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2D NMR Study of the DNA Duplex d(CTCTC*A*ACTTCC)•d(GGAAGTTGAGAG) Cross-Linked by the Antitumor-Active Dirhodium(II,II) Unit at the Cytosine–Adenine Step[†]

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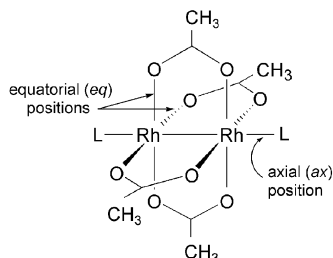
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ABSTRACT: The 2D NMR analysis in solution of the DNA duplex d(CTCTC*A*ACTTCC)•d(GGAAGTTGAGAG) binding to the dirhodium unit *cis*-[Rh₂(μ-O₂CCH₃)₂(η¹-O₂CCH₃)]⁺ showed that an unprecedented *intrastrand* adduct, dsII, is formed with the dirhodium unit cross-linking in the major groove residues C5 and A6 (indicated with asterisks), also corroborated by enzyme digestion studies. Formation of the dirhodium complex dsII destabilizes significantly the duplex as indicated by the substantial decrease in its melting temperature (Δ*T*_m = −22.9 °C). The reduced thermal stability of dsII is attributed to the decreased stacking of the bases and the complete disruption and/or weakening of the hydrogen bonds within the base pairs in the immediate vicinity of the metalation site (C5•G20 and A6•T19), but the effects due to the metal binding are more severe for the base pairs in the 5′ direction to the lesion site. The NMR spectroscopic data indicate that Watson–Crick hydrogen bonding is completely disrupted for the C5•G20 site and considerably weakened for A6•T19. In dsII, the bases C5 and A6 bind to *eq* positions of the dirhodium unit *cis*-[Rh₂(μ-O₂CCH₃)₂(η¹-O₂CCH₃)]⁺, which retains one monodentate and two bridging acetate groups, presumably due to steric reasons. Binding of A6 takes place via N7, whereas binding of the C5 base takes place via the exocyclic N4 site, resulting in the *anti*-cytosine rotamer with respect to site N3 in its metal-stabilized rare iminooxo form.

In spite of the phenomenal success of cisplatin and related platinum compounds as anticancer drugs (1–3), a need has risen for new types of antitumor agents due to the limitations of platinum drugs, i.e., natural and acquired resistance of tumor cells, in addition to the numerous deleterious side effects (4–7). In this vein, dirhodium(II) tetracarboxylate derivatives (8) emerged as one of the most promising classes of antitumor-active transition-metal compounds, with Rh₂(O₂CCH₃)₄ being the first member in the series. Dirhodium tetraacetate exhibits a “paddlewheel” structure with four equatorial (*eq*) acetate groups bridging the dirhodium unit in a symmetrical fashion and two axial (*ax*) sites available to donor ligands (Chart 1) (8).

Chart 1: Structure of Dirhodium Tetraacetate



Pioneering studies that emanated in the 1970s showed that dirhodium carboxylate compounds Rh₂(O₂CR)₄ (R = Me, Et, Pr) exhibit significant *in vivo* antitumor activity against L1210 tumors (9, 10), Ehrlich ascites (11–13), and sarcoma 180 and P388 tumor lines (14). Although the exact mechanism of action of dirhodium compounds has not yet been elucidated, the most probable targets are DNA (15), RNA (16, 17), and enzymes (18) involved in DNA and RNA synthesis. Studies of the binding affinity of Rh₂(O₂CCH₃)₄ to several biomolecules by employing ¹⁴C-labeled Rh₂(O₂CCH₃)₄ indicated that Rh₂(O₂CCH₃)₄ is only slightly reactive toward native calf thymus DNA and polyguanylic (poly-G) and polycytidylic (poly-C) acids, but binds fairly well to denatured DNA and polyadenylic acid (poly-A) (11, 12). The interactions of Rh₂(O₂CCH₃)₄ with adenine nucleos(t)ides (11, 12, 19, 20) are established by axial binding of the adenine bases (via N7 and N1 if available (19)) and formation of hydrogen bonds between the exocyclic adenine NH₂ groups and the oxygen atoms of carboxylate ligands (for binding via N7), as evidenced by the reported crystal structures of Rh₂(O₂CCH₃)₄(1-MeAdo)₂¹ (21) and [Rh₂(O₂CCH₃)₂(NHCOCF₃)₂(9-methyladeninium)₂](NO₃)₂ (22). In

¹ Abbreviations: Ado, adenosine; CD, circular dichroism; COSY, double-quantum-filtered correlation spectroscopy; dap, 1,12-diazaperylene; dsI, native d(CTCTCAACTTCC)•d(GGAAGTTGAGAG); dsII, metalated d(CTCTC*A*ACTTCC)•d(GGAAGTTGAGAG) (dsD); DTolF, anion of *N,N'*-p-tolylformamidate; ESI-MS, electrospray ionization mass spectrometry; HPLC, high-performance liquid chromatography; HSQC, heteronuclear single-quantum correlation; MALDI-MS, matrix-assisted laser desorption/ionization mass spectrometry; NOESY, 2D nuclear Overhauser spectroscopy; ssI, native d(CTCTCAACTTCC); ssII, native d(GGAAGTTGAGAG); ssIII, metalated d(CTCTC*A*ACTTCC); *T*_m, melting temperature of a DNA duplex.

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the presence of DTolF groups, however, 9-ethyladenine (23, 24) and d(ApA) (25) bind to the dirhodium unit $[\text{Rh}_2(\text{DTolF})_2(\text{CH}_3\text{CN})_6]^{2+}$ in a bridging fashion through positions N7 and N6 of the adenine bases. Furthermore, in the cytosine and 1-methylcytosine complexes with $\text{Rh}_2(\text{HNOCCF}_3)_4$, the pyrimidine ring binds axially to the dirhodium core via site(s) N3 or both N3 and O2, and the structure is further stabilized by intramolecular hydrogen bonds between the nucleobase exocyclic groups and the dirhodium trifluoroacetamidate moieties (26).

In spite of earlier claims (11, 12), recent studies revealed that guanine nucleobases (27–29), guanine dinucleotides (30–32), and double-stranded DNA (33) also bind to dirhodium units, although the reactivity of $\text{Rh}_2(\text{O}_2\text{CCH}_3)_4$ is lower as compared to other dirhodium compounds (33, 34). In two studies, the reactions of dirhodium(II) complexes with single-stranded oligonucleotides were monitored by mass spectrometry and enzymatic digestion studies (35, 36). To gain further insight into the interactions of dirhodium complexes with DNA, we pursued the goal of structurally characterizing in solution a dirhodium–oligonucleotide duplex by 2D NMR spectroscopy. Herein, we report the structural characterization of the duplex $\text{d}(\text{CTCTC}^*\text{A}^*\text{ACTTCC})\cdot\text{d}(\text{GGAAGTTGAGAG})$ with the dirhodium cross-link at the C^*A^* site (dsII; C^* and A^* indicate the metalated sites). The interactions of the same duplex with the dirhodium complex *cis*- $[\text{Rh}_2(\mu\text{-O}_2\text{CCH}_3)_2(\eta^1\text{-O}_2\text{CCH}_3)(\text{dap})(\text{CH}_3\text{OH})](\text{O}_2\text{CCH}_3)$ containing the intercalating group dap have been the subject of another detailed NMR spectroscopic investigation (37).

EXPERIMENTAL PROCEDURES

Materials. The compound $\text{Rh}_2(\text{O}_2\text{CCH}_3)_4$ was prepared by standard procedures (8). The DNA oligonucleotide $\text{d}(\text{CTCTCAACTTCC})$ (ssI) and its complementary strand $\text{d}(\text{GGAAGTTGAGAG})$ (ssII) were purchased from Integrated DNA Technologies, Inc. (Coralville, IA) as crude materials. The enzymes Phosphodiesterase I (SVP) and II (BSP) were obtained from Sigma.

Oligonucleotide Purification. The crude oligonucleotide materials for ssI and ssII were purified by anion exchange HPLC chromatography. A Dionex DNAPac PA-100 (22 × 250 mm) anion exchange column was used on a Beckman Coulter System Gold instrument equipped with a 168 diode array detector for large-scale purification of oligonucleotides using the following elution conditions: 25 mM ammonium acetate in a $\text{CH}_3\text{CN}/\text{H}_2\text{O}$ (10:90) solution (eluent A) and eluent A + 1.0 M NaCl (eluent B) starting with 28% eluent B, which was increased to 33% eluent B for 19 min to 60% eluent B for the next 18 min to 100% eluent B for the final 5 min with a 10 mL/min flow rate. The fractions collected from the HPLC purification were reduced in volume using a centrifugal evaporator (Jouan, Winchester, VA) and desalted on a Sephadex G-25 size exclusion column. The collected fractions from the size exclusion column were filtered through a 0.22 μm membrane (Fisher Scientific), and their concentrations were determined by UV spectroscopy (Shimadzu UV 1601PC spectrophotometer) at 260 nm prior to the samples being lyophilized.

