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Penetrative DNA intercalation and G-base selectivity of an organometallic tetrahydroanthracene Ru\textsuperscript{II} anticancer complex†

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The organometallic Ru\textsuperscript{II} arene complex \[(\eta^6-tha)Ru(en)Cl\]\textsuperscript{+} (I), where tha = tetrahydroanthracene and en = ethylenediamine, is potently cytotoxic towards cancer cells. We have used a combination of HPLC, ESI-MS, 1D- and 2D-NMR, including \[^1H,^1H\] ROESY, NOESY, \[^1H,^{15}N\] HSQC (using \[^{15}N\]-I), and \[^1H,^{15}P\] experiments to elucidate the role of the non-aromatic, bulky rings of tha in adducts with the DNA hexamer d(CGGCCG), since DNA is a potential target for this drug. Reactions of I with single-stranded d(CGGCCG) gave rise to ruthenation at each of the three G bases, whereas reactions of the duplex (d(CGGCCG))\textsubscript{2} with 1 mol equiv. I led to exclusive ruthenation of G\textsubscript{3} and G\textsubscript{6} (and G\textsubscript{4}, G\textsubscript{5}) and not G\textsubscript{2} (or G\textsubscript{7}). Addition of a second mol equiv. of I gave di-ruthenated adducts (major sites G\textsubscript{3}/G\textsubscript{6}, G\textsubscript{4}/G\textsubscript{5}, G\textsubscript{6}/G\textsubscript{7}), and on reaction with a third mol equiv. tri-ruthenation (G\textsubscript{3}, G\textsubscript{4}/G\textsubscript{5}/G\textsubscript{7}). The NMR data are indicative of the coordinative binding of Ru-tha specifically to G\textsubscript{3} and G\textsubscript{6}, together with penetrative intercalation of the bulky non-coordinated tha rings B and C of I, selectively between two base pairs G\textsubscript{3}/C\textsubscript{6},G\textsubscript{4}/G\textsubscript{5} and G\textsubscript{4}/C\textsubscript{5},C\textsubscript{6}/G\textsubscript{7}. Intercalation at GpC base steps by tha has a lower energy penalty compared to intercalation at GpG base steps, thereby allowing accommodation of tha. Mono-intercalation of tha reduced the strength of H-bonding between en-NH and GO\textsubscript{6}. These differences in structural distortions compared to cisplatin induced by the coordinative binding of Ru-tha to GN7 may contribute to the differences in mechanism of action, including protein recognition of the metallated lesions, and lack of cross resistance.

Introduction

There is currently much interest in the design and mechanism of action of ruthenium anticancer complexes.\textsuperscript{1–7} Structural distortions of DNA are thought to play a major role in the anticancer activity of many ruthenium and other transition metal complexes.\textsuperscript{1,8–23} It is apparent that distortions induced in DNA by \[(\eta^6-arene)Ru(en)Cl\]\textsuperscript{+} organometallic ruthenium(ii) arene anticancer complexes\textsuperscript{1,2,21,24–28} (en = ethylenediamine) differ significantly from those induced by cisplatin.\textsuperscript{9,10} In particular the arene appears to play a significant role in DNA interactions and the cytotoxicity of complexes shows a strong dependence on the arene.\textsuperscript{19,22,23,29,31} The cytotoxic activity of these ruthenium arene complexes appears to increase with the size of the coordinated arene,\textsuperscript{1,2,24,25} p-cymene (Ru-cym) < biphenyl (Ru-bip) < dihydroanthracene (Ru-dha) < tetrahydroanthracene (Ru-tha, I, Fig. 1), the latter complex being as cytotoxic as the clinical platinum drug cisplatin. If the arene is extended, the possibility arises of intercalation between DNA base-pairs. Studies of the interactions of ruthenium complexes Ru-cym and Ru-bip with the 6-mer duplex d(CGGCCG)\textsubscript{2} revealed that the binding of Ru to GN7 is accompanied by strong H-bonding between GO\textsubscript{6} and en-NH, and that the arene ligand distorts the duplex either via steric interactions (Ru-cym) or via intercalation\textsuperscript{19,22,23,32,38} (Ru-bip), which may explain why the IC\textsubscript{50} values of complexes Ru-cym and Ru-bip are similar.\textsuperscript{1,2,24,25} Further work\textsuperscript{21} revealed

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Fig. 1 Structures and NMR numbering schemes for \[(\eta^6-tha)Ru(en)Cl\]\textsuperscript{+} (I) and for the 5’-d(CpGp) fragment of the hexamer d(CGGCCG); single-strand (ss) I and duplex II.
a unique mode of binding of Ru-bip to a 14-mer DNA duplex: the monofunctional fragment \( ((\eta^5\text{-biphenyl})\text{Ru(en)}) \\) is highly specific for GN7, although mobile at elevated temperature (migrating to other G residues). The uncoordinated phenyl ring of bip can be involved in π–π stacking with DNA bases either via classical intercalation\(^9,15-38\) between the bases, or with a partially extruded T base\(^23,39,40\).

