermore, molecular docking was used to interpret experimental observations and to propose a three-dimensional (3D) model for the binding region of ctDNA-ABT-888 complex.

2. Materials and methods

2.1. Materials

Veliparib (purity > 99%) was purchased from Selleck (USA) [<http://www.selleckchem.com>]. Highly polymerized ctDNA sodium salt was purchased from Sigma-Aldrich Corporation

(Milwaukee, USA) and was used without further purification. EB, Hoechst 33258, deuterium oxide (D_2O , 99.9% purity), and dimethyl sulfoxide-d6 (DMSO-d6) were purchased from J&K Scientific Ltd. (Beijing, China). All reagents were of analytical reagent grade, and thrice-distilled water were used throughout the experiment.

2.2. Sample preparation

ctDNA was dissolved in 10 mM phosphate buffer solution (PBS) of pH 7.4 and stored at 4 °C. Concentration of nucleotide was determined using UV-1800 Shimadzu ultraviolet (UV)-spectrophotometer (Japan). UV absorbance at 260 and 280 nm of ctDNA solution yielded a ratio of 1 : 1.9 ($\epsilon_{260} = 6600 \text{ L mol}^{-1} \text{ cm}^{-1}$), indicating that ctDNA was sufficiently free of protein.^{19,20} Stock standard solution ($5.0 \times 10^{-3} \text{ M}$) of ABT-888 was prepared by dissolving the compound in DMSO. $1.0 \times 10^{-3} \text{ M}$ EB and $1.0 \times 10^{-3} \text{ M}$ Hoechst 33258 were prepared in thrice-distilled water.

2.3. Apparatus and methods

2.3.1. UV-visible spectroscopy. Absorption spectra of ABT-888 and the ctDNA-ABT-888 complex were recorded on a UV-1800 Shimadzu ultraviolet (UV)-spectrophotometer (Japan) using 1 × 1 cm quartz cuvettes. Absorbance of ABT-888 alone and that of ctDNA-ABT-888 complex with various concentrations of ctDNA ($0\text{--}2.3 \times 10^{-4} \text{ M}$) and fixed concentration of ABT-888 ($8.0 \times 10^{-5} \text{ M}$) was determined at wavelength ranging from 220 to 350 nm.

2.3.2. ¹H NMR spectroscopy. Solution ¹H NMR spectra were recorded on a Bruker Avance-600 spectrometer at 298 K. 2.4 mg ABT-888 and 5.0 mL DMSO-d6 + D_2O (2 : 3) were mixed thoroughly and added into the NMR tube for ¹H NMR measurement of ABT-888. As a comparison, drug-DNA complex was prepared by mixing 2.4 mg ABT-888 and 2.8 mg ctDNA in 5.0 mL DMSO-d6 and D_2O (2 : 3). In all NMR experiments, 0.5 mL solutions were added into the NMR tube. ¹H NMR spectra were processed using TopSpin 3.5 software (Bruker BioSpin, Ltd.).

2.3.3. Steady-state fluorescence spectroscopy. Steady-state fluorescence spectra were recorded on Varian Cary Eclipse fluorescence spectrophotometer (Santa Clara, CA, USA) equipped with 1.0 cm quartz cells. Experiments were conducted in PBS buffer (10 mM, pH 7.4) and allowed to equilibrate thermally at 295/303/310 K at about 30 min before measurements. Fluorometric experiments were carried out by varying the concentrations of ctDNA (0 M to $1.3 \times 10^{-4} \text{ M}$) while fixing ABT-888 concentration at $5.0 \times 10^{-6} \text{ M}$. All solutions were excited at 271 nm, which is λ_{max} for free ABT-888, as identified by UV-visible spectroscopy. Emission spectra were recorded within wavelength range of 300 nm to 500 nm. Both excitation and emission slits were set at 5 nm. To eliminate the possibility of re-absorption and inner filter effect arising from UV absorption, fluorescence intensities were corrected for absorption of excited light and re-absorption of excitation light according to the following formula:^{21,22}

$$F_c = F_m e^{(A_1 + A_2)/2} \quad (1)$$

where F_c and F_m are the corrected and measured fluorescence intensities, respectively; A_1 and A_2 are the absorbance values of ABT-888 at excitation and emission wavelengths, respectively.

2.3.4. Time-resolved fluorescence spectra. Time-resolved fluorescence decay measurements were conducted by time-correlated single-photon counting technique in Horiba Jobin Yvon FluoroLog spectrofluorometer (HORIBA, Les Ulis, France). Time-resolved ABT-888-fluorescence quenching by ctDNA was recorded by fixing 271 nm as excitation wavelength and 360 nm as emission wavelength at room temperature. ABT-888 concentration was fixed at 4.0×10^{-5} M, and ctDNA concentration varied from 0 M to 2.5×10^{-4} M.

2.3.5. Competitive displacement assay. EB and Hoechst 33258 are normally used to probe DNA-binding mode in drug-DNA interactions.²³ EB displacement experiments were conducted by keeping EB (8.0×10^{-6} M) and ctDNA (1.2×10^{-4} M) concentration constant while varying ABT-888 concentrations from 0 M to 1.8×10^{-4} M. EB-ctDNA complex was excited at 481 nm, and fluorescence spectra were recorded between 550 and 750 nm. In a similar experiment, Hoechst dye displacement assay was conducted under similar concentrations of dye and ctDNA-like EB. In Hoechst 33258 displacement assay, Hoechst 33258-ctDNA complex emission spectra were recorded from 400 nm to 600 nm with fixed excitation at 343 nm.

2.3.6. Viscosity measurements. Viscosity experiments utilized a Discovery HR-2 hybrid rheometer (TA Instruments, New Castle, DE, USA) and TA Instruments TRIOS V3.3.1 software for data acquisition and analysis. The shear rate used was 50.0 s^{-1} , and duration was 100 s. Three measurements were used to calculate average viscosity of the samples. Viscosity values were corrected for buffer solution. Concentration of ctDNA (7.7×10^{-5} M) in solution was fixed while varying ABT-888 concentrations. Values of relative specific viscosity are presented as $(\eta/\eta_0)^{1/3}$ versus ABT-888-to-ctDNA ([ABT-888]/[ctDNA]) concentration ratios, where η is viscosity contributions of ctDNA in the presence of ABT-888, and η_0 is viscosity of ctDNA alone.