Preparation and Isolation of the Metalated Duplex dsII. A solution of $\text{Rh}_2(\text{O}_2\text{CCH}_3)_4$ in H_2O was added to an

oligonucleotide solution of $\text{d}(\text{CTCTCAACTTCC})$ (ssI) in H_2O in a 5:1 ratio (5 mM final concentration of oligonucleotide), the metal–DNA mixture was incubated at 37 °C for ~4–5 h, and the progress of the reaction was monitored over time by anion exchange column HPLC (Figure S1a in the Supporting Information). The major adduct of the reaction (eluting at ~22 min) was separated from the reaction mixture by anion exchange HPLC chromatography using a Dionex DNAPac PA-100 (9 × 250 mm) column. The same eluents used for the purification of the free oligonucleotides were used for the purification of the metalated adduct, with a gradient of 20–35% eluent B over a 30 min period and 35–100% eluent B over the next 20 min with a reduced flow rate of 3 mL/min. The collected fractions were desalted on a Sephadex G-25 size exclusion column and lyophilized. The dried and desalted major adduct from the anion exchange HPLC column (Figure S1a) was reinjected into a MetaChem Inertsil ODS-3 (10 × 250 mm) reversed-phase HPLC column and was further separated into several peaks, A–E (Figure S1c), under the following elution conditions: 25 mM ammonium acetate in H_2O (eluent A)/25 mM ammonium acetate in MeCN (eluent B) with a gradient of 5–10% eluent B over 10 min, 10–20% eluent B over the next 20 min, and finally 20–100% eluent B in the remaining 20 min. Samples were collected for the metalated species A–E (Figure S1c), and each metalated adduct was added to the complementary strand (ssII). The molecular mass of each single-stranded adduct A–E collected from the reversed-phase column is $[\text{M} + \text{Rh}_2(\text{O}_2\text{CCH}_3)_3]$, $\text{M} = \text{ssI}$, as determined by ESI-MS (Figure S2 in the Supporting Information).

The concentrations of the solutions of the single-stranded metalated species (Figure S1c) were estimated by measuring the absorbance at 260 nm. On the basis of the calculated concentrations, a quantity of the purified complementary strand $\text{d}(\text{GGAAGTTGAGAG})$ (ssII) was added to each metalated strand. The resulting solutions of the double-stranded adducts were injected into the anion exchange column under the same conditions employed for the isolation of the single-stranded adducts except for the gradient, which was 28–31% eluent B over a 6 min period, 31–55% eluent B over the next 10 min, and 55–100% B for the final 5 min (Figure S3 in the Supporting Information). The concentrations of the collected fractions dsA–dsE were measured by UV spectroscopy, and the samples were reduced in volume, desalted, and lyophilized. Typically, the color of the dirhodium samples was pink or light purple depending on the concentration of the adduct. As indicated by detailed NMR experiments, adduct dsD (dsII) exhibits the most prominent changes among the modified double-stranded oligonucleotides studied (dsA–dsE) and thus is the focus of this report.

NMR Experiments. The final sample of dsII was dissolved in a 40 mM KHPO_4 buffer solution (pH 6.9) containing 300 mM NaCl and 0.2 mM DSS, lyophilized twice with D_2O , and redissolved in 99.96% D_2O . For observation of the exchangeable protons, the dsI and dsII samples were dissolved in 90% $\text{H}_2\text{O}/10\%$ D_2O . The final concentration of each sample used for collection of the NMR data was 1 mM (duplex), and the volume was 260 μL in a Shigemi tube (Shigemi, Inc., Allison Park, PA). The 2D NMR data sets were collected at 15 °C. NMR experiments were performed on either a Varian Inova 500 MHz spectrometer equipped with a 5 mm indirect detection probe or a Varian 600 MHz

spectrometer equipped with a triple-axis gradient pentaprobe. All chemical shifts were referenced to internal DSS (sodium 2,2-dimethyl-2-silapentane-5-sulfonate) (38). NOESY experiments in D₂O were performed by using a Varian pulse sequence (noesy.c 14.1) with a mixing time of 280 ms, a 10 ppm sweep width, in the hypercomplex mode, 2150 complex points, 512 *t*₁ blocks, 32–64 scans per *t*₁ block, 3 or 9 s delay times, and water suppression by presaturation during relaxation delay. 2D [¹H–¹H] DQF-COSY (double-quantum-filtered correlation spectroscopy) spectra were typically collected with a 10 ppm sweep width, 1792 data points in the *t*₂ dimension, 500 *t*₁ increments, and 32 scans per *t*₁ increment using the Varian dqcosy.c 14.1 pulse sequence. The chemical shifts of the imino and amino protons were assigned by collecting Watergate-NOESY spectra at 15 °C with a mixing time of 200 ms (39). The assignments of the H2 protons of the adenosine residues and the ¹³C atoms of the aromatic rings were performed by a combination of short-range (one-bond) and long-range [¹H, ¹³C] heteronuclear single-quantum correlation (HSQC) experiments (40). All 2D NMR data were processed using the program nmrPipe (41) with the 90° phase-shifted sine-bell apodization function and zero-filling. The baseline was corrected with first- and second-order polynomials. The program SPARKY (<http://www.cgl.ucsf.edu/home/sparky/>) was used for the peak assignments of dsI and dsII.

Enzyme Digestion Studies. Enzymatic digestion experiments were carried out using Phosphodiesterase I from *Crotallus adamanteus* venom (SVP) (3' ↔ 5' DNA exonuclease) and Phosphodiesterase II from bovine spleen (BSP) (5' → 3' DNA exonuclease). Phosphodiesterase I (SVP) digestions were performed at room temperature with no additional buffer using 5 times less enzyme than the incubations with Phosphodiesterase II (BSP) digestions, which were performed for 2 h at 37 °C with no additional buffer.

Mass Spectroscopy. Electrospray mass spectra of the isolated dirhodium adducts were acquired on an MDS Sciex API QStar Pulsar (Toronto, Ontario, Canada) fitted with a Protana (Odense, Denmark) nanoelectrospray source. All spectra were acquired in the negative ion mode in 50% H₂O and 50% 2-propanol with 5 mM ammonium acetate. The spray voltage was between –1050 and –1100 V. The nozzle-skimmer potential was set to –5 V to minimize fragmentation in that region.

Thermal Denaturation Experiments. The thermal melting profiles for dsI (unmetalated duplex) and dsII (metalated duplex) were obtained by monitoring the solutions at 260 nm on a CARY 300 Bio UV–vis spectrometer (Varian) equipped with a six-cell sample changer and temperature control. Capped 10 mm path length cuvettes were used, and one of the cuvettes was filled with buffer as a reference. The same salt and buffer conditions used for preparation of the NMR samples were used for the thermal denaturation experiments. Stock solutions of dsI and dsII were prepared at a concentration of 2 μM, and 900 μL of the stock solution was used for each measurement. The temperature was increased from 5 to 90 °C at a rate of 0.3 °C/min and then decreased to 5 °C for 5 min; the first derivative of each melting profile was used to calculate the melting temperatures *T*_m for dsI and dsII.

Circular Dichroism Spectropolarimetry. Circular dichroism (CD) spectropolarimetry was employed to investigate the average conformation of dsII. Data were collected on an AVIV 62DS spectropolarimeter. In a typical experiment, a 2 μM solution of duplex dissolved in the NMR buffer solution was added to a 10 mm cuvette, and the data were recorded between 200 and 350 nm with a bandwidth of 1 nm and an averaging time of 5 s.