Arene–base stacking may play a role in determining the rates of reactions of Ru\(^{11}\) arene complexes with DNA, as appears to be the case for mononucleotides. We have shown previously that the tha and bip arenes can exert slightly different effects on the chemical reactivity of these Ru\(^{11}\) complexes and on distortions induced in DNA\(^26,29,30\). For example the rate of reaction of Ru-bip with cGMP is ca. \(2 \times\) slower than that of I,\(^27\) but Ru-bip induces similar unwinding of DNA as complex I (14\(^+\)).\(^38\) Both of these complexes can potentially intercalate into DNA, leading to “downstream” effects on DNA processing and repair mechanisms. Ru-bip is an aromatic intercalator,\(^22,23,41\) in which all the protons of the extended ring are within the aromatic plane, and can form π–π interactions with bases; however, the extended rings A and C of tha in I are bulky non-aromatic groups; the three rings A, B and C are not in the same plane and the H5,8 and H9,10 protons are located above or below the arene ring plane by nearly 0.9 Å. The extended rings B and C cannot form π–π interactions with bases in the same way that aromatic intercalators do and so Ru-tha is more sterically demanding than Ru-bip. It is reasonable to predict that the intercalative interactions between duplex DNA and Ru-tha or Ru-bip should be quite different. The Ru-tha complex I is 10 times more toxic to cancer cells than Ru-bip.\(^24,25\)

Bulky substituents at the sites of DNA lesions may activate nucleotide-excision repair;\(^42,43\) however, reports of intercalation by bulky molecules are rare. Gomez-Pinto \textit{et al.}\(^44\) have shown that intercalation of a modified nucleotide containing a cholesterol derivative into a DNA decamer induces DNA distortions which are different from those induced by aromatic intercalators.\(^45,46\)

Threading intercalation has attracted recent attention because the intercalator occupies and interacts strongly with both the minor and major grooves of DNA simultaneously. This has been observed for polyaromatic intercalators,\(^7-9,47-49\) dinuclear metallointercalators,\(^50-53\) and platinum complexes with aromatic side-arms, such as acridine-9-ylihexaurea.\(^54-56\) As a result, threading intercalative interactions promise high DNA binding affinity and specificity, a slow rate of dissociation, and an enhanced ability to block DNA–protein interactions.\(^50,57\)

It is important to understand the mode of interaction between \([((\eta^5\text{-tha})\text{Ru(en)})\text{Cl}^-]\) (I) and DNA since a major contributor to its high potency appears to be the lack of repair of the lesions formed on DNA by this complex.\(^29\) \textit{i.e.} lack of recognition by repair enzymes. In the present work we have investigated the role of the extended non-aromatic bulky tha in interactions of \([((\eta^5\text{-tha})\text{Ru(en)})\text{Cl}^-]\) (I) with the single-stranded hexamer d(CGGCCG) and double-stranded duplex d(CGGCCG)\(_2\) using a wide variety of experimental techniques including HPLC, ESI-MS and 1D \(^1\)H and 2D \([\text{H},\text{H}]\) ROESY, NOESY, \([\text{H}, \ ^{15}\text{N}]\) HSQC (using \(^{15}\text{N}-\text{I}\)), and \([\text{H}, \ ^{31}\text{P}]\) NMR spectroscopy.

### Results

Scheme 1 indicates the reaction pathways that were followed during the course of this study. HPLC enabled separation of single DNA strands even for reactions of the duplex. The 1 : 1, 2 : 1 and 3 : 1 I/I\(_{-}\)mixtures, 1 : 1 and 2 : 1 I/II mixtures were studied by HPLC and ESI-MS, and 1 : 1, 2 : 1 and 3 : 1 I/II mixtures were studied by 1D \(^1\)H and 2D \([\text{H},\text{H}]\) TOCSY NMR experiments. The 1 : 1 I/II mixture was studied by 1D \(^1\)H, 2D \(^1\)N-decoupled \([\text{H},\text{H}]\) ROESY, NOESY, \(^{15}\text{N}-\text{edited}\) \([\text{H},\text{H}]\) TOCSY and NOESY, 2D \(^1\)N-decoupled \([\text{H},\ ^{15}\text{N}]\) TOCSY and HSQC NMR experiments using \(^{15}\text{N}-\text{en} labelled I. A near-complete NMR spectral assignment of the NOESY NMR spectrum of the 1 : 1 I/II mixture was achieved, although its complexity precluded full structure determination. Assignments were made possible by the known selectivity of I for guanines, and the localization of structural perturbations to residues close to the ruthenated G residue. Thus, sequential assignments along each strand always led to cross-peaks largely identical to those of the non-ruthenated duplex. Despite extensive overlap of NOE cross-peaks, little ambiguity in the assignments of individual resonances was found, with cross-validation of signal assignments from related connectivities being possible, with the help of \([\text{H},\ ^{15}\text{P}]\) HSQC, \([\text{H},\ ^{15}\text{N}]\) HSQC and \(^{15}\text{N}-\text{edited}\) \([\text{H},\ ^{15}\text{H}]\) NOESY.

![Diagram](Image 311x164 to 547x429)

**Scheme 1** (A) Reaction of single-stranded (ss) hexamer I (0.1 mM) with 1 mol equiv. of I in H\(_2\)O, 310 K for 48 h, gives three mono-ruthenated I (I-Ru-G\(_2\), I-Ru-G\(_3\), I-Ru-G\(_3\)). (B) Reaction of double-stranded (ds) hexamer II (0.3 mM, 0.1 M NaClO\(_4\)) with 1.1 mol equiv. of I in 90% H\(_2\)O/10% D\(_2\)O gives rise to two mono-ruthenated duplexes II-Ru-G\(_2\) and II-Ru-G\(_3\) as products. Addition of a second mol equiv. of I results in di-ruthenated duplexes, including II-Ru-G\(_2\)-G\(_2\), II-Ru-G\(_2\)-G\(_3\) and II-Ru-G\(_3\)-G\(_2\) as main products. Addition of a third mol equiv. of I results in two tri-ruthenated duplexes II-Ru-G\(_2\)-G\(_2\)-G\(_2\) and II-Ru-G\(_2\)-G\(_3\)-G\(_2\) as main products. Ru = \([(\eta^5\text{-tha})\text{Ru(en)}]\)\(^+\), I. For structure of I, see Fig. 1.
experiments and HPLC-MS data. 2D $\text{^15}$N-decoupled $[^1\text{H}, \text{^1}\text{H}]$ ROESY and NOESY and $\text{^15}$N-decoupled $[^1\text{H}, \text{^1}\text{H}]$ HSQC NMR data were also acquired for 2 : 1 and 3 : 1 I/II mixtures, but the spectra were too complex for interpretation.