2.3.7. CD spectroscopy. Far UV-CD measurements (230–320 nm) were conducted on a Hitachi-F7000 fluorescence spectrometer, which used a quartz cell with path length of 0.1 cm at 298 K, equipped with R3788 photomultiplier. Concentration of ctDNA was kept at 3.5×10^{-4} M while ABT-888 concentration varied from 0 to 8.0×10^{-5} M. All observed CD spectra were corrected for buffer signal, and results were expressed as ellipticity. Each CD spectrum was scanned at 60 nm min^{-1} , and data points were recorded at 1 nm intervals for three-scan accumulations.

2.3.8. FT-IR spectra. IR spectra within $1850\text{--}900 \text{ cm}^{-1}$ with resolutions of 4 cm^{-1} and 128 scans were recorded on a Nicolet-6700 FT-IR (Thermo, US) spectrometer with a smart OMNI-sampler accessory. Background atmospheric spectrum was collected before each measurement. Corresponding absorbance contributions of the buffer and free ABT-888 solutions were recorded and digitally subtracted from the spectra of free ctDNA and ABT-888-ctDNA complex solutions.¹⁸ All spectra data were

analyzed using first derivative combined with Savitzky–Golay filter smoothing.

2.3.9. Molecular docking analysis. MGL tools 1.5.4 with AutoGrid4 and AutoDock4 (ref. 24 and 25) were used to set up and perform blind docking calculations between the drug and DNA sequence. The 3D structure of ABT-888 was constructed from a 2D structure and geometry was optimized using the Discover Minimization tool in Materials Studio 6.0 software (Accelrys Co. Ltd, California, USA). Crystallographic structure of DNA duplex was obtained from the Protein Data Bank, using identifier 1BNA [(CGCGAATTCGCG)2] (<http://www.rcsb.org/pdb>). Before docking analysis, DNA structure was modified to remove water molecules and to add polar hydrogen atoms and Gasteiger charges. DNA was enclosed in a box with number of grid points in x -, y -, and z -axes, $60 \times 68 \times 126 \text{ \AA}^3$, and grid spacing of 0.375 \AA . Grid center was set to positions of 14.780, 20.976, and 8.807. Parameters were used throughout docking. After grid map generation, 200 Lamarckian genetic algorithm runs were carried out using 250 000 energy evaluations and 27 000 generations. All other parameters were measured at default setting. The lowest energy docked conformation, according to AutoDock scoring function, was selected as binding mode.

3. Results and discussion

3.1. UV-visible absorption studies

The strength and nature of binding interaction between ABT-888 and ctDNA can be obtained from the absorption titration method by following the changes in the absorbance and shift in the wavelength of DNA-drug complex.^{26–28} Most of small molecules noncovalently interact with DNA through three distinct mode types: (I) intercalation, (II) partially intercalated binding, and (III) groove binding, which can be stabilized by hydrophobic, electrostatic, and hydrogen bonding interactions.^{18,29,30} The absorption spectra of ABT-888 in the absence and presence of increasing concentration of ctDNA were recorded and are shown in Fig. 1. The inset shows the characteristic maximum absorption of ctDNA and ABT-888 centered at 271 and 296 nm in the wavelength range of 230–350 nm, respectively. With the gradual addition of ctDNA ($0\text{--}22.8 \times 10^{-5}$ M) to the fixed concentration of ABT-888 (8.0×10^{-5} M), there was a continuous increase of the absorption spectrum of ABT-888. Fig. 1 clearly shows the occurrence of the hyperchromicity of absorbance at 271 nm along with a blue shift of about 9 nm. In addition, the absorption peak at 296 nm gradually disappeared with the increasing concentration of ctDNA. These changes may be a result of the interaction of ABT-888 with ctDNA. Some investigators have reported that the binding of small molecules to the base pairs of DNA through intercalation usually results in hypochromic (or hyperchromic) effect and a red shift of the small molecules absorption band.²⁰ Thus, our results show that ABT-888 is likely to interact with ctDNA through mixed-ligand type binding modes. Since the UV-visible spectroscopic results did not provide some defined evidence regarding the nature of binding between ABT-888 and ctDNA, some other experiments



were needed to confirm the strength and nature of this interaction and its clear-cut binding mechanism.

3.2. ^1H NMR spectroscopy of ctDNA-ABT-888 system

NMR technique is a leading tool for probing and understanding at molecular level the binding between a small molecule and its target receptor. Specifically, ^1H is the most commonly used atomic nuclei for DNA binding study given that it is not limited by molecular size and does not require isotope-labeling.³¹⁻³³ The aforementioned three distinct types of ligand-DNA interaction mode are easily differentiated by ^1H NMR signals based on changing line widths and chemical shifts.³¹ In type I binding, intercalation of molecules into base pairs of DNA results from total line broadening of ^1H NMR signal.³⁴ For type II binding, observation includes line broadening and upfield chemical shift of the signal. These changes may arise from two reasons: (1) weak restrictions of molecular tumbling in the DNA complex; (2) slow rate exchange between various DNA binding sites and the unbound state of molecules. For type III binding, binding of molecules with DNA grooves is not exhibited in line broadening and upfield shift of the ^1H NMR signal.

In this experiment, ^1H NMR measurements were carried out to understand how interaction mode is formed in the ctDNA-ABT-888 system. Fig. 2 shows ABT-888 ^1H NMR spectra obtained at 298 K in the absence and presence of ctDNA. Compared with Fig. 2A and B clearly shows line broadening of H-a, b, c, and d proton peaks after adding ctDNA, indicating that addition of ctDNA influences the microenvironment around ABT-888 protons.³⁵ Table 1 shows the upfield shift of proton resonances of a, b, c, and d hydrogen atoms in the presence of ctDNA, indicating that π - π stacking interactions between ABT-888 and base pairs of ctDNA possibly exist, and magnetic shielding effects resulted in increased electron density in the surroundings of these protons after binding. Zhang *et al.*³² suggested that magnetic shielding effects may be caused by hydrogen bond formation between small drug molecules and ctDNA. There was a phenomenon that changes in the chemical shifts of the

Table 1 Chemical shifts (ppm) of protons of ABT-888 in the system

System	H _a	H _b	H _c	H _d
ABT-888	8.045	7.973	7.611	3.642-3.700
	8.033	7.950	7.599	(multiple splitting peaks)
ctDNA-ABT-888	8.028	7.955	7.590	3.623-3.682 (multiple splitting peaks)

hydrogen of the benzimidazolyl moiety of ABT-888 were greater than those of the hydrogen of pyrrolidyl moiety in the presence of ctDNA (Table 1). This difference implies an unequal participation of the two moieties of ABT-888 in the formation of the complex and the benzimidazolyl moiety may be more intimate contact with ctDNA. Broadening and upfield shift of peaks of ABT-888 in the presence of ctDNA indicate that ABT-888 indeed bound to ctDNA through partial intercalation binding.