Molecular Modeling. The sander program in Amber (42) (version 9) was used to minimize the manually docked dirhodium–DNA complex with the generalized Born solvation model and a nonbonding cutoff of 60.0 Å (Supporting Information). Unknown parameters and charges were developed on the basis of the X-ray crystal structure of dirhodium tetraacetate (43) and the TPSS (44) optimized complex *cis*-[Rh₂(μ-O₂CCH₃)₂(η¹-O₂CCH₃)(A-N7)(C-N4)(H₂O)]⁺ (Supporting Information). TPSS calculations with density-fitting basis sets (45) were performed using Gaussian 03 (46) with the 6-31G(d') basis set (47) for C, N, O, and H and the all-electron DGDZVP basis set (48) for Rh. Force constants were set to reasonable values on the basis of similar terms in the ff03 force field. The new parameters were validated by comparing the modified ff03-optimized geometry of dirhodium tetraacetate with the crystal structure (43) and the modified ff03-optimized geometry of the dirhodium–DNA adduct with the TPSS-optimized *cis*-[Rh₂(μ-O₂CCH₃)₂(η¹-O₂CCH₃)(A-N7)(C-N4)(H₂O)]⁺ structure.

RESULTS

Determination of *T*_m. UV melting profiles were carried out to determine the distortion induced by the dirhodium binding in dsII as compared to the native DNA duplex (dsI). The *T*_m values of dsI and dsII, under identical experimental conditions, are 47.1 ± 1 and 24.2 ± 1 °C, respectively (Figure S4 in the Supporting Information). The decrease in the *T*_m value (Δ*T*_m = –22.9 °C) for dsII indicates that the double-stranded oligonucleotide is considerably destabilized due to the metal binding and corroborates the presence of a covalent *intrastrand* dirhodium adduct (49).

CD Data. The circular dichroism spectrum of dsII exhibits a slight shift of the apex point as compared to that of dsI (Figure S5 in the Supporting Information), which indicates that the overall conformation of dsII is similar to that of dsI (B-DNA).

Enzyme Digestion Studies. Exonuclease digestions followed by MS studies were performed on a sample of ssIII (single-stranded adduct D), which has the formula [M + Rh₂(O₂CCH₃)₃], M = ssI, as determined by ESI-MS (Figure S2), before the enzyme digestions. The successive cleavage of nucleotides by both Phosphodiesterases I and II from the 3' and 5' ends of ssIII (single-stranded adduct D), respectively, is inhibited at the A6 and C5 residues (Figure S6 in the Supporting Information), which strongly supports dirhodium binding to these sites (50–52) and corroborates the conclusions from the 2D NMR investigation of dsII.

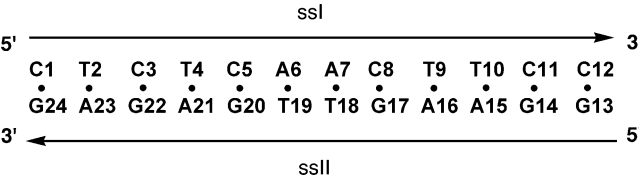
NMR Characterization of dsI. The combined use of NOESY and DQF-COSY experiments allowed the assignment of the protons for dsI (Chart 2), which was performed by standard procedures based on the proximity of the sugar protons H1', H2', H2'', and H3' to the protons on the same and the 3'-adjacent residues in a typical B-DNA conformation

Table 1: ¹H NMR Chemical Shifts (ppm) of the Protons in the Dirhodium Adduct dsII at 15 °C^{a,b}

Base	H6/H8	H2/H5/Me	H1'	H2'	H2''	H3'	H4'	H5'	H5''	N4Hb/N4He ^c	GH1/TH3
C1	7.92	5.94	5.87	2.28	2.59	4.65	4.11	3.80	3.86	7.77/7.19	
T2	7.67	1.67	6.19	2.29	2.59	4.90	4.29				13.88 (−0.09)
C3	7.66	5.66	6.08	2.18	2.54	4.81	4.19			8.36 (0.09)/7.13	
T4	7.36	1.56	6.16 (0.20)	2.05	2.24 (0.19)	4.81	4.24		4.07		<i>d</i>
C5* ⁱ	7.93 (0.44)	6.23 (0.56)	5.70 (0.54)	2.42 (0.44)	2.47 (0.25)	4.44 (−0.35)	4.21 (0.17)	4.07		8.76 (0.19) ^e	
A6* ⁱ	8.74 (0.55)	7.17	6.16 (0.30)	2.75	2.99	5.02	4.40	4.12	4.08		
A7	8.29 (0.17)	7.79 (0.22)	6.23 (0.19)	2.67	2.89	5.03	4.52		4.25		
C8	7.26	5.15	5.78	1.97	2.50	4.67	4.24			7.86 (0.12)/6.65	
T9	7.44	1.48	6.08	2.18	2.59	4.88	4.20	4.11	4.11		14.10
T10	7.47	1.65	6.12	2.20	2.56	4.88	4.23				13.84 (−0.07)
C11	7.64	5.77	6.08	2.28	2.48	4.87	4.21			8.52/7.04	
C12	7.74 (0.20)	5.87 (0.28)	6.22	2.30	<i>f</i>	4.56	4.18 (0.19)	4.07	4.07	<i>g</i>	
G13	<i>f</i>		5.59	2.40	2.53	4.76	4.13	3.62			<i>g</i>
G14	7.85		5.35	2.66	2.71	4.97	4.31	4.08	4.00		12.81 (−0.02)
A15	8.14	7.37	5.94	2.71	2.91	5.07	4.44	4.20	4.12		
A16	8.01	7.53	6.06	2.63	2.89	5.05	4.47		4.24		
G17	7.38		5.77	2.41	2.65	4.84	4.40				12.73 (−0.09)
T18	7.22	1.12	6.08	2.13	2.58	4.83	4.29	4.23	4.16		13.87 (0.03)
T19	7.32	1.56	<i>f</i>	<i>f</i>	<i>f</i>	4.89	4.20				13.11 (−0.67)
G20	<i>h</i>		5.47	2.30	2.48	<i>h</i>	<i>h</i>				11.19 (−1.38)
A21	8.09	7.67	5.85	2.66	2.71	5.00	4.37				
G22	7.75		5.47	2.53	2.65	4.97	4.32				12.60 (−0.09)
A23	8.06	7.83	6.13	2.62	2.89	5.01	4.42		4.13		
G24	7.65		6.00	2.25	2.42	4.63	4.15	4.11			<i>g</i>

^a The acetate resonances for *a*CH₃, *b*CH₃, *c*CH₃ of the dirhodium unit appear at 1.77, 1.93, and 1.99 ppm, respectively. ^b Numbers in parentheses indicate the difference between the chemical shift of the dirhodium adduct dsII and the unmodified duplex dsI resonance [$\Delta\delta = \delta(\text{dsII}) - \delta(\text{dsI})$]; the $\Delta\delta$ values for the nonexchangeable protons are listed if $|\Delta\delta| \geq 0.15$. The other $\Delta\delta$ values are omitted for the sake of simplicity. Positive and negative values of $\Delta\delta$ indicate downfield and upfield shifts, respectively. ^c N4Hb and N4He are the cytosine hydrogen-bonded and exposed amino protons. ^d Thymine imino proton not detected. ^e N4He of C5* is displaced due to metal binding. ^f Proton not detected due to weak or overlapped cross-peaks. ^g Imino and amino protons of frayed terminal bases not detected (74). ^h Proton not detected due to lack of stacking (58, 69) and flexibility (71, 100) or fast conformational exchange (72, 73) of the base. ⁱ The asterisk denotes the residues binding to the dirhodium core.

Chart 2: Sequential Numbering Scheme for dsI



(53) (Figures S7–S10 in the Supporting Information); the values for the chemical shifts of the protons are listed in Table S1 in the Supporting Information. An intense cross-peak between the H2 protons of residues A6 and A7 (Figure S8) indicates that the duplex dsI is in the B-DNA conformation (53).

NMR Spectroscopic Data for dsII. Assignment of Nonexchangeable Protons. NOESY, DQF-COSY, and HSQC spectra were collected and analyzed to assign the resonances for the nonexchangeable protons of dsII. The cross-peaks between the cytosine protons H5 and H6 for dsII were readily identified by their presence in the DQF-COSY spectrum (Figure S11 in the Supporting Information) (54). The chemical shifts of the protons for dsII and the notable changes as compared to the corresponding protons in dsI are listed in Table 1. A global indication of the localization of the structural distortions in dsII is derived by comparing the chemical shifts for the nonexchangeable protons of dsI to those of dsII; these changes are summarized in the graph (Figure 1).