HPLC and ESI-MS characterization of products from ss-DNA I + 1

A 100 µM aqueous solution of I was incubated with I at 310 K at Ru : I molar ratios of 1 : 1, 2 : 1 and 3 : 1 for 48 h in the dark, and these were then analyzed by HPLC. The low ionic strength (5.1 × 10$^{-4}$ M) ensures that this self-complementary oligonucleotide remains largely single-stranded (calculated melting temperature 264 K) under these conditions.

New peaks were observed for each reaction (Fig. 2 and Table S1†), and the adducts associated with them were identified subsequently by ESI-MS. The peaks for the observed negative ions are listed in Table S1†. Reaction at a Ru : I molar ratio of 1 : 1 resulted in three mono-ruthenated products together with three di-ruthenated products. Reaction at a Ru : I molar ratio of 2 : 1 resulted in three di-ruthenated products together with a tri-ruthenated product. Reaction at a Ru : I molar ratio of 3 : 1, gave only one main HPLC peak corresponding to a tri-ruthenated product.

HPLC and ESI-MS characterization of products from reaction of duplex II + 1

An aqueous solution of I was incubated with duplex II in 0.1 M NaClO$_4$ at ambient temperature for 24 h at a Ru : II molar ratio of 1 : 1 in an NMR tube in the dark. This gave rise to HPLC peaks which were identified by ESI-MS as ss-DNA I and two mono-ruthenated single-stranded products (see Fig. 2(d), Table S1†), with relative peak area ratios of 2 : 1. Another equimolar amount of I was then added to give a Ru : II molar ratio of 2 : 1, and was kept at ambient temperature in the dark for 48 h. This gave HPLC peaks which were identified by ESI-MS as two mono-ruthenated and two di-ruthenated single-stranded products (see Fig. 2(e) and Table S1†).

NMR characterization of products

Assignments of the $[^1\text{H}]$ NMR peaks for the ruthenated DNA duplexes were made on the basis of established methods developed for studying right-handed B-DNA duplexes by NMR spectroscopy.

The assignments of the $[^1\text{H}]$ NMR resonances of free DNA duplex II have been reported by Lam and Au-Yeung and the $[^1\text{H}]$ and $[^3]\text{P}$ chemical shifts are listed in Table S2†. The resonance assignments of backbone $[^3]\text{P}$ and sugar ring $[^1\text{H}]$ (H4′ and H3′) resonances for free and mono-ruthenated DNA duplexes were achieved by $[^1\text{H}, ^[3]\text{P}]$ HSQC NMR experiments and the H4′ and H3′ proton resonances were assigned by correlation to their respective $[^3]\text{P}$ resonance. The assignments of ruthenated G*H8 and I′-en-NH2 $[^1\text{H}]$ (NHu and NHd, see Fig. 1 for labelling) resonances of mono-ruthenated DNA duplexes were achieved by reference to 2D $[^1\text{H}, ^[15]\text{N}]$ HSQC and $[^15]$N-edited $[^1\text{H}, ^[1\text{H}]$ NOESY NMR data. The NMR chemical shifts of the $[^1\text{H}]$ and $[^3]\text{P}$ resonances associated with these two mono-ruthenated duplex adducts are listed in Table S3 (II-Ru-G′), Table S4 (II-Ru-G′) and Table S2 (free II). The assignments of H1,2, H2,3, H9,10, H5,6, H6,7 and I'-en-NH2 (NHu and NHd, see Fig. 1 for labelling) $[^1\text{H}]$ NMR resonances of $[^1\text{H}]$ (I′)-en-NH2) $[^1\text{H}]$ (I′) were achieved by 2D $[^1\text{H}, ^[15]\text{N}]$ HSQC, $[^15]$N-edited $[^1\text{H}, ^[1\text{H}]$ NOESY and $[^1\text{H}, ^[1\text{H}]$ NOESY experiments and are listed in Table I and Fig. 5. The assignments of H2,3 and H1,4 NMR resonances of I′ were achieved by correlations to the I′-en-NH2 resonances in $[^1\text{H}, ^[1\text{H}]$ NOESY NMR data, where the H9,10 protons were assigned by correlation to the H1,4 resonances, and H5,8 protons by correlation to the H9,10 and H6,7 resonances.

NMR of 1.1 : 1, 2 : 1 and 3 : 1 I/II reactions

Fig. S1 shows the imino and aromatic region of the 800 MHz 1D $[^1\text{H}]$ NMR spectrum of DNA duplex II in the absence (Fig. S1A†) and presence of 1 mol (Fig. S1B†), 2 mol (Fig. S1C†) and 3 mol (Fig. S1D†) equiv. of I. Reaction with 1.1 mol equiv. of I resulted in the formation of a number of new peaks for II (especially near 8.5 ppm (G*H8), 13.0–13.6 ppm (imino) and

![Fig. 2](image-url)