3.3. Analysis of fluorescence quenching

Fluorescence titration experiment was used to study relative binding of ABT-888 to ctDNA. Fluorescence signal was not observed for the sample with ctDNA alone in the buffer solution. However, ABT-888 exhibited strong fluorescence emission peak at 360 nm with excitation wavelength at 271 nm. In the present study, ABT-888 and ctDNA interaction was investigated by gradually adding ctDNA to ABT-888 solution (5.0 μM). Fig. 3A shows the remarkable decreasing trend of emission intensity of ABT-888 at 360 nm with increasing concentrations of ctDNA. Quenching rate was 53.5%, and there was a slight blue shift (2 nm) at the maximum wavelength of ABT-888 fluorescence emission when the solution of ctDNA was added. Sameena and Enoch²⁷ suggested that such a blue shift showed the increase in the hydrophobicity around the macromolecule due to the formation of new non-fluorescent complex between small molecule and macromolecule. Therefore, the results indicated that ctDNA could quench the intrinsic fluorescence of ABT-888, and that binding of this compound to ctDNA indeed exists. Results further support the findings of UV absorption and ^1H NMR.

Different quenching mechanisms are usually classified as dynamic or static quenching, and the two can be distinguished by analyzing fluorescence data at different temperatures. Decrease in fluorescence intensity is described by the Stern-Volmer equation:^{36,37}

$$F_0/F = 1 + K_{\text{SV}}[Q] \quad (2)$$

where F_0 and F denote fluorescence intensities in the absence and presence of quencher, respectively; $[Q]$ is the quencher concentration; K_{SV} is the Stern-Volmer quenching constant, which is determined by linear regression of a plot of F_0/F against $[Q]$.

Table 2 provides the K_{SV} obtained using the Stern-Volmer equation. By analyzing results of K_{SV} values, dynamic or static

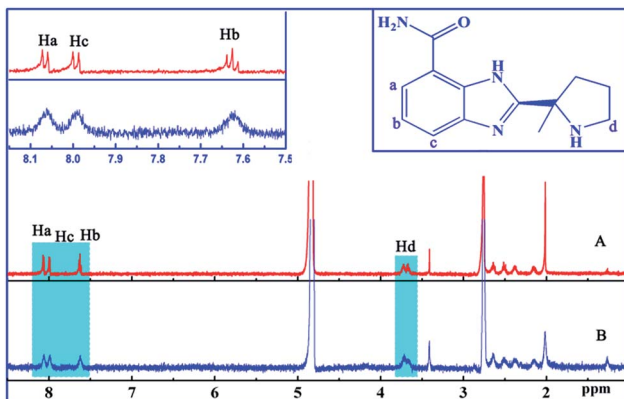


Fig. 2 ABT-888 ^1H NMR spectra in the absence (A) and presence (B) of ctDNA in DMSO-d6 : D₂O (2 : 3) at 298 K.



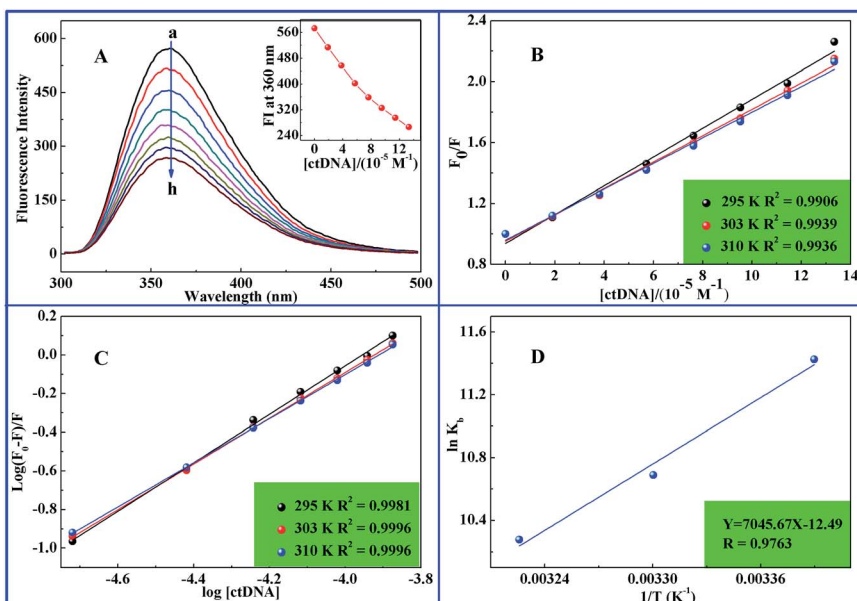


Fig. 3 (A) Fluorescence spectra of ABT-888 (5.0×10^{-6} M) in the absence and presence of various concentrations of ctDNA. (B) Stern–Volmer plots constructed on the basis of fluorescence quenched ABT-888 with increasing concentrations of ctDNA at three different temperatures. (C) Plot of $\log(F_0 - F)/F$ vs. $\log[ctDNA]$ for the ctDNA–ABT-888 system at three different temperatures. (D) Van't Hoff plot of ctDNA–ABT-888 system.

Table 2 Association constants and thermodynamic parameters at different temperatures for the ctDNA–ABT-888 interaction

T (K)	$K_{SV} (\times 10^3 \text{ M}^{-1})$	SD ^a	n	$K_b (\times 10^4 \text{ M}^{-1})$	SD ^a	$\Delta G (\text{kcal mol}^{-1})$	$\Delta H (\text{kcal mol}^{-1})$	$\Delta S (\text{cal mol}^{-1} \text{ K}^{-1})$
295	9.46	0.0113	1.25	9.17	0.0232	-6.68	-14.00	-24.80
303	8.66	0.0095	1.18	4.39	0.0352	-6.48		
310	8.40	0.0105	1.14	2.91	0.0331	-6.31		

^a SD is the standard deviation of three measurements.

quenching were differentiated. Dynamic quenching depends upon diffusion. Since the higher temperature leads to the larger diffusion coefficients, the K_{SV} increases with rising the temperature. In contrast, increased temperature is likely to cause the decrease of complex stability, and thus lower values of the static quenching constants were resulted.²⁰ Fig. 3B and Table 2 present the decreasing trend of K_{SV} values with increasing temperature. Therefore, fluorescence quenching mechanism of ABT-888 by ctDNA may be controlled by static rather than dynamic quenching along with ctDNA–ABT-888 complex formation.