In regards to the aromatic protons of the dsII bases, the resonance for A6H8 exhibits a substantial downfield shift of $\Delta\delta = +0.55$ ppm (Table 1, Figures 1 and 2) as compared to that of A6H8 for dsI (Figure S7), which indicates that the

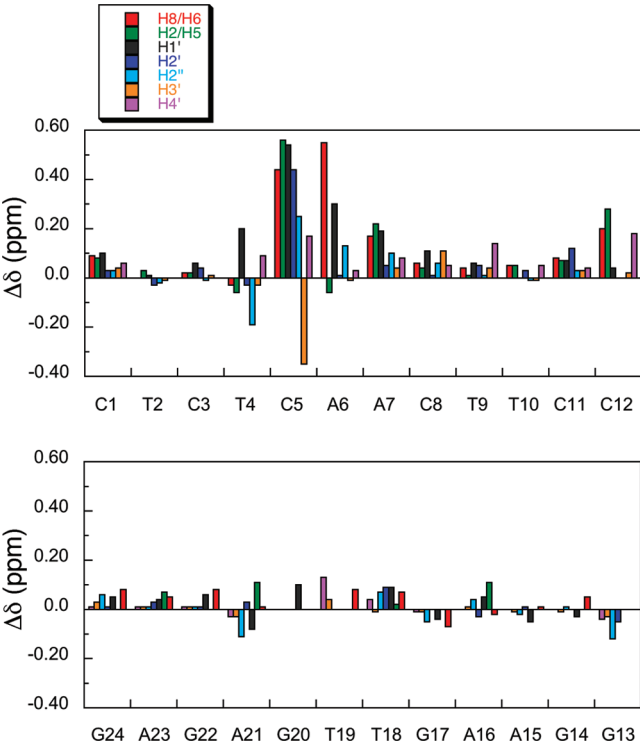


FIGURE 1: Differences in the chemical shifts of the nonexchangeable protons upon binding to the dirhodium unit [$\Delta\delta = \delta(\text{dsII}) - \delta(\text{dsI})$]. Bars in the positive and negative regions indicate downfield and upfield shifts, respectively.

binding of the dirhodium unit takes place at site N7 of the A6 residue. Comparable downfield shifts of the H8 resonance of adenosine ($\Delta\delta \approx +0.55$ ppm) have been recorded in its complex with dirhodium tetraacetate (55). In the case of the

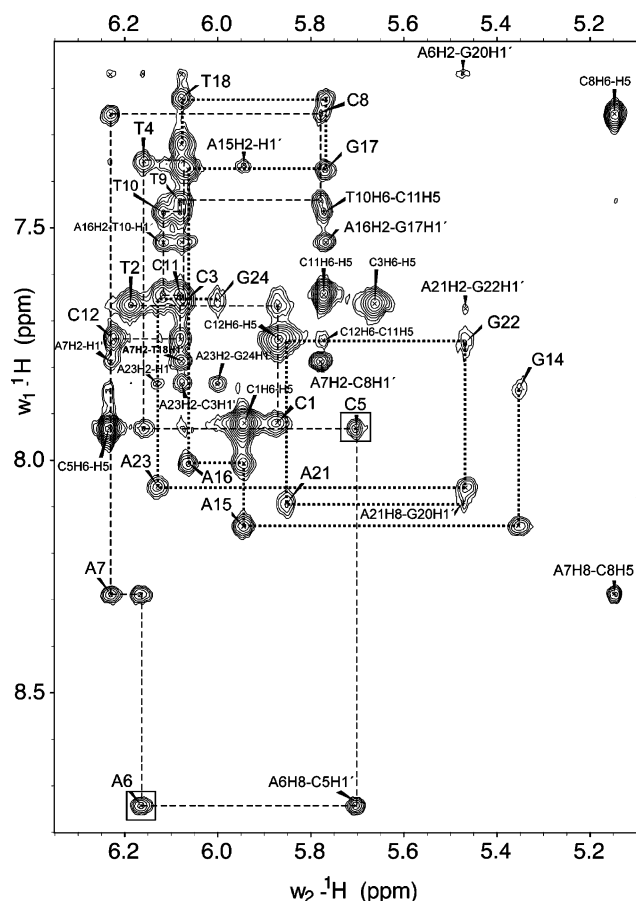
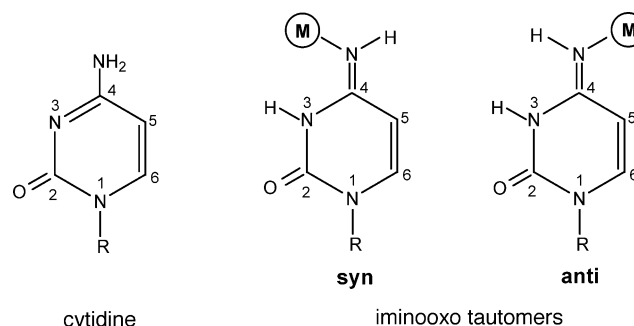


FIGURE 2: Contour plots of the NOESY spectrum for duplex dsII in D₂O (collected at 15 °C, 280 ms mixing time) exhibiting the correlation between the aromatic protons of the bases and the H1' protons of the sugar rings.

interaction of the duplex dsl with the dirhodium complex *cis*-[Rh₂(μ-O₂CCH₃)₂(η¹-O₂CCH₃)(dap)(CH₃OH)](O₂-CCH₃) containing the intercalating group dap (dap = 1,12-diazaperylene), binding also takes place via site N7 of A6, but the downfield shift of A6H8 (Δδ = +0.14 ppm) is counterbalanced by the upfield shift of the resonances due to the intercalation of the dap group between residues A6 and A7 (37). Considerable downfield shifts of the H8 purine resonances of DNA oligonucleotides have also been recorded in platinum–DNA adducts, wherein the metal binds to the N7 sites of the guanine residues (56–58).

In the duplex dsII, notable changes for residue C5 are the $\Delta\delta = +0.56$ and $+0.44$ ppm downfield shifts of the H5 and H6 resonances, respectively, as compared to those of the corresponding protons in dsI. Additionally, the resonances of the deoxyribose protons H1', H2', H2'', and H4' for residue C5 shift downfield by $+0.54$, $+0.44$, $+0.25$, and $+0.17$ ppm, respectively, whereas that of H3' moves upfield by $\Delta\delta = -0.35$ ppm as compared to those of dsI (Table 1). These appreciable chemical shift changes indicate that the C5 residue is bound to the dirhodium unit, inducing considerable structural changes to C5 of dsII. The downfield shifts of the H1', H2', H2'', and H4' sugar protons of C5 (Figure 1) indicate that most likely the C5 residue deoxyribose ring is close to the dirhodium unit, resulting in metal-induced downfield shifts of the protons (59). The upfield shift of the C5H3' sugar proton may be mainly attributed to the neighboring base diamagnetic ring current anisotropy (60). The substantial downfield shifts of C5H5 and C5H6 strongly suggest that the exocyclic site N4 of C5 is binding to the

Chart 3: Neutral and Metal-Stabilized Rare Cytidine Tautomers



rhodium atom (61–63). In particular, characteristic downfield shifts of the C5H5 resonance (63–66) indicate the presence of the *anti* rotamer (67, 68) of the base with respect to N3 in its metal-stabilized rare iminooxo form (Chart 3, vide infra).

For dsII, the sequential contacts that give rise to intra- and internucleotide ($nH1'-(n+1)H6/H8$) NOEs involving base H6/H8 to deoxyribose H1' protons are observed for all the bases, but are absent in the complementary strand opposing the metal lesion site; e.g., the cross-peaks T19H6-H1', T19H1'-G20H8, and G20H8-H1' are absent, and G20H1'-A21H8 is weak (Figure 2). This is an indication of distortion of the stacking interactions between T19 and G20 (58, 69, 70). Moreover, the absence of any NOE contacts for G20H8 (complementary to C5*) is attributed to reduced stacking (58, 69), which results in dynamic behavior or fast conformational exchange (flexibility) of this base on an intermediate NMR time scale (69-73). The aromatic-H1' NOE cross-peaks for the bases C1, C12, and G13 are also missing (Figure 2), most likely due to terminal fraying effects (the changes in the chemical shifts of the C12 residue protons (Table 1) are attributed to fraying effects (74, 75)). The NOE cross-peaks between T4H6 and C5H6, T4H6 and C5H5, and C5H6 and A6H8, which are the bases binding to the dirhodium unit or flanking the binding sites, are very weak or absent in the NOESY spectrum, indicating that the stacking between the aforementioned bases in dsII has been disrupted as compared to that in dsI (58, 69). The intense cross-peak between the H2 protons of residues A6 and A7, however, indicates that the metalated duplex dsII in the 3' direction of the lesion site remains in the B-DNA conformation (53).