Fig. 2  HPLC chromatograms for reaction of $[(\eta^5\text{-tha})\text{Ru(en)}\text{Cl}]$ (I) with single-stranded (ss) d(CGGCCG) (I) (0.1 mM in H$_2$O) at Ru : I mol ratios of (a) 1 : 1, (b) 2 : 1, and (c) 3 : 1, and for reaction of I with duplex d(CGGCCCG)$_2$ (II) (0.34 mM, 0.1 M NaClO$_4$, 90% H$_2$O/10% D$_2$O) at a Ru : II mol ratio of (d) 1.1 : 1 and (e) 2 : 1. The mono-ruthenated duplex II-Ru$_3$ elutes as mono-ruthenated ss-DNA I (I-Ru-G′ and I-Ru-G′; species I-Ru-G′ and I-Ru-G′ are identical, as are I-Ru-G′ and I-Ru-G′); di-ruthenated duplex II-Ru$_2$ elutes as mono-ruthenated ss-DNA I (I-Ru-G′ and I-Ru-G′, see (d), and di-ruthenated ss-DNA I (I-Ru, see (e)). It is notable that G′ is readily ruthenated for single strand I (see I-Ru-G′ in (a)) but not for duplex II in (d). Little ruthenation on G′ was observed when 1 mol equiv. of I was added to mono-ruthenated duplexes II-Ru-G′ and II-Ru-G′ (e). Ru = ($[^1\text{H}]$-tha)-Ru(en)$[^{-2}]$ (I′), and is bound to G′N7 or G′N7; for DNA sequence, see Fig. 1 and Scheme 1.
6.3–6.7 ppm (NHu-I', I' is the bound complex), \{[(\eta^4-tha)-Ru(en)]^2+\}, Fig. S1B†). Two imino \(^1\)H NMR resonances were shifted to low field by \(+0.04\) ppm (G\(^*$\), mono-ruthenated G\(^*$\) base) and \(+0.14\) ppm (G\(^*\)), and two imino \(^1\)H resonances were shifted to high field by \(-0.07\) ppm (G\(^*$\), mono-ruthenated G\(^*$\) base) and \(-0.04\) ppm (G\(^0\)), relative to the free duplex II (Figs. S1 and S2, Tables S3, S4 and S2†). Reaction of the second mol equiv. of I with the mono-ruthenated duplexes resulted in a notable increase in intensities of the new peaks, especially G\(^*$\)H8 (near 8.5 ppm), NHu-I' (6.3–6.7 ppm), H5 and H1' resonances; the intensities of imino and H8, H6 resonances of free II all decreased (Fig. S1C†). Reaction of the third mol equiv. of I with the di-ruthenated duplexes resulted in an increase in the intensities of new peaks, e.g. at 8.8 ppm (Fig. S1D†). The resonances of CH5, CH6, H1' and G\(^*$\)H8 moved to low field by up to +0.3 ppm, and the peaks for imino protons almost disappeared.

**2D \(^{1}\)H, \(^1\)H TOCSY NMR of 1.1 : 1, 2 : 1 and 3 : 1 I/II reactions**

The 2D TOCSY NMR spectrum of the 1.1 : 1 I/II reaction mixture clearly showed the existence of cross-peaks for the two mono-ruthenated duplexes, as seen for example in the aromatic region in Fig. 3B. Two sets of H5-H6 cross-peaks were detected for C\(^4\), C\(^6\), C\(^7\) and C\(^10\) residues. The proportions of Ru-IHa and Ru-Ib at 283 K were determined by integration of the TOCSY cross-peak volumes of C\(^5\)-H5/C\(^6\)-H6 of Ru-IHa, and C\(^5\)-H5/C\(^6\)-H6 of Ru-Ib, and the HPLC peak areas for I-Ru-G\(^*$\) and I-Ru-G\(^0\) (see Fig. 2d). This gave a Ru-IHa : Ru-Ib ratio of 2 : 1 (±10%). Other species account for less than 10% of the total DNA. The 2D TOCSY NMR spectrum of the 2 : 1 I/II reaction mixture shows that peaks for other new species are present but not all can be assigned due to the complexity of the spectrum (Fig. 3C). It was notable that the intensities of the CH5-CH6 cross-peaks for C\(^5\)/C\(^11\) residues of free II decreased remarkably, but those of the CH5-CH6 cross-peaks for C\(^v\), C\(^u\) and C\(^w\) residues of the ruthenated species increased markedly. The 2D TOCSY NMR spectrum of the 3 : 1 I/II reaction mixture shows that the CH5-CH6 cross-peaks for C\(^5\)/C\(^11\) residues of free II almost completely disappeared (Fig. 3D).

**2D \(^{1}\)H, \(^{15}\)N HSQC NMR of the 1.1 : 1 I/II reaction**

These experiments allowed detection of NMR peaks specifically for the \{[(\eta^4-tha)-Ru(en)(\(^{15}\)N-en)]^2+\} fragment. These are commonly difficult to resolve in normal \(^1\)H NMR experiments. One major new species was detected by 2D \(^{1}\)H, \(^{15}\)N HSQC NMR analysis of the 1.1 : 1 mixture of duplex II and \(^{15}\)N-I (the \(^{15}\)N-en labelled complex I) at 283 K in 90% H\(_2\)O/10% D\(_2\)O (Fig. S3†). Peaks were assignable to en-NH\(_u\) resonances (the NH protons oriented...
towards the coordinated arene ring, see Fig. 1 and Table S5†) of mono-ruthenated duplexes Ru-IIa and Ru-IIb (Ru = \{(η⁶-tha)Ru(en)}²⁺ (I²)). No cross-peaks for en-NHu resonances of Ru-IIa and Ru-IIb were detectable after the equilibrium mixture had been freeze-dried and re-dissolved in D₂O at 283 K. The en-NHd resonances of both Ru-IIa and Ru-IIb were not observed in either H₂O or D₂O solutions. In contrast, the en-NHu and en-NHd resonances of unreacted I were detected in 90% H₂O (Fig. S3). The assignments are listed in Table S5†.