To further substantiate the quenching mechanism, fluorescence lifetime measurements were carried out on a single-photon counting setup given that lifetime would be unaffected in ground-state quenching but would decrease in the excited-state type.³⁸ Time-resolved fluorescence decay of ABT-888 was ascertained in the absence and presence of various concentrations of ctDNA (Fig. 4). All time-resolved fluorescence data were analyzed by bi-exponential iterative fittings. Fitting parameters are listed in Table 3. Goodness of fit was evaluated from reduced chi-square (χ^2) values, and in general, $\chi^2 < 1.3$ is

acceptable. Mean fluorescence lifetimes (τ_{avg}) were calculated from the decay times (τ_1 , τ_2) and normalized pre-exponential factors (α_1 , α_2) using the following equation:³⁹

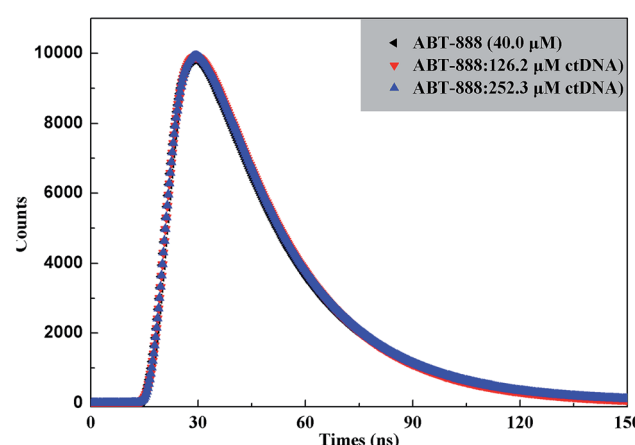


Fig. 4 Time-resolved fluorescence decay of ctDNA–ABT-888 systems in the absence and presence of various concentrations of ctDNA.

Table 3 Fluorescence decay fitting parameters of ctDNA–ABT-888 (lifetimes in ns) in the presence of various concentrations of ctDNA

Sample	τ_1	τ_2	α_1	α_2	τ_{avg}	χ^2
ABT-888 (40.0 μM)	0.5095	2.4677	0.0665	0.9335	2.3375	1.0598
ABT-888 + ctDNA (126.2 μM)	0.5922	2.4799	0.0660	0.9340	2.3553	1.1574
ABT-888 + ctDNA (252.3 μM)	0.5041	2.4838	0.0573	0.9427	2.3703	1.1897

$$\tau_{\text{avg}} = \alpha_1 \tau_1 + \alpha_2 \tau_2 \quad (3)$$

As mentioned in Table 3, $\chi^2 < 1.3$ indicates acceptable quality of exponential fits. ABT-888 exhibits bi-exponential fluorescence decay in buffer solution with the lifetime of 2.3375 ns. With gradually increasing ctDNA concentrations, the lifetime value increases to 2.3703 ns. The result shows that a short lifetime increment of ~ 33 ps appeared in the decay profile. Thus the τ_{avg} , estimated from the lifetime decay measurements, almost does not change in the presence of increasing ctDNA concentrations. The results are practically the same as the steady-state fluorescence data that the quenching is mainly static quenching mechanism. Hence, both steady-state fluorescence and time-resolved fluorescence decay measurements confirmed that static type fluorescence quenching was caused mainly by ground-state complex formation.

3.4. Determination of binding constants and number of binding sites

From variations in intrinsic fluorescence of small molecules in the presence of ctDNA, binding constant (K_b) between small molecules and ctDNA can be possibly calculated.^{19,20} Thus, binding constant (K_b) and a number of binding sites (n) in ABT-888 and ctDNA were determined from the above fluorimetric titration. Parameters were calculated using the following equation:^{40,41}

$$\log(F_0 - F)/F = \log K_b + n \log[Q] \quad (4)$$

where F_0 and F denote fluorescence emission intensity of ABT-888 in the absence and presence of ctDNA; $[Q]$ is ctDNA concentration. Fig. 3C presents the plots of $\log(F_0 - F)/F$ versus $\log[Q]$, and Table 2 summarizes K_b values and n at three various temperatures. As shown in Table 2, values of K_b are 9.17, 4.39, and $2.91 \times 10^4 \text{ M}^{-1}$ at 295, 303, and 310 K respectively, suggesting that binding constants decreased with increasing temperature. K_b was lower than the reported values for some typical intercalators, such as EB ($2.6 \times 10^6 \text{ M}^{-1}$) and acridine orange ($4.0 \times 10^5 \text{ M}^{-1}$),⁴² but higher than *N,N*-bis(3 β -acetoxy-5 α -cholest-6-yl- idene) hydrazine ($4.7 \times 10^3 \text{ M}^{-1}$), which binds to DNA in groove-binding mode.⁴³ Thus, results indicate that DNA-binding mode of ABT-888 is not simple intercalation or groove-binding; such observation is consistent with UV-absorption and ^1H NMR results.

3.5. Determination of thermodynamic parameters

Thermodynamic analysis is normally used to determine the essence of interaction between small molecules and

biomolecules. Interaction forces between small molecules and biomolecules may involve van der Waals interactions, hydrophobic forces, hydrogen bonds, electrostatic interactions, etc.^{25,40} To characterize the binding force involved in ABT-888 and ctDNA, thermodynamic parameters, such as enthalpy change (ΔH), entropy change (ΔS), and free energy change (ΔG) of the interaction, were determined using the following equations:⁴¹

$$\ln K_b = -\Delta H/RT + \Delta S/R \quad (5)$$

$$\Delta G = \Delta H - T\Delta S \quad (6)$$

where R is the gas constant, and temperatures are same as those in quenching mechanism studies. ΔH and ΔS values were obtained from slope and intercept of the linear plot (eqn (5)) based on $\ln K_b$ versus $1/T$, and ΔG values were estimated from eqn (6) at corresponding temperatures (Fig. 3D). ΔH was $-14.00 \text{ kcal mol}^{-1}$, and ΔS was $-24.80 \text{ cal mol}^{-1} \text{ K}^{-1}$ (Table 2), indicating that hydrogen bond and van der Waals play major roles in ABT-888-to-ctDNA binding. Per the rules summarized by Ross and Subramanian,⁴⁴ negative values of ΔH and ΔG revealed that the interaction was spontaneous and predominantly enthalpically-driven reaction.

3.6. Competitive displacement assay

To corroborate the interacting mode, competition binding experiments were carried out using two well-known DNA binding dyes, EB and Hoechst 33258. Dye-competition-binding study can confirm whether this dye and an unknown drug compete for the same binding locus on the DNA. EB is a classic intercalating dye that can produce bright, intense fluorescence when it binds to DNA.^{45,46} Changes observed in the fluorescence intensities of the ctDNA–EB complex are associated with quenching of its luminescence. Such events may indicate displacement of intercalating EB and efficient interaction of the studied compound with the DNA helix. Hoechst 33258 binds strongly to the minor groove of double-stranded B-DNA with specificity for A-T rich sequence.⁴⁷ Addition of excess unknown drug in solution containing Hoechst 33258 and DNA reduces fluorescence intensity of Hoechst 33258–DNA complex when the drug also binds to the groove region.