The H2 protons of the adenine bases for dsI and dsII were assigned by performing long- and short-range HSQC and NOESY spectra. In the lr-HSQC spectra (Figure S12, top, in the Supporting Information), three-bond correlations are detected through cross-peaks between the H2 protons of adenine bases and the $^{13}\text{C}4$ and $^{13}\text{C}6$ carbon atoms in the respective aromatic rings. In the sr-HSQC spectra (Figure S12, bottom), one-bond correlations are detected, through cross-peaks between the H2 protons and the $^{13}\text{C}2$ carbon atoms. In the sr-HSQC spectrum of dsII (Figure S12, bottom, right), all six H2 protons were identified except for partial overlap between A23H2 and A7H2; the two protons were rather easily identified in the corresponding NOESY spectrum by the correlation peaks with other protons (Figure 2). Overall, there were small changes in the chemical shifts of the carbon atoms of adenine bases for dsII; the resonances

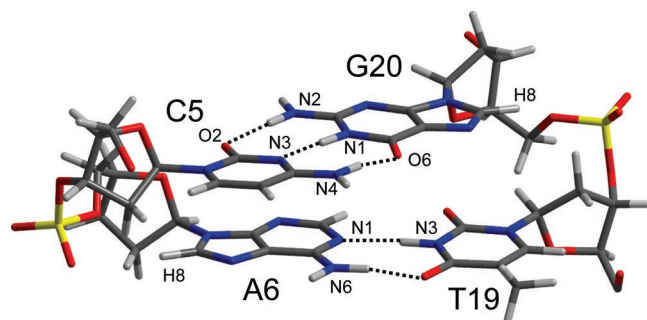


FIGURE 3: Watson–Crick hydrogen bonding between the partner bases C5 and G20 and A6 and T19 in dsI.

of the H2 protons, however, were downfield shifted by 0.1–0.2 ppm, except for A6H2, which exhibited a ~ 0.1 ppm upfield shift (Table 1, Figure 1).

The intrasidue NOE cross-peaks between the base H8/H6 protons and the sugar H1' protons of dsII (except for bases T19 and G20, for which these cross-peaks are missing, Figure 2) are weaker than those detected between the cytosine atoms H5 and H6, e.g., C3, C5, C8, and C11 (Figure 2), which indicates that the H8/H6 to H1' distances for the bases are longer than the fixed length of 2.4 Å between H5 and H6 in cytosine (53), establishing that in dsII each base adopts an *anti* conformation about its glycosyl bond (74). Moreover, the intensities of the C5H6–H3' and A6H8–H3' cross-peaks in the NOESY spectrum of dsII, when examined qualitatively, are comparable in intensity to the aromatic–H3' cross-peaks for the other bases of the duplex, indicating that the deoxyribose rings at the metal binding site are S-type (56, 57) as in B-DNA.

Assignment of the Exchangeable Protons. Wateregate-NOESY spectra in H₂O/D₂O (90:10) were collected to assign the exchangeable protons of dsII. Characteristic features of the G–C base pair Watson–Crick hydrogen bonding in double-stranded DNA (Figure 3) are the cross-peaks between both the N4Hb (hydrogen-bonded) and N4He (exposed) amino protons of the cytosine base and the N1H imino proton of the partner guanine base (Figure 4, top; e.g., C3N4He–G22N1H, C3N4Hb–G22N1H, C5N4He–G20N1H, C5N4Hb–G20N1H, C8N4He–G17N1H, C8N4Hb–G17N1H) (74). In dsI, both amino protons of each cytosine residue give NOE cross-peaks with the H5 resonance of the same base except for the terminal residues (Figure S13, top, in the Supporting Information), and thus, the guanine N1H resonances are linked to the backbone cytosine protons. In the case of dsII, however, for residue C5, only the C5H5–N4Hb cross-peak is observed (Figure S13, bottom) as opposed to the presence of both C5H5–N4Hb and C5H5–N4He cross-peaks for dsI (Figure S13, top). Moreover, the N1H proton of G20 has a cross-peak with C5N4Hb only, but not with C5N4He (Figure 4, bottom). The previous findings imply that one of the exocyclic amino group protons of C5 has been displaced, due to the dirhodium binding to site N4 of the C5 residue (68). The resonance for C5N4Hb in dsII is downfield shifted by $\Delta\delta = +0.19$ ppm as compared to that of dsI and appears at 8.76 ppm (Table 1), which is essentially the downfield extreme of the typical range for hydrogen-bonded exocyclic amino protons (6.6–9.0 ppm) (53). The cross-peak between G20N1H and C5N4Hb (Figure 4, bottom) implies that C5 and G20 are still close to each other. As indicated earlier, the metal adopts an *anti* orientation with respect to N3, which

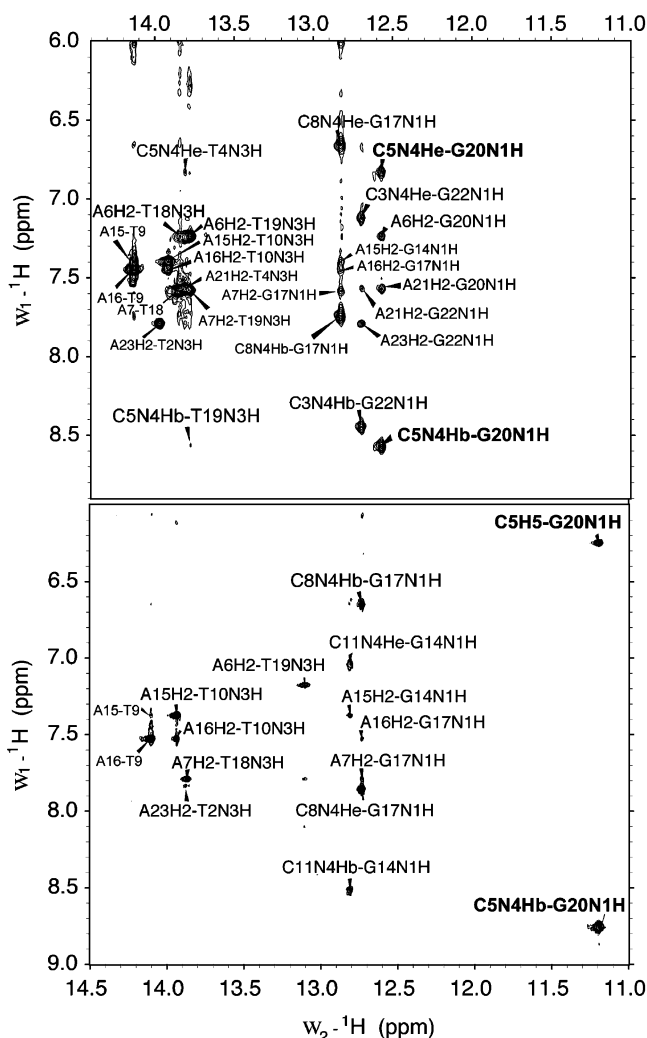


FIGURE 4: Contour plots of the Wateregate-NOESY spectra (collected at 15 °C, 200 ms mixing time) of duplexes dsI (top panel) and dsII (bottom panel) showing NOE correlations between the imino and amino protons.

is required for base C5 to remain close to G20. The considerable downfield shift of the C5 amino proton N4Hb ($\Delta\delta = +0.19$ ppm) is attributed to the inductive effect of the metal binding at the N4 site (63, 76) or may reflect a decrease in the extent of stacking interactions involving C5 (74, 77). In the 3' direction of the lesion site, both the base pair cytosine N4H–guanine N1H partner cross-peaks are observed (e.g., C8N4Hb–G17N1H, C8N4He–G17N1H, C11N4Hb–G14N1H, C11N4He–G14N1H), although weaker than in dsI. In contrast, in the 5' direction of the lesion site, the C3N4Hb–G22N1H and C3N4He–G22N1H cross-peaks are absent (Figure 4, bottom), although both C3 amino protons were found to have NOE cross-peaks to the H5 proton of the same base (Figure S13, bottom); this indicates that the Watson–Crick hydrogen bonding in C3–G22, which is two base pairs away from the lesion site in the 5' direction, is destabilized (58, 74, 78).