2D [¹H, ³¹P] HSQC NMR of 1.1 : 1 I/II reaction

The backbone phosphate ³¹P (−0.6 to −1.4 ppm) to sugar ring H₃' (5.3–4.6 ppm) and H₄' (4.6–4.0 ppm) HSQC connectivities for free duplex II and the 1.1 : 1 I/II reaction are shown in Fig. 4 and the assignments are listed in Tables S3 and S4.† Compared to free duplex II, the ³¹P/H₄' cross-peaks for C₄ (peak e) and G₈* (peak i) residues, ³¹P₆⁺/H₃'₉₈ cross-peaks for G₂⁻G₄* (peak d), G₈*-C₄ (peak f) and C₄-C₆ (peak h) residues were shifted to give new peaks, but ³¹P/H₄' cross-peaks for G₈*, C₁₀ and C₈ (peak g) and ³¹P₆⁺/H₃'₉₈ cross-peaks for C₄-G₄* (peak j) residues were too broad to assign. Decreased intensities of ³¹P/H₄' and ³¹P₆⁺/H₃'₉₈ cross-peaks were found for G₈/G₉* (peak b), G₉/G₈* (peak c), and C₄/C₉* (peak d) residues. These results are consistent with the HPLC-MS and 2D TOCSY NMR data.

2D [¹H, ¹⁵N]-edited [¹H, ¹⁵N] NOESY NMR of products from 1.1 : 1 I/II reaction

Assignments for ¹H NMR peaks of mono-ruthenated duplexes Ru-IIa and Ru-IIb in the spectra of the 1.1 : 1 I/II reaction are listed in Tables 1, S3 and S4,† and intermolecular NOEs in Tables S6 and S7.† For Ru-IIa, a large low-field shift of the G₉/H₈ resonance was observed, as was also the case for H₈ of the neighbouring G² base and H₅ and H₆ of the neighbouring C⁴ base, relative to free duplex II (Fig. S4 and Tables S2 and S3†). The largest changes in deoxyribose H¹' chemical shifts occur for G¹*, G³ and C⁶ residues, with the smallest changes for the neighbouring C⁴ and C¹⁰ residues (Fig. S4 and Table S3†). NOE cross-peaks were found between G²⁻H₈ and I'-en-NHd, I'-en-NHu, H₂.3, H₁.4 and H₉.10, protons, between G¹*-H¹', G⁹⁻H²'/ H₄' and I'-H₉.10 and H₁.4, protons, between C⁴⁻H⁵, C₆⁻H₆ and I'-H₉.10 and H₁.4, protons, and between C⁶⁻H¹' and I'-H₉.10, H₅.8 protons (Figs. 5, S6 and Table S6†). NOE cross-peaks were also found between protons of bases G², G³ and C¹⁰ and I'. In particular, NOE cross-peaks were observed between G'H²*, C₉(o), H¹', I' and H₆.7 (Figs. 5, S6 and Table S6†). Sequential connectivities for base-to-sugar ¹H NMR resonances were obtained, but those in the G²-C⁴*, G₈*-C₄ and G₉⁻C₉ steps were extremely weak or absent. The interruption or weakening of NOE connectivities between sequential DNA nucleotides is consistent with the binding of \{(η⁶-tha)Ru(en)}²⁺ (I²) at G₈* in the adduct Ru-IIa.

For adduct Ru-IIb, large low-field shifts were observed for the G₉/H₈ resonance and for H₅ and H₆ resonances of the neighbouring C⁴ residue (Fig. S5 and Tables S2 and S4†). The H₆ resonance of C⁴ in the complementary strand, which is paired with G², shifted slightly to low field, but the H₅ resonance shifted to high-field relative to free duplex II. The largest changes in H¹' chemical shifts were found for G¹*, and for C¹, C₃, and G₈*. NOE cross-peaks were found between G₉⁻H₈ and I'-en-NHd, I'-en-NHu, H₂.3, H₁.4, H₉.10 and H₅.8, protons, between G¹*-H¹', and I'-H₉.10, H₅.8 and H₆.7, protons, and between G¹*⁻H₄', G₈*⁻H₅* and I'-H₉.10 and H₅.8 protons (Figs. 5, S6 and Table S7†). NOE cross-peaks were also detected between protons of the bases C⁴, G₉, C¹₀ and bound fragment I' (Figs. 5, S6 and Table S7†). Particularly of note were cross-peaks observed between C⁴⁻H²', C₉-o⁻H₆.7 and I'-H₅.8. Sequential base-to-sugar connectivities were obtained, but those in the C₄-C⁴*, C₉⁻C₈ and C₉⁻C₉ steps were extremely weak or absent. The interruption or weakening of NOE connectivities between sequential DNA nucleotides is consistent with the binding of I' at G₈* in the adduct Ru-IIb.

Only one set of signals was observed for the bound fragment \{(η⁶-tha)Ru(en)}²⁺ (I²) in the two ruthenated duplexes Ru-IIa and Ru-IIb (Figs. 5, S6† and Table 1). Compared to the unbound chloro form of I, peaks for I'-H₁/H₄ and H₂/H₃ of the coordinated arene (see Fig. 1 for labelling) were shifted to low-field, the largest shift being for I'-H₁/H₄ (Table 1). Peaks for I'-H₉/H₁₀, H₅/H₈ and H₆/H₇ of the non-coordinated rings were shifted to low-field by +1.00, +0.12 and +0.09 ppm, respectively, the largest shift being for I'-H₉/H₁₀ (Δ δ = +1.00 ppm). Two sets of slightly
low-field-shifted or unchanged signals for I'-en CH$_2$ of both Ru-IIa and Ru-IIb were detected. Two sets of signals for both I'-en-NHd and en-NHu protons of Ru-IIa and Ru-IIb were observed, and the peaks for I'-NHd and NHu were shifted to low-field, the largest shift being for I'-NHu. One set of signals from unreacted ruthenium complex I was observed in the 1.1 : 1 I/II reaction mixture (Fig. S7† and Table 1). These results are consistent with ROESY experiments (data not shown).