In this study, EB was first selected as fluorescence probe to explore the binding mode between ABT-888 and ctDNA, and molar ratio ([ctDNA]/[EB]) of 14.5 was used as basis for EB fluorescence intensity, which did not further increase upon sequential addition of DNA to EB solution. Fig. 5A displays the effect of increasing concentration of ABT-888 on fluorescence



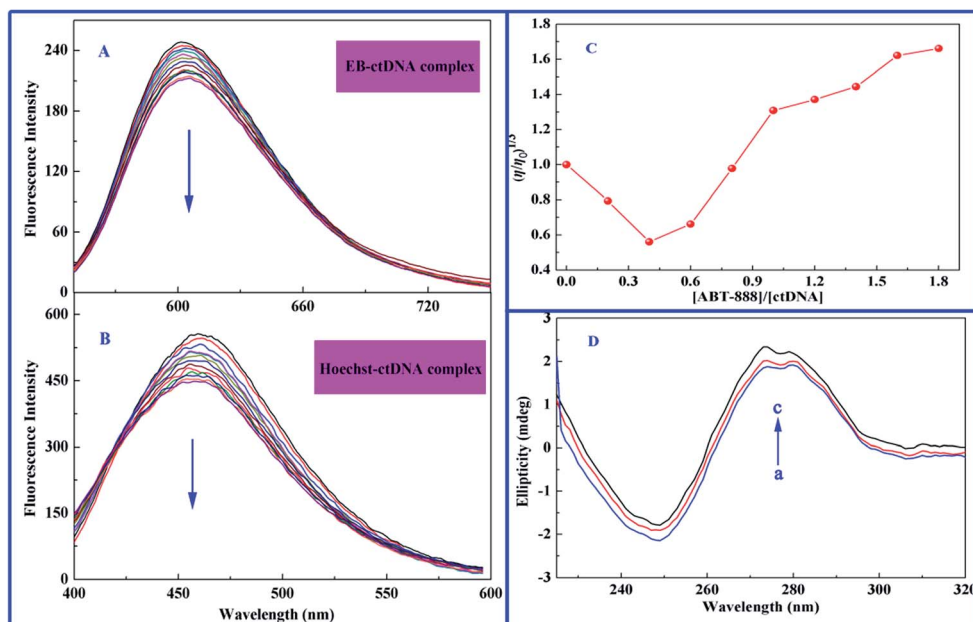


Fig. 5 Competition displacement assays between ABT-888 and various dyes. (A) Fluorescence displacement spectra of EB-ctDNA complex with increasing concentrations of ABT-888 excited at 481 nm. (B) Fluorescence displacement spectra of Hoechst-ctDNA complex with increasing concentrations of ABT-888 excited at 343 nm. (C) Effect of increasing amounts of ABT-888 on relative viscosity of ctDNA at pH 7.4 and 298 K. $C[ctDNA] = 7.7 \times 10^{-5}$ M. (D) CD spectra of ctDNA in the presence of increasing amounts of ABT-888 at pH 7.4 and 298 K. $C[ctDNA] = 3.5 \times 10^{-4}$ M, $C[ABT-888] = 0$; 4.0 and 8.0×10^{-5} M correspond to the curves from a to c, respectively.

spectra of EB-ctDNA complex. Apparently, fluorescence intensity of EB-ctDNA complex at 601 nm remarkably decreased with increasing concentrations of ABT-888 until measurements reached 1.8×10^{-4} M. Such result suggests that ABT-888 could interact with DNA through a similar intercalative mode and was able to replace the intercalation-bound EB dye from double-stranded DNA. Under the same molar ratio of dye-ctDNA (14.5), Hoechst 33258 competition displacement assay was also conducted. Upon subsequent addition of ABT-888, maximum emission wavelength (464 nm) significantly decreased in the Hoechst-ctDNA system (Fig. 5B), suggesting that some Hoechst 33258 molecules were released from Hoechst 33258-ctDNA after an exchange with ABT-888. Results indicate that groove binding is also an interaction mode between ABT-888 and ctDNA. Thus, ABT-888 is not a simple intercalator or groove binder. Instead, ABT-888 and ctDNA bind with each other through a mixed type or partial intercalated mode of DNA binding.

3.7. Viscosity measurements

Viscosity measurement is a hydrodynamic method of determining the binding mode of drugs with DNA.⁴⁸ A classical intercalation binding requires separation of adjacent base pairs to accommodate binding drug molecules into the double helix of DNA, leading to an increased contour length of DNA and DNA viscosity.³² By contrast, a drug molecule that interacts with DNA by groove binding or electrostatic interaction does not change the relative viscosity of DNA, because molecules do not alter the axial length of DNA upon binding.^{31,32} In addition, a partial or non-classical intercalation of drug molecules could bend or

kink DNA helix, resulting in decreased effective length of DNA molecule with concomitant decrease in its viscosity. Fig. 5C shows the relative viscosity $(\eta/\eta_0)^{1/3}$ versus ABT-888-concentration-to-ctDNA mole ratios ($r_i = [ABT-888]/[ctDNA]$). At lower r_i values, relative viscosity showed a decreasing trend, which then increased at higher mole ratios up to the initial viscosity. We then conclude that at low amounts of ABT-888, drugs bind to ctDNA by partial intercalation mode, whereas at higher drug concentrations, intercalation of ABT-888 into the adjacent base pairs of ctDNA increases the effective length of DNA.

3.8. CD study

CD titration is one of the most direct means of examining the binding modes of drug molecules to DNA. Within visible range, a majority of drug molecules do not show any ellipticity in CD spectra. However, right-hand B form of DNA produced a characteristic spectrum with a positive band at 275 nm because of base-stacking and a negative band at 245 nm because of polynucleotide helicity.⁴⁹ Simple groove-binding and electrostatic interaction of drug molecules with DNA show little or no perturbations on base stacking and helicity bands, whereas intercalation changes the intensities of both bands, stabilizing the right-handed B conformation of DNA.⁵⁰ As shown in Fig. 5D, upon addition of ABT-888 to ctDNA, characteristic CD spectrum of ctDNA undergoes changes in both bands. Moreover, ABT-888 increase moderately intensities of the positive band at 275 nm and decrease moderately that of the negative band at 245 nm. However, interestingly, ABT-888 did not affect the peak shapes of ctDNA. Observation of increased CD signals at 275 nm is an

important evidence of interaction of ABT-888 with the base pairs, resulting in the conformational changes in ctDNA.⁵⁰ Possibly, different changes in both bands resulted from binding of ABT-888 with ctDNA *via* mixed binding or partial intercalation mode; such result is consistent with conclusions of above competition experiments.