The N1H imino protons of the guanine and the N3H imino protons of the thymine residues participate in Watson–Crick hydrogen bonding in native DNA (Figure 3, G–C and A–T pairs, respectively) and appear in the range 12.0–14.5 ppm for dsI (Figure 5, top; Table S1). In general, the imino protons of dsII exhibit moderate upfield shifts (-0.1 ppm) as compared to the resonances for the corresponding imino

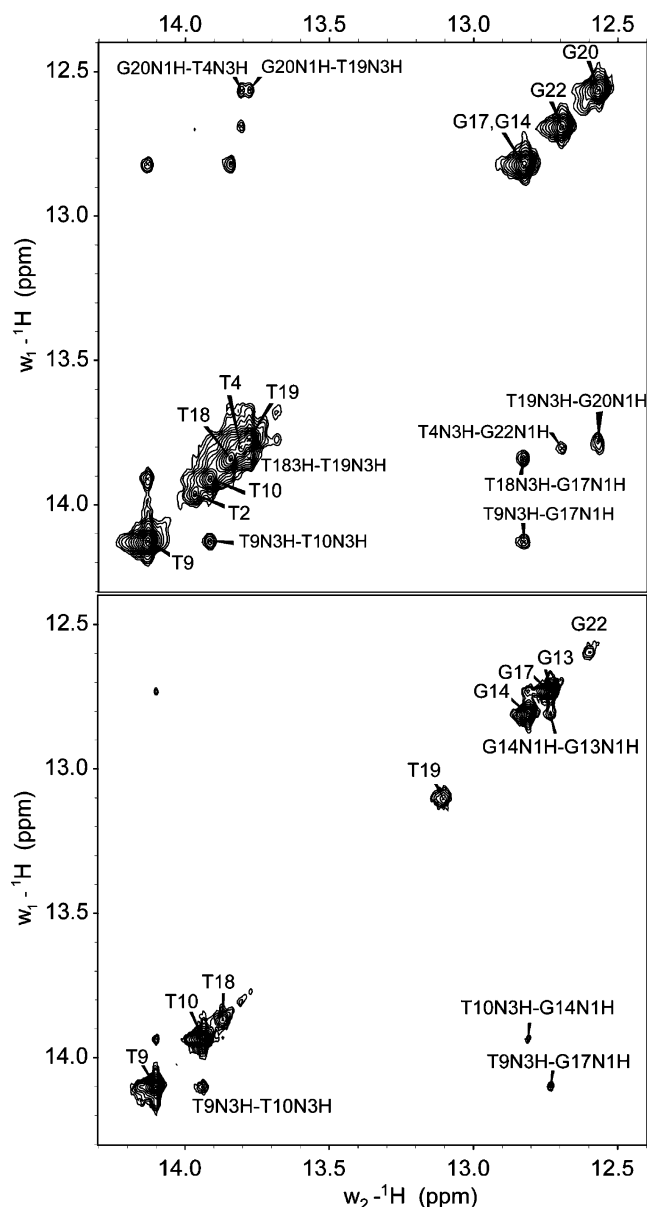


FIGURE 5: Contour plots of the Watergate-NOESY spectra (collected at 15 °C, 200 ms mixing time) of duplexes dsI (top panel) and dsII (bottom panel) showing the imino–imino proton NOE correlations. For dsII, G20N1H is further upfield from the region shown.

protons of dsI, except for the substantial upfield shifts in the resonances of the imino protons for T19N3H ($\Delta\delta = -0.67$ ppm) and G20N1H ($\Delta\delta = -1.38$ ppm) (Figure 4, bottom; Table 1), which are the partner bases to the binding sites in dsI (Figure 3). The substantial upfield shift of T19N3H indicates weakening of the hydrogen bonds to A6 and in the case of G20N1H complete disruption of G20N1H–C5N3 (Figure 3); the latter is supported by the value of $\delta = +11.2$ ppm for G20N1H (74, 77, 79, 80) and the broadness of the G20N1H resonance at 2 °C in 90% H₂O/10% D₂O (78,81), although substantially upfield-shifted imino guanine protons have been recorded in spite of the protons being involved in hydrogen bonding (82).

As expected for dsI, the nonterminal imino protons exhibit NOE cross-peaks to the neighboring imino protons in the same strand (Figure 5, top; e.g., T9N3H–T10N3H, G17N1H–T18N3H, T19N3H–G20N1H) as well as to the imino

protons in the adjacent base pairs (e.g., T4N3H–G20N1H and T4N3H–G22N1H in the 3' and 5' directions, respectively, and T9N3H–G17N1H). In the case of dsII, however, the pattern of connectivities for the imino protons is disrupted in the 5' direction of the lesion site; e.g., T4N3H–G20N1H, T4N3H–G22N1H, and T19N3H–G20N1H are absent (Figure 5, bottom). The G17N1H–T18N3H cross-peak is absent, and the T9N3H–G17N1H cross-peak is much weaker than in dsI.

Evidence of Watson–Crick hydrogen bonding between DNA complementary strands also stems from the correlation peaks between the H2 proton of the adenine residue and the N3H imino protons of the thymine residue in the same base pair and the adjacent ones (Figure 4, top; e.g., A6H2–T19N3H, A6H2–T18N3H, A7H2–T19N3H, A15H2–T9N3H, A16H2–T9N3H, A16H2–T10N3H, A21H2–T4N3H) as well as to the adjacent guanine N1H imino protons, e.g., A15H2–G14N1H, A16H2–G17N1H, A21H2–G20N1H, A21H2–G22N1H, A23H2–G22N1H, A6H2–G20N1H, A7H2–G17N1H (53). For dsII (Figure 4, bottom), the correlation peaks for the base pairs A6H2–T19N3H and A7H2–T18N3H are observed, although much weaker, but A6H2–T18N3H, A7H2–T19N3H, A6H2–G20N1H, A21H2–G22N1H, and A23H2–G22N1H between neighboring base pairs and A21H2–T4N3H between partner bases are absent (A23H2–T2N3H is weak). On the contrary, the A15H2–T10N3H, A16H2–T10N3H, A15H2–T9N3H, A16H2–T9N3H, A15H2–G14N1H, and A16H2–G17N1H correlations for the bases in the 3' direction of the lesion site are observed, although weaker than in dsI (Figure 4, bottom). The previous findings suggest that the base stacking and the hydrogen bonding are considerably disrupted in the 5' direction of the lesion site, which corroborates the absence of the imino–imino correlation peaks in the same direction (vide supra).

Dirhodium Unit Acetate Groups. As indicated by the mass spectrum of ssIII (Figure S2, adduct D) and the acetate region of the NOESY spectrum for dsII (Figure S14 in the Supporting Information), only three acetate groups remain attached on the dirhodium core in dsII. The acetate groups on the dirhodium unit of dsII, referred to as *a*CH₃, *b*CH₃, and *c*CH₃, resonate at 1.77, 1.93, and 1.99 ppm, respectively (the acetate groups in Rh₂(O₂CCH₃)₄ resonate at 1.89 ppm in D₂O). The different chemical shifts of the three dirhodium acetate groups in dsII indicate that, as expected, they are not equivalent. On the basis of the values of their chemical shifts for dsII, the *b*CH₃ and *c*CH₃ groups are bridging the dirhodium core and *a*CH₃ is monodentate. The previous NMR assignments are supported by the chemical shifts of acetate groups in dirhodium compounds with one dangling and two bridging acetate groups; typically, the monodentate groups resonate upfield (~ 1.7 ppm) from the bridging acetate ligands (83, 84). The presence of a monodentate acetate group in dsII is also corroborated by the modeling studies (steric reasons). For dsII, an NOE cross-peak is observed between acetate *b*CH₃ and A6H8 (Figure S14), which is one of the binding residues. The monodentate acetate group *a*CH₃ exhibits NOE cross-peaks to C3H6 and T4H6 in the 5' direction of the lesion site, and the bridging group *c*CH₃ to T18H1', T18CH₃, and G17H8 in the 3' direction of the lesion site (Figure S14); these cross-peaks indicate that the acetate group *a*CH₃ is directed toward the 5' end of ssI, whereas the acetate group *c*CH₃ is situated toward the 5' end of the complementary strand ssII (Figure 6).