**Discussion**

Complex I selectively ruthenates guanine bases in single strand DNA I or duplex II (Fig. 2a–c) with a similar pattern to that observed for the biphenyl (bip) and p-cymene (cym) complexes. Complex I is as reactive towards duplex II as the Ru-bip complex, and much more reactive than the Ru-cym complex. This pattern of reactivity was observed previously with calf thymus DNA, for which $t_{50\%}$ values of 10 min, 10 min and 3.5 h for complexes I, Ru-bip and Ru-cym, respectively, were found. Precipitation of adducts was observed when >1 mol equiv. of Ru-bip or Ru-cym complex was added to duplex II (0.2 mM). However, no such behaviour was observed in the present work. Addition of up to 3 mol equiv. of complex I to duplex II, even at the higher concentration of 0.3 mM, did not result in precipitation. This suggests that the nature of the arene influences intermolecular interactions. However, precipitation of adducts was observed when >3 mol equiv. of complex I was added to duplex II (0.3 mM), and was also the case when the reaction mixture of I + II (3 : 1) was kept at 277 K for long periods (ca. four weeks).

Intermolecular interactions are probably also influenced by the order of occupation of the Ru sites and the extent of arene intercalation (for tha and bip).

**Determination of binding sites by NMR**

The $^{31}$P chemical shift changes determined from the 2D [$^1$H, $^{31}$P] HSQC NMR experiment are consistent with ruthenation at N7 of the G residues of the 6-mer DNA duplex II by I. Binding of Ru-bip to the phosphate of 5'-GMP causes a low-field shift of the $^{31}$P NMR resonance by up to +5.11 ppm. However, the binding of Ru-bip to N7 of 5'-GMP, 5'-IMP or 5'-cGMP caused low-field $^{31}$P NMR shifts of less than 1 ppm. Similarly, ruthenation of 5'-GMP by trans-[RuCl$_3$(DMSO)$_4$] giving Ru-OPO$_3$ coordination causes a +5.8 ppm $^{31}$P downfield shift, and direct Pt-OPO$_3$ binding to IMP produces a $^{31}$P downfield shift of about +3.5 ppm. The formation of N7-ruthenated complexes of 5'-GMP and 5'-IMP, N6- or N4-ruthenated complexes of 5'-AMP or 5'-cAMP by [Ru(ni)(NH$_3$)$_3$]$^{13+}$ gave rise to little change in $^{31}$P resonances of the nucleotides. Therefore, it is evident that direct coordination of Ru$^{II}$ to a phosphate oxygen induces a $^{31}$P chemical shift change of ca. +5 ppm, while coordination to GN7 and no direct binding to phosphate oxygen induces a chemical shift change in the range of 0–1 ppm. In the present case, the most affected signals are assigned to the phosphate groups of residue C$^1$ ($\Delta\delta$ +0.09 ppm) and residue G$^7$ ($\Delta\delta$ –0.05 ppm) (Fig. 4 and Tables S3 and S4). Other $^{31}$P resonances are shifted by less than 0.03 ppm. Significant changes were observed for $^{31}$P, H$^3$ and H$^4$ resonances of residues G$^{3\alpha}$, C$^\beta$, C$^\delta$ and G$^{4\beta}$; minor
changes observed for the corresponding resonances of C'\text{/}C" and G'\text{/}G" further indicated that the binding site was N7 of G* or G*. Selective binding to N7 of the G residues of double-stranded DNA duplex II was evident from the \textsuperscript{15}N-edited [\textsuperscript{H}, \textsuperscript{H}] NOESY NMR spectrum and confirmed by H NMR chemical shift changes (Figs. 5, S6 and Tables S3 and S5†). NOE connectivities between ruthenated G*H8 and I'-NHu or I'-NHz were observed in the \textsuperscript{15}N-edited [\textsuperscript{H}, \textsuperscript{H}] NOESY and \textsuperscript{15}N-decoupled [\textsuperscript{H}, \textsuperscript{H}] NOESY NMR spectra. Binding of I to 9-ethylguanine and 5'-GMP\textsuperscript{7} via N7 causes a low-field shift of the H8 NH NMR resonance by up to +0.6 ppm. Binding of Ru-cym and Ru-bip complexes to 6-mer single strand DNA I or duplex II\textsuperscript{11} via N7 causes low-field shifts of the H8 NH NMR resonance of +0.49 to +0.66 ppm, and +0.28 to +0.58 ppm, respectively. Similar shifts were observed for the H8 resonances of G bases in the hexamer, and allow assignment of the binding sites as G* (Δδ H8 + 0.59 ppm) in Ru-IIa and G* (Δδ H8 + 0.46 ppm) in Ru-IIb present in the 1.1 : 1 reaction mixture of I + II (Figs. S1, S4 and S5, Tables S3 and S4†). With the binding fragment ([η\textsuperscript{5}-thia]-Ru(en)\textsuperscript{2} (I'), the mono-ruthenated duplex Ru-IIa is assigned as II-Ru-G'\textsuperscript{1}(I), Ru-IIb as II-Ru-G'\textsuperscript{1}(I) (for DNA sequence, see Scheme 1).

The ruthenation of duplex II by complex I mainly caused low-field shifts of imino proton resonances of G residues G* and G' in II-Ru-G'\textsuperscript{1}(I), but high field shifts of imino proton resonances of G residues G* and G' in II-Ru-G'\textsuperscript{1}(I) (Fig. S2 and Tables S3 and S4†). In contrast, the imino proton resonances of the mono-intercalated duplexes and di-intercalated duplex ruthenated with Ru-bip are broad and weak, implying that the base-pairs are disrupted in the duplex with an increase in dynamic mobility of the bases.\textsuperscript{23} High-field shifts of imino proton resonances were found for mono-ruthenated species in the 1 : 1 reaction mixture of Ru-cym complex + II, and platination of the 14-mer duplex d(TATGTACCATGTAT/d(ATACATGTTACATA) also causes high field shifts of G imino proton resonances.\textsuperscript{25,59}

Structural perturbations induced by ruthenation with complex I are larger than those observed for Ru-bip and Ru-cym complexes,\textsuperscript{23} and are localized to within a few (±2) base-pairs of the ruthenation site in all cases for complex I, while only the two adjacent bases (C' and C'\textquoteright or C" and C") are affected by ruthenation at G* or G** in all cases for Ru-bip and Ru-cym adducts. Not only were low field shifts of the H5 and H6 resonances observed for C' in II-Ru-G'\textsuperscript{1}(I) and C" in II-Ru-G'\textsuperscript{1}(I), but also for H1' of G', G', C', C" and C" in II-Ru-G'\textsuperscript{1}(I), and of C', C" and G" in II-Ru-G'\textsuperscript{1}(I) (Tables S3 and S4†).