3.9. FT-IR studies

FT-IR spectroscopy has emerged as an efficient tool for characterization of drug-biomolecules interaction and for identification of biomolecule conformations.¹⁷ Earlier experiments demonstrated that infrared spectra of DNA consist primarily of three principal regions.⁵¹ The first region (~ 1750 – 1600 cm^{-1}) contains absorption bands assigned to ring stretching vibrations of nitrogenous bases ($\text{C}=\text{O}$, $\text{C}=\text{N}$, $\text{C}=\text{C}$) and bending vibrations of exocyclic NH_2 in the DNA duplex.⁵² The second region (~ 1600 – 1500 cm^{-1}) results primarily from purine and pyrimidine ring vibrations. The last region (~ 950 – 1250 cm^{-1}) is mostly due to stretching vibrations of symmetric and asymmetric PO_2 groups in the phosphodiester-deoxyribose backbone.⁵³ Specifically, vibrational bands of free DNA at 1710, 1663, 1608, and 1491 cm^{-1} were assigned to guanine, thymine, adenine, and cytosine nitrogenous base, respectively.^{17,53}

In this study, diverse IR spectra were obtained by addition of different amounts of ABT-888 to ctDNA solution, as evident in Fig. 6A–C. A discernible shift in G band from 1712 to 1718 and 1719 cm^{-1} were observed with increasing ABT-888 concentration, suggesting that ABT-888 interacted with the G base of ctDNA. Similarly, major shifting of the T band from 1665 to 1655 and 1649 cm^{-1} was observed, suggesting that ABT-888 probably interacted with O-2 atoms of T base.³⁰ A (1612 cm^{-1}) and C bands (1494 cm^{-1}) also showed obvious shifts, which

correspond respectively to NH_2 scissoring and N–H bending changes in purine and pyrimidine ring.²⁹ Major shifts in IR bands specify ABT-888 intercalation between A/T and G/C base pairs. The band at 983 cm^{-1} also shifted to 1006 and 1013 cm^{-1} , and a new absorption band appeared at 950 cm^{-1} , indicating that phosphate backbone was weak influenced by ABT-888 interaction with ctDNA. In view of free ctDNA and its complexes spectra, these changes are clearer in the first derivative spectrum (Fig. 6D and E). These feature changes imply that ABT-888 induces local conformational changes in ctDNA to accommodate molecules between base pairs. Result was consistent with that suggested by above CD spectra.

3.10. Molecular docking analysis

Molecular docking is an accepted method in search of therapeutic agents designed to interact with a specific biomolecule. Especially in studies of binding of small molecules to DNA, docking underwent substantial development in the last few years. The method allows for virtual screening on large libraries of compounds, ranking of results, and proposing structural hypotheses on how small molecules bind to DNA.^{19,20,54} Molecular docking calculations were performed using Auto-Dock strategy to search the entire DNA surface for binding locus while simultaneously optimizing conformation of base pairs. Geometry-optimized ABT-888 molecule was successfully docked onto DNA segments. A total of 11 multimember conformational clusters were gathered from 200 docking runs, in which case the highest populated cluster contained almost half of the obtained conformations and was lowest on energy scale. Table 4 displays binding free energies [kcal mol^{-1}] and remarks for the top five conformations in this highest cluster. Data analysis showed that interaction is basically non-

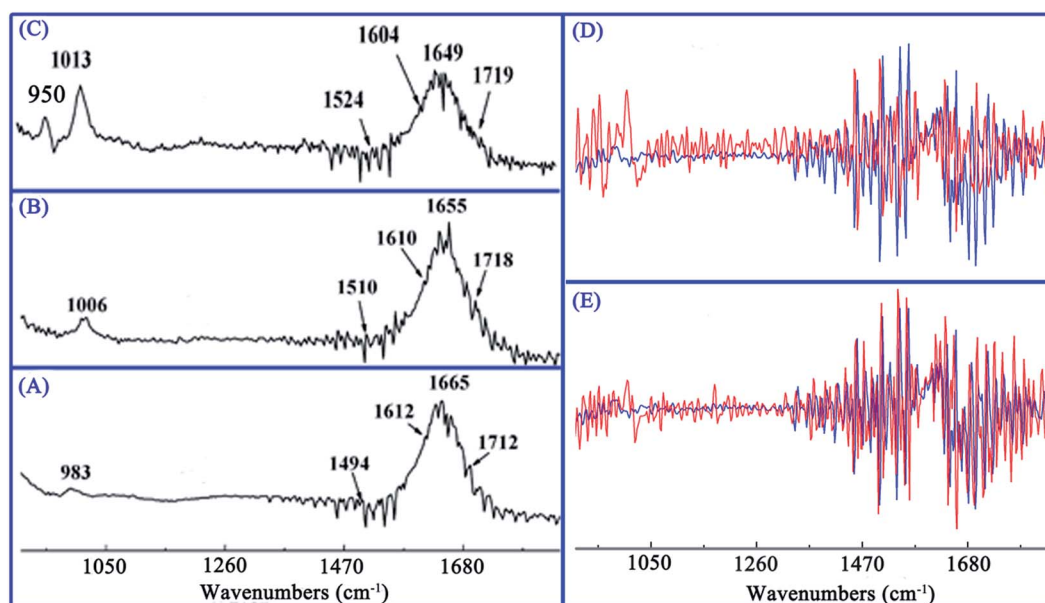


Fig. 6 Effects of DNA interaction with ABT-888. FT-IR spectra of free ctDNA (A) and its complexes with ABT-888 at different molar ratios ($r = [\text{ctDNA}]/[\text{ABT-888}]$) of 7/1 (B), 4/1 (C), and $[\text{ctDNA}] = 4.4 \times 10^{-4}$ M. Overlaid view of first derivative spectra of ctDNA and complexes at molar ratios of 7/1 (D), 4/1 (E), ctDNA (red), and complex (blue).