DISCUSSION

The structural characterization of the duplex d(CTCTC*A*ACTTCC)·d(GGAAGTTGAGAG) with the dirhodium lesion site indicates that considerable distortions are induced to the DNA structure upon binding to the metal, especially at the base pairs located at and near the binding sites, but the global conformation of dsII appears to be mainly that of dsI, i.e., B-DNA. This is supported by the CD data of dsII, the intense cross-peak between the H2 protons of the dsII residues A6 and A7 (53), and the NOE proton connectivity pattern along the two strands of the modified duplex (53, 74), except for bases T19 and G20, which are the partner bases to the metal binding sites. The considerable downfield shifts of the aromatic proton resonances for C5 and A6 ($\Delta\delta \approx 0.5$ ppm) of dsII relative to dsI are consistent with the helix being right-handed, as usually observed for GG intrastrand cisplatin DNA duplexes (85). The duplex dsII, however, has been considerably destabilized as indicated by the substantial decrease in the melting temperature by $\Delta T_m = -22.9$ °C, as compared to dsI. The reduced thermal stability of dsII is attributed to the decreased stacking of the bases and the complete disruption and/or weakening of the hydrogen bonds within the base pairs (86–88) due to metal binding (vide infra). The decrease in the T_m value for dsII ($\Delta T_m = -22.9$ °C) suggests the presence of a covalent *intrastrand* dirhodium adduct (49), which is corroborated by the NMR data. The formation of intrastrand cisplatin cross-linked adducts at GG sites in various DNA sequences induces a decrease in the melting temperatures of the host duplexes by $\Delta T_m = 15$ –27 °C (88), which is comparable to that for the dirhodium adduct dsII. When the duplex dsI binds to the dirhodium unit *cis*-[Rh₂(μ -O₂CCH₃)₂(dap)]²⁺, the net decrease in the melting point of dsI is considerably smaller ($\Delta T_m = -6.0$ °C) due to intercalation of the dap group between residues A6 and A7, apart from covalent binding of the A6 residue to the metal via N7 (37).

The halt of the Phosphodiesterases I and II at the A6 and C5 residues of ssIII, respectively (Figure S6), strongly supports dirhodium binding to these bases. The large downfield shift of A6H8 ($\Delta\delta = +0.55$ ppm, Table 1), which is comparable to that of the H8 resonance of adenosine ($\Delta\delta \approx +0.55$ ppm) in its complex with dirhodium tetraacetate (55), is attributed to the inductive effect of the metal (89) and indicates that site N7 of the A6 residue is binding to the dirhodium unit (Figure 6). In the case of the duplex dsI with the dirhodium complex *cis*-[Rh₂(μ -O₂CCH₃)₂(η^1 -O₂-CCH₃)(dap)(CH₃OH)](O₂CCH₃), binding takes place via site N7 of A6 and the dap group intercalates between residues A6 and A7 (37). Apparently, for the given DNA sequence, site A6N7 is the preferable binding site in the presence (37) or absence (present study) of intercalation. In the crystal structure of dirhodium tetraacetate with tRNA^{phe}, however, axial binding of the A36 base takes place via site N1 (90), but in the case of dsII, the change in the chemical shift of A6H2 is negligible (Table 1), thus excluding N1 binding for base A6.

In the present dirhodium adduct dsII, C5 is the other base that establishes a coordination bond with the dirhodium unit (Figure 6). This is strongly supported by the enzyme digestion data, as well as by the considerable changes of the chemical shifts for the protons of C5 (Table 1). Although

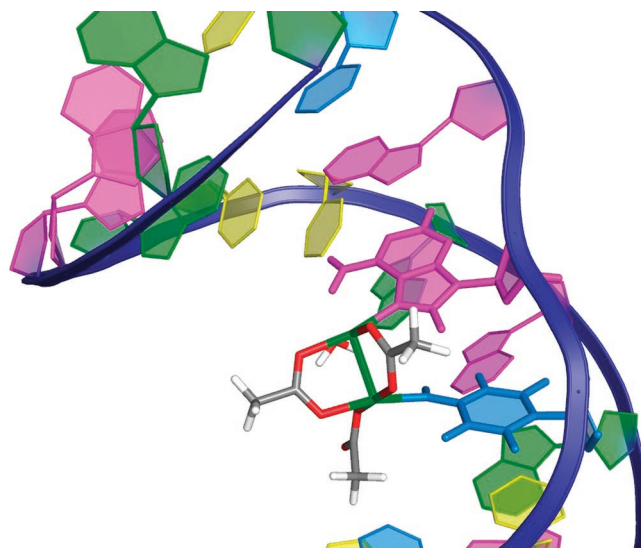


FIGURE 6: Model of dsII depicting the binding sites (C5N4 and A6N7) of the dirhodium unit *cis*-[Rh₂(μ -O₂CCH₃)₂(η^1 -O₂CCH₃)]⁺. DNA base colors: A, pink; C, light blue; G, green; T, yellow. Blue ribbon = phosphodiester backbone. Atom colors: Rh, green; O, red; C, gray; H, white.

cytosine has a number of potential binding sites for the metal (Chart 3; e.g., N3, O2, N3/O2, N3/N4 (91)), metal binding to N7 of A6, which is in the major groove of DNA, and the substantial downfield shift of C5H5 ($\Delta\delta = +0.56$ ppm) (61–63) render the C5 exocyclic amino group N4, also located in the major groove (92), the suggested cytosine binding site of the dirhodium unit (Figure 6). Characteristic downfield shifts of the C5H5 resonance are typical of the *anti* cytosine rotamer (with respect to N3) in its metal-stabilized rare iminoxo form (Chart 3) (64–66); e.g., for the platinum complex *trans*-[Pt(NH₃)₂(1-MeC-N4)₂](NO₃)₂, downfield shifts of ~ 0.5 – 0.7 ppm are observed for the cytosine H5 protons of the *anti/anti* rotamers (63). These appreciable downfield shifts of the cytosine resonances are attributed to the magnetic anisotropy of the d⁸ Pt(II) atom in a square planar arrangement, which deshields protons situated directly above or below the coordination plane (63). The presence of the *anti* rotamer of cytosine is corroborated by the cross-peak between the N1H proton of G20 and C5N4Hb (Figure 4, bottom), which indicates close proximity of C5N4Hb and G20N1H in dsII (4.3 Å in the model, Figure 7), although Watson–Crick hydrogen bonding between G20 and C5 is disrupted (e.g., in the model for dsII, G20O6–C5N4Hb = 6.5 Å and G20N2H–C5O2 = 5.2 Å, and the partner bases G20/C5 are no longer coplanar, Figure 7); this is supported by the significant upfield shift of the imino proton G20N1H resonance ($\Delta\delta = -1.38$ ppm), the value of its chemical shift ($\delta = +11.2$ ppm, Table 1) (74, 77–80), and its broadness in 90% H₂O/10% D₂O at 2 °C (78, 81). On the other hand, the resonance for C5N4Hb in dsII is downfield shifted by $\Delta\delta = +0.19$ ppm as compared to that in dsI and appears at 8.76 ppm (Table 1), which is essentially the downfield extreme of the typical range for hydrogen-bonded exocyclic amino protons (6.6–9.0 ppm) (53). The considerable downfield shift of the C5 amino proton N4Hb ($\Delta\delta = +0.19$ ppm), however, is not due to G20O6–C5N4Hb hydrogen bonding (Figure 3), but rather is attributed to the inductive effect of the metal binding at the N4 site (63, 76); e.g., the chemical shifts of the N4 exocyclic amino protons are downfield

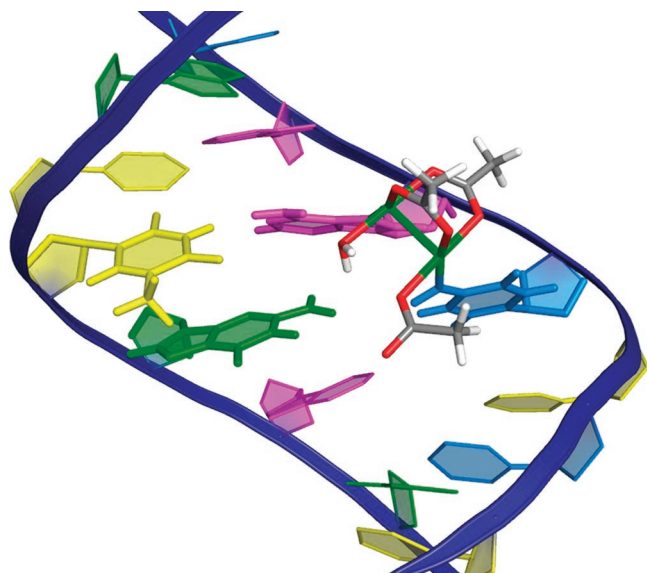


FIGURE 7: Model of dsII depicting the partner bases C5•G20 and A6•T19 and the bases C5 and A6 binding to the dirhodium unit *cis*-[Rh₂(μ-O₂CCH₃)₂(η¹-O₂CCH₃)₂]⁺. DNA base colors: A, pink; C, light blue; G, green; T, yellow. Blue ribbon = phosphodiester backbone. Atom colors: Rh, green; O, red; C, gray; H, white.

shifted by ~1.2 ppm for the isomers of the platinum complex *trans*-[Pt(NH₃)₂(1-MeC-N4)₂](NO₃)₂ as compared to free cytosine (spectra collected in DMSO-*d*₆) (63, 76). The metalation of the exocyclic amino group N4 of cytosine is known to stabilize the rare iminooxo form of the base and induce protonation of site N3 (Chart 3; protonation of C5N3H inhibits formation of the G20N1H–C5N3 hydrogen bond, Figures 3 and 7) (68, 91), thus leading to rare nucleobase tautomers.