**Intercalation**

Literature reports show that intercalation into DNA base pairs can often be recognised by distinctive features,\textsuperscript{19,22,23,49,54,61,65} including (a) upfield 1H NMR shifts of resonances of the intercalator; (b) NOE cross-peaks between protons of the intercalator and DNA bases at sites of intercalation; (c) the interruption or weakening of NOE connectivities between sequential DNA nucleotides; (d) the absence or weakening of the correlation peaks of H3\textsubscript{p} -2J\textsubscript{p}p\textsubscript{n+1} and H3\textsubscript{p} -2J\textsubscript{p}p\textsubscript{n} at sites of intercalation steps and the large chemical shift perturbations at the intercalation steps; and (e) the weakening of the strength of H-bonding between en-NH and GO6 in the case when the ruthenium complex Ru-bip with an extended arene ring system was involved.\textsuperscript{22,23}

It is notable that no large high-field shifts of proton resonances of I' were detected, but large low-field shifts up to +1.00 ppm were observed for protons H9,10, H5,8 and H6,7 of rings B and C in the mono-ruthenated duplexes II-Ru-G'\textsuperscript{(I)} and II-Ru-G'\textsuperscript{(I)} (Table 1). These shifts are inconsistent with shielding effects from the ring-currents of nucleobases which form a sandwich with the intercalated non-coordinated rings of bound I', and so do not provide evidence for intercalative binding.\textsuperscript{19,49,54,65} For example, upfield shifts of between −0.4 and −1.0 ppm have been reported for Ru-bip intercalated into 6-mer or 14-mer duplex DNA,\textsuperscript{22,23} and upfield shifts of −0.1 to −1.0 ppm for the intercalated dap (1,12-diazaperylene) ligand of the dirhodium(II) carboxylate complex [Rh\textsubscript{2}(dap)(CH\textsubscript{3}COO)\textsubscript{3}(CH\textsubscript{2}OH)\textsubscript{4}] into a 12-mer duplex DNA.\textsuperscript{29} Such large low-field shifts of the bulky thia intercalator have not been observed for other bulky intercalators, for example, large high-field shifts have been observed for bulky intercalated cholesterol groups.\textsuperscript{44} However, similar large low-field shifts for proton resonances of I'-9EtG were found for the adduct [(η\textsuperscript{5}-thia)Ru(en)(9EtG)],\textsuperscript{28} the H5,8 and H6,7 resonances slightly shifted to high-field, but the H9,10 resonances remained unchanged. In the case of [(η\textsuperscript{5}-thia)Ru(en)(5'-GMP)] (I'-GMP),\textsuperscript{22,23} H9,10 and H5,8 resonances are shifted to low-field by +0.83 and +0.50 ppm, respectively, and the H6,7 resonances shifted to high-field by −0.10 ppm. In the present case of the mono-ruthenated duplexes II-Ru-G'\textsuperscript{(I)} and II-Ru-G'\textsuperscript{(I)}, H9,10, H5,8 and H6,7 resonances shifted to low-field by +1.00, +0.12 and +0.09 ppm, respectively. Thus it is reasonable that the resonances of intercalated non-aromatic rings B and C of thia in the mono-ruthenated duplexes II-Ru-G'\textsuperscript{(I)} and II-Ru-G'\textsuperscript{(I)} shift to low field.

The single crystal X-ray structure of (I'-GMP)\textsuperscript{28} shows that ring C of I' is tilted towards the purine by 27.8° and lies directly over the purine base, indicative of strong intramolecular π–π stacking between ring C and the purine ring with a centroid–centroid separation of 3.45 Å and dihedral angle of 3.3°. Intercalation of the non-coordinated rings of I’ into the DNA duplex is also consistent with circular and linear dichroism data.\textsuperscript{26,29} Due to excessive resonance broadening, the resonances for protons that intercalate between purine rings are difficult to assign. Weak to intermediate intensity NOE cross-peaks were found not only between the rings of bound I' and H1' or H8 protons of G* or C' in II-Ru-G'\textsuperscript{(I)}, but also between the rings of I' and G" (Figs. 5, S6 and Table S6†). This can happen if the intercalation occurs not only at the GpC\textsuperscript{4} base step, but also at the GpC\textsuperscript{10} base step. The intermediate intensity cross-peaks observed between G'H2' or C"H1' and I'-H6,7 protons, indicate that the extended rings of I' intercalate deeply and are located between the middle of G" and C\textsuperscript{10} bases. Analogous NOE cross-peaks between the rings of bound I' and H1', H8, H2' and H2' of G* in II-Ru-G'\textsuperscript{(I)}, and also between rings of I' and C', C" and G" were detected, indicating that intercalation occurs between G* and C", and between G' and C' as well (Figs. 5 and S6, and Table S7†). The intermediate intensity cross-peaks observed between G" or C' and I'-H1,4 protons, indicate that the coordinated arene ring of I' is located between the middle of G" and C" bases. Additionally, intermediate intensity cross-peaks observed between C'H2' and I'-H6,7 and C'H2" and I'-H5,8 protons indicate that the extended rings of
I’ intercalate deeply and are located near to the C’ base. The interruption of NOE connectivity pathways between the corresponding base pairs (G’-C’*, G*-C’ and G’-C’0 steps in II-Ru-G’(I’), and C*-G’, C’-G’* and C’-G’ steps II-Ru-G’(I’)) is consistent with these intercalation sites.