Table 4 Docking summary of ABT-888 with DNA in the highest populated cluster by AutoDock program generating different molecule conformers (top 5)

Sub rank	Binding energy (kcal mol ⁻¹)	Electrostatic energy (kcal mol ⁻¹)	van der Waals	Inhibition constant (nmol)	Number of H-bonds	Cluster rmsd	Reference rmsd
1	-9.38	-1.39	-8.58	132.71	3	0	25.89
2	-9.38	-1.36	-8.62	132.77	2	1.18	25.54
3	-9.38	-1.39	-8.59	133.04	2	0.04	25.91
4	-9.38	-1.39	-8.59	133.15	2	0.05	25.90
5	-9.38	-1.40	-8.58	133.16	2	0.04	25.90

covalent interaction, that is, van der Waals and electrostatic interactions exist between ABT-888 and DNA. Interactions may be either through phosphate backbone atoms of DNA or *via* the DNA bases.

Optimum binding conformation between ABT-888 and DNA with the lowest binding energy of -9.38 kcal mol⁻¹ was selected for discussion and shown in Fig. 7. As shown in Fig. 7, it was obvious that the benzimidazolyl moiety of ABT-888 bound into the base pairs of DNA by parallel way, while the pyrrolidyl moiety was perpendicular to the base pairs. And the hydrogen atoms (a, b, c) of the benzimidazolyl moiety located inside, which shows that the benzimidazolyl moiety is closer to the inside of DNA duplex. The result was coincided with the deduction of ¹H NMR study. Fig. 7 also shows that ABT-888 binds mainly to minor grooves in A-T base pairs region of DNA. ABT-888 bends base pairs by intermolecular hydrogen bonds, namely, the ABT-888 three N-H groups, which forms three hydrogen bonds with DT7, DA17 and DT20, respectively. The O-2 oxygen of DT7 is at a distance of 2.13 Å from the NH₂ amide H atom of ABT-888. Similarly, O-3 oxygen of DT17 is 1.95 Å from the NH imidazole H atom of ABT-888, and O-3 oxygen of DT20 with the NH pyrrole H atom is 1.96 Å. Based on above analysis, groove binding interaction may be play a major role in binding of ABT-888 to DNA, whereas other binding types, including classic intercalation and partial intercalation, cannot be ruled out. Thus, molecular docking studies shed light on binding modes between ABT-888 and DNA and complement experimental observations.

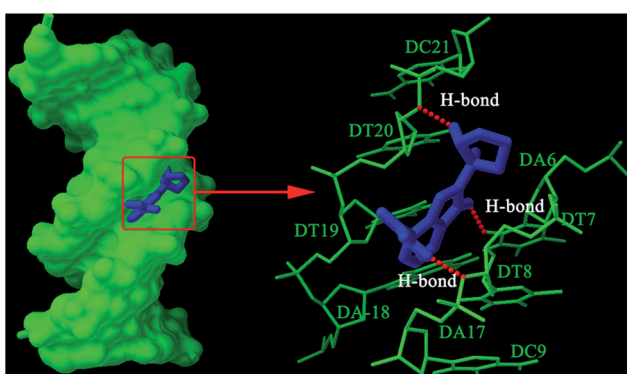


Fig. 7 Molecule docked poses of ABT-888 with DNA (left) and hydrogen bonds (red dotted lines) between ABT-888 and bases (right).

4. Conclusions

The present study attempted to analyze binding properties of ABT-888 to ctDNA. From these multispectroscopic experiments and molecular docking studies, results showed that mixed-ligand type binding occurs between ABT-888 and DNA with a binding constant of 9.17×10^4 M⁻¹ at 295 K. Calculated thermodynamic parameters suggested that binding of ABT-888 to ctDNA is driven mainly by hydrogen bonds and van der Waals force. Competition fluorescence displacement experiments were studied using EB and Hoechst 33258 as probes for intercalation and minor groove binding modes, respectively, and these experiments indicated that ABT-888 can compete simultaneously for binding sites on ctDNA with the two probes. In viscometric titration, relative viscosity of ctDNA decreased and increased steadily at low and high ABT-888 concentration regions respectively, suggesting transition from predominantly partial intercalation to predominantly classic intercalation. Changes in CD and FT-IR spectra indicated that ABT-888 induces conformational transformation of ctDNA. Theoretical calculations applying AutoDock have been carried out, and calculational results can reasonably complement experimental findings when ABT-888 interacted with DNA by groove binding mode.

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References

- 1 C. Jones and E. R. Plummer, *Br. J. Radiol.*, 2008, **81**, 2–5.
- 2 Y. Drew and R. Plummer, *Drug Resist. Updates*, 2009, **12**, 153–156.
- 3 L. Gatti and F. Zunino, in *Chemosensitivity: Volume II: In vivo Models, Imaging, and Molecular Regulators*, 2005, pp. 127–148.
- 4 P. Borst, S. Rottenberg and J. Jonkers, *Cell Cycle*, 2008, **7**, 1353–1359.
- 5 J. R. Marchand, A. Carotti, D. Passeri, P. Filippioni, P. Liscio, E. Camaioni, R. Pellicciari, A. Gioiello and A. Macchiarulo,



Biochim. Biophys. Acta, Proteins Proteomics, 2014, **1844**, 1765–1772.

6 B. Basu, S. K. Sandhu and J. S. de Bono, *Drugs*, 2012, **72**, 1579–1590.

7 M. Rouleau, A. Patel, M. J. Hendzel, S. H. Kaufmann and G. G. Poirier, *Nat. Rev. Cancer*, 2010, **10**, 293–301.

8 L. M. Wagner, *OncoTargets Ther.*, 2015, **8**, 1931–1939.

9 H. Farmer, N. McCabe, C. J. Lord, A. N. Tutt, D. A. Johnson, T. B. Richardson, M. Santarosa, K. J. Dillon, I. Hickson and C. Knights, *Nature*, 2005, **434**, 917–921.

10 L. Virág and C. Szabó, *Pharmacol. Rev.*, 2002, **54**, 375–429.

11 S. Bogliolo, C. Cassani, M. Dominoni, V. Musacchi, P. L. Venturini, A. Spinillo, S. Ferrero and B. Gardella, *Expert Opin. Invest. Drugs*, 2016, **25**, 367–374.

12 T. Penning, G. Zhu, V. Gandhi, J. Gong, X. Liu, Y. Shi, V. Klinghofer, E. Johnson, C. Donawho and D. Frost, *J. Med. Chem.*, 2009, **52**, 514–523.

13 X. Li, J. Delzer, R. Voorman, S. M. de Moraes and Y. Lao, *Drug Metab. Dispos.*, 2011, **39**, 1161–1169.

14 J. P. Palma, Y. C. Wang, L. E. Rodriguez, D. Montgomery, P. A. Ellis, G. Bukofzer, A. Niquette, X. Liu, Y. Shi and L. Lasko, *Clin. Cancer Res.*, 2009, **15**, 7277–7290.