A theoretical study that has been performed on N4-metalated cytosine with Hg(I) and Pt(II) showed that the metalation of the amino group of the nucleobases improves the protonation energy of the nucleobase aromatic ring by 10–14 and 30–34 kcal/mol, respectively (93). Protonation of site N3 of cytosine, which is induced by metalation of N4, may lead to stabilization of mispairs such as a CH⁺•G Hoogsteen base pair or a G[–]•CH⁺ ion pair, with significant biological consequences (68, 93). In the case of the dirhodium adduct dsII, however, the CH⁺•G Hoogsteen base pair (94) is ruled out due to the presence of the NOE cross-peak between G20N1H and C5N4Hb (Figure 4, bottom), and a G[–]•CH⁺ ion pair is not considered because G20N1H continues to be protonated at the pH of the 2D NMR data collection. The observation of G20N1H cross-peaks, although this proton is not hydrogen bonded, is consistent with slow exchange on the NMR time scale or minimal exposure to the solvent (74, 95), most likely due to shielding from the minor groove by G20(NH2) (Figure 7). It is notable that the rate of exchange of the GN1H imino proton has been correlated to the relative position of the kink (bend) of the helix with respect to the G base (96–98).

The conformational changes and the Watson–Crick hydrogen-bonding disruption of the C5•G20 base pair are also indicated by the absence of the G20N1H–A6H2 NOE cross-peak (Figure 4), as opposed to the 3′ flanking base pair A6•T19, which still retains hydrogen bonding, albeit significantly weaker than in dsI, on the basis of the substantial upfield shift in the chemical shift of the imino

proton T19N3H ($\Delta\delta = -0.67$ ppm, Table 1) and the intensity of the cross-peak A6H2–T19N3H (Figure 4, bottom) (74, 77). The absence of T19H1′–G20H8 and G20H8–H1′ (Figure 2) and the imino T19N3H–G20N1H (Figure 5, bottom) and the weak G20H1′–A21H8 cross-peaks suggest disruption of the stacking interactions among T19, G20, and A21 (58, 69), which together with the absence of any NOE contacts for G20H8 (95) corroborate high flexibility of residue G20 (69–73). Lack of the sequential *n*H1′–(*n* + 1)H6/H8 NOE cross-peaks at the C16–C17 and C17–G18 steps of the d(GCCG*G*ATCGC)•d(GCGATC-CGCG) duplex bearing a cisplatin unit cross-linking the two G* bases is attributed to an unusual position of C17 with respect to both its neighbors; i.e., the G4*–C17 base pair slides back and forth along a vector perpendicular to the local helix axis (99, 100). Furthermore, the substantial downfield chemical shifts of the deoxyribose protons H1′, H2′, H2′′, and H4′ (+0.54, +0.44, +0.25, and +0.17 ppm, respectively) for the partner base C5 (Table 1) indicate that the C5 deoxyribose ring may be close to the dirhodium unit (59), whereas the upfield shift of the C5H3′ sugar proton may be attributed to its positioning in the shielding cone of residue A6 (100); the absence (or weakness) of the NOE cross-peaks between the aromatic protons for T4, C5, and A6 and the missing contacts A6H2–T18N3H and A7H2–T19N3H indicate that stacking interactions are disrupted for T4, C5, and A6 as well (58, 69). The hydrogen bonding for the base pairs T4•A21 and C3•G22 is disrupted, and that of T2•A23 is weak (vide supra). It is obvious from the aforementioned observations that major disruptions are induced in the stacking and the Watson–Crick hydrogen bonding three base pairs away from the lesion site in the 5′ direction. On the contrary, the hydrogen-bonding interactions between the base pairs in the 3′ direction to the lesion site A6•T19 are still present (A7•T18, C8•T17, T9•A16, T10•A15, C11•G14 base pairs), although weaker than in dsI (Figure 4); the stacking interactions are preserved in the 3′ direction to the lesion site but disrupted to an extent among G17, T18, and T19 on the basis of the absence of the imino–imino cross-peaks (Figure 5, bottom). Considering that binding takes place at base A6 at the center of the duplex and the flanking base C5 in the 5′ direction to it, there are four base pairs above C5•G20 (two C•G base pairs and two A•T base pairs) and six base pairs below A6•T19 (three C•G base pairs and three A•T base pairs), which explains the different stabilities of each stem, the severe effect of the metal binding on the shorter stem in the 5′ direction to the lesion site, and its higher exposure to the solvent (DNA breathing) (74, 95).

CONCLUSION

The detailed 2D NMR data analysis performed on dsII revealed that the duplex d(CTCTC*A*ACTTCC)•d(GGAAGTTGAGAG) forms an unprecedented *intrastrand* adduct with the dirhodium unit binding residues C5 and A6 (also corroborated by enzyme digestion studies), which significantly destabilizes the duplex as indicated by the substantial decrease in its melting temperature ($\Delta T_m = -22.9$ °C) as compared to that of dsI. The reduced thermal stability of dsII is attributed to the decreased stacking of the bases and the complete disruption and/or weakening of the hydrogen bonds within the base pairs in the immediate vicinity of the modification site (C5•G20 and A6•T19) as

compared to those in the parent duplex, but the effects due to the metal binding are more severe for the base pairs in the 5' direction to the lesion site. The NMR spectroscopic data indicate that Watson–Crick hydrogen bonding is completely disrupted for the dsII lesion site C5•G20 and considerably weakened for the site A6•T19. In dsII, the bases C5 and A6 bind to *eq* positions of the dirhodium unit $cis\text{-}[\text{Rh}_2(\mu\text{-O}_2\text{CCH}_3)_2(\eta^1\text{-O}_2\text{CCH}_3)]^+$, which retains one monodentate and two bridging acetate groups, presumably due to steric reasons. Binding of A6 takes place via N7, whereas binding of the C5 base takes place via the exocyclic N4 site, resulting in the *anti* cytosine rotamer with respect to site N3 in its metal-stabilized rare iminoxo form. Formation and stabilization of rare base forms and mispairs can be one of the factors that contribute to the antitumor activity of dirhodium compounds.

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SUPPORTING INFORMATION AVAILABLE

Figure S1, HPLC spectra of the reaction mixture of $\text{Rh}_2(\text{O}_2\text{CCH}_3)_4$ with ssI, Figure S2, ESI-MS spectra of isolated single-stranded dirhodium adducts A–E from purification by consecutive anion exchange and reversed-phase column chromatography, Figure S3, anion exchange column chromatograms of the isolated dirhodium adducts dsA–dsE, Figure S4, thermal denaturation curves of dsI and dsII, Figure S5, CD spectra of the double-stranded DNA species dsI, dsC (green), dsD (dsII), and dsE, Figure S6, ESI-MS spectra of the enzyme digests of ssIII (single-stranded adduct D) with Phosphodiesterases I and II, Figure S7, section of the NOESY spectrum of dsI in D_2O at 15 °C, 280 ms mixing time, in the region of the aromatic to H1' protons, Figure S8, aromatic to aromatic proton region from the NOESY spectrum at a 280 ms mixing time for dsI in D_2O at 15 °C, Figure S9, expansion of the aromatic to H3' proton region of the NOESY spectrum for dsI in D_2O at 15 °C at a 280 ms mixing time, Figure S10, expansion of the aromatic to H4' proton region of dsI from the NOESY spectrum in D_2O at 15 °C and a 280 ms mixing time, Figure S11, superimposed DQF-COSY spectra for dsI and dsII, Figure S12, expansion of the Ir-HSQC and sr-HSQC spectra for dsI and dsII, Figure S13, expansions of NOE correlations between imino and aromatic protons for dsI and dsII from the Watergate-NOESY spectra at 15 °C, Figure S14, NOESY spectrum for dsII in D_2O at 15 °C and a 280 ms mixing time in the dsII proton acetate region, computational details for the molecular modeling, Figure S15, TPSS-optimized complex $cis\text{-}[\text{Rh}_2(\mu\text{-O}_2\text{CCH}_3)_2(\eta^1\text{-O}_2\text{CCH}_3)(\text{A-N7})(\text{C-N4})\text{-(H}_2\text{O})]^+$, Table S1, ^1H NMR chemical shifts of the protons in dsI at 15 °C, and full author list for ref 46. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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