15 D. Sarkar, P. Das, S. Basak and N. Chattopadhyay, *J. Phys. Chem. B*, 2008, **112**, 9243–9249.

16 D. Sahoo, P. Bhattacharya and S. Chakravorti, *J. Phys. Chem. B*, 2010, **114**, 2044–2050.

17 P. Ghosh, G. P. Devi, R. Priya, A. Amrita, A. Sivaramakrishna, S. Babu and R. Siva, *Appl. Biochem. Biotechnol.*, 2013, **170**, 1127–1137.

18 D. K. Jangir, S. Kundu and R. Mehrotra, *PLoS One*, 2013, **8**, 69933.

19 S. U. Rehman, T. Sarwar, H. M. Ishqi, M. A. Husain, Z. Hasan and M. Tabish, *Arch. Biochem. Biophys.*, 2015, **566**, 7–14.

20 I. Ahmad, A. Ahmad and M. Ahmad, *Phys. Chem. Chem. Phys.*, 2016, **18**, 6476–6485.

21 J. Liu, Y. Yue, J. Wang, X. Yan, R. Liu, Y. Sun and X. Li, *Spectrochim. Acta, Part A*, 2015, **145**, 473–481.

22 D. Wu, J. Yan, J. Wang, Q. Wang and H. Li, *Food Chem.*, 2015, **170**, 423–429.

23 D. Pastré, O. Piétrement, A. Zozime and E. L. Cam, *Biopolymers*, 2005, **77**, 53–62.

24 Y. Yue, Y. Sun, X. Yan, J. Liu, S. Zhao and J. Zhang, *Chemosphere*, 2016, **161**, 475–481.

25 Y. Yue, J. Liu, R. Liu, Y. Sun, X. Li and J. Fan, *Food Chem. Toxicol.*, 2014, **71**, 244–253.

26 S. Yousuf, D. Radhika, I. V. M. V. Enoch and M. Easwaran, *Spectrochim. Acta, Part A*, 2012, **98**, 405–412.

27 Y. Sameena and I. V. Enoch, *J. Lumin.*, 2013, **138**, 105–116.

28 S. Yousuf, R. Alex, P. M. Selvakumar, I. V. Enoch, P. S. Subramanian and Y. Sun, *ChemistryOpen*, 2015, **4**, 497–508.

29 L. Han, Y. Zhou, X. Huang, M. Xiao, L. Zhou, J. Zhou, A. Wang and J. Shen, *Spectrochim. Acta, Part A*, 2014, **123**, 497–502.

30 L. Wang, G. Zhang, J. Pan, C. Xiong and D. Gong, *J. Photochem. Photobiol. B*, 2014, **141**, 253–261.

31 S. U. Rehman, T. Sarwar, M. A. Husain, H. M. Ishqi and M. Tabish, *Arch. Biochem. Biophys.*, 2015, **576**, 49–60.

32 Y. Zhang, G. Zhang, F. Peng, Y. Ma and J. Zhou, *Spectrochim. Acta, Part A*, 2012, **96**, 1012–1019.

33 A. Anantharaman, R. R. Priya, H. Hemachandran, A. Sivaramakrishna, S. Babu and R. Siva, *Spectrochim. Acta, Part A*, 2015, **144**, 163–169.

34 W. Zhong, J. S. Yu, Y. Liang, K. Fan and L. Lai, *Spectrochim. Acta, Part A*, 2004, **60**, 2985–2992.

35 H. Guo, C. Cai, H. Gong and X. Chen, *Spectrochim. Acta, Part A*, 2011, **79**, 92–96.

36 X. Li and S. Wang, *New J. Chem.*, 2015, **39**, 386–395.

37 Y. Yue, Y. Sun, Q. Dong, R. Liu, X. Yan, Y. Zhang and J. Liu, *Luminescence*, 2016, **31**, 671–681.

38 P. Mandal, M. Bardhan and T. Ganguly, *J. Photochem. Photobiol. B*, 2010, **99**, 78–86.

39 S. Ghosh, P. Kundu and N. Chattopadhyay, *J. Photochem. Photobiol. B*, 2016, **154**, 118–125.

40 Y. Yue, Q. Dong, Y. Zhang, X. Li, X. Yan, Y. Sun and J. Liu, *Spectrochim. Acta, Part A*, 2016, **153**, 688–703.

41 Y. Yue, J. Liu, R. Liu, Q. Dong and J. Fan, *Spectrochim. Acta, Part A*, 2014, **124**, 46–51.

42 Y. Cao and X.-W. He, *Spectrochim. Acta, Part A*, 1998, **54**, 883–892.

43 R. Vijayabharathi, P. Sathyadevi, P. Krishnamoorthy, D. Senthilraja, P. Brunthadevi, S. Sathyabama and V. B. Priyadarisini, *Spectrochim. Acta, Part A*, 2012, **89**, 294–300.

44 P. D. Ross and S. Subramanian, *Biochemistry*, 1981, **20**, 3096–3102.

45 Y. Song, J. Kang, J. Zhou, Z. Wang, X. Lu, L. Wang and J. Gao, *Spectrochim. Acta, Part A*, 2000, **56**, 2491–2497.

46 M. B. Lyles and I. L. Cameron, *Biophys. Chem.*, 2002, **96**, 53–76.

47 M. M. Ramana, R. Betkar, A. Nimkar, P. Ranade, B. Mundhe and S. Pardeshi, *Spectrochim. Acta, Part A*, 2016, **152**, 165–171.

48 S. U. Rehman, T. Sarwar, M. A. Husain, H. M. Ishqi and M. Tabish, *Arch. Biochem. Biophys.*, 2015, **576**, 49–60.

49 M. Tao, G. Zhang, J. Pan and C. Xiong, *Spectrochim. Acta, Part A*, 2016, **155**, 28–37.

50 J. Kang, X. Wang and S. Dong, *Toxicol. Mech. Methods*, 2006, **16**, 515–523.

51 S. H. Brewer, S. J. Anthireya, S. E. Lappi, D. L. Drapcho and S. Franzen, *Langmuir*, 2002, **18**, 4460–4464.

52 C. Le-Tien, R. Lafortune, F. Shareck and M. Lacroix, *Talanta*, 2007, **71**, 1969–1975.

53 B. Zhousun, J. Liquier, E. Taillandier, J. Sun, T. Garestier, C. Hélène and S. M. Gryaznov, *Nucleic Acids Res.*, 1997, **25**, 1782–1787.

54 A. Bykowska, R. Starosta, J. Jezierska and M. Jeżowskabojczuk, *RSC Adv.*, 2015, **5**, 80804–80815.