The absence of the H3’/Pn+1 cross-peaks linking the C’-G’** step and the H3’-P cross-peaks of G’*, C’ and C’, the low field shifts for H3’/Pn+1 cross-peaks linking G’-G’*, C’-C’ and G’*-C’, and for H3’-P cross-peaks of C’/C’0 and G’, and the large chemical shift perturbations at the G’-G’* and C’-G’** steps, together indicate that the intercalation occurs between G’pC’ or C’pG’ base steps (Fig. 4). Previous work has shown that the intercalation sites of the non-coordinated phenyl ring of Ru-bip in mono-ruthenated duplexes** are also between G’pC’ or C’pG’ base steps.

No cross-peaks for en-NHu resonances of Ru-Ha and Ru-IIb were detected after the 1.1 : 1 I/II reaction mixture had been freeze-dried and re-dissolved in D2O. This suggests that the hydrogen bond between G’*O6 and en-NH of I’ is weakened (Fig. S3 and Table S5†), which is consistent with intercalation of the non-coordinated rings of I’ into duplex DNA II. Similarly weakened hydrogen bonds were also observed when the biphenyl ring of Ru-bip intercalates into the hexamer duplex.22 The strength of the H-bond between G’*O6 and en-NH is related to the decay rate of the en-NH signals when II-Ru adducts are dissolved in 99% D2O.22 For the non-intercalated adduct with Ru-cym, the half-life was 72 h. However, those of the mono- and di-intercalated Ru-bip adducts were only 5 h and <0.1 h, respectively.

NMR studies show that the arenene–nucleobase π−π stacking of I’ with hexamer duplex is different from that of Ru-bip.22 Only a very few weak NOE contacts between protons of ring B of biphenyl and H1’ and H2’/H2’+2 protons of G’+ or C’ of the hexamer duplex are observed.22 The protons Ho’, Hp’ and Hm’ of bound Ru-bip in the DNA duplex adducts were consistently shielded relative to free Ru-bip, consistent with base stacking of the non-coordinated ring B between base pair G’*pC’. However, in the present case, not only were weak to intermediate intensity NOE contacts detected between protons of rings B and C of tha and protons of G’+ and C’ or G’* and C’, but also intermediate intensity NOE contacts were detected between protons of ring C of tha and protons of G’ and C’0 or C’ and G’ DNA bases which pair with G’* or C’*. G’ or C’ in the complementary DNA strand (Fig. 5, Tables S6 and S7†), respectively. This indicates that rings B and C of tha are involved in a penetrative intercalation between two pairs of bases, G’/C’0;C’/G’ or G’/C’;C’/G’, respectively.

It is interesting that large low-field shifts, but not large high-field shifts are observed for the proton resonances of intercalated rings B and C of I’, indicating that the intercalative interactions between Ru-tha and Ru-bip with the DNA duplex are significantly different from one another. The ring current shifts are position-related: upfield shifts only arise when the protons are above or below the ring plane. In contrast, resonances for protons located close to the plane and beyond the confines of the ring are shifted to low field.66

Modelling of II-Ru-G’(I’) and II-Ru-G’(I’)

In order to attempt to rationalize the NOE and chemical shift information gathered for II-Ru-G’(I’) and II-Ru-G’(I’), two molecular models were constructed in which the duplex d(CGCGCG)2 was ruthenated at N7 of G’ or G’ by I’, as shown in Fig. 6. The II-Ru-G’(I’) (Fig. 6a–c) and II-Ru-G’(I’) (Fig. 6d–f) models are mostly consistent with the NMR observations, and overwhelming evidence indicates that a penetrative interaction occurs between rings B and C of I’ and duplex DNA at G’/C’0;C’/G’ or G’/C’;C’/G’, respectively. The model also provides a useful indication of how I’ may lie in relation to its DNA binding site. The H5,8 and H9,10 protons of I’ are located above or below the ring plane and the distances between H5,8 or H9,10 protons and purine or pyrimidine rings of G or C bases are nearly 0.9 A shorter than the case of aromatic intercalators, such as Ru-bip. Compared to the structural findings for aromatic intercalators, both structures (II-Ru-G’(I’) and II-Ru-G’(I’)) suggest distortion of bases-pair planes around the site at which the tha has penetrated and the system is likely to be highly dynamic in and around the intercalation sites. The dynamics of II-Ru-G’(I’), in which the Ru binds to a terminal, potentially frayed nucleotide, would be expected to be significantly different compared with those for II-Ru-G’(I’), in which the Ru binds to the internal base.

The data imply that in both cases the tha of I’ in the models has swung round so that the ring system points across to the opposite strand rather than penetrating so deeply into the strand to which the Ru centre is attached (G’* or G’*0). This supports the NOE contacts observed between protons of rings A and B of I’ and H1’ and H8 or H6 protons of G’, C’ or C’, G’ and in particular between H6,7 protons of ring C of I’ and the H1’, and H2’ protons of residue C’0, G’ or G’, C’ of the complementary.

Fig. 6 Molecular models of duplex II ruthenated at N7 of G’ or N7 of G’ with [(η6-tha)Ru(en)]2+. (a)–(c) II-Ru-G’ showing the intercalation of the tetrahydroanthracene ligand between G’/C’0;C’/G’. (d)–(f) II-Ru-G’ in which the non-arene rings are intercalated between G’/C’;C’/G’. In each case side and top views of the intercalation site are shown as well as the whole duplex (bottom). Colour code: tha green, Ru purple, P yellow, O red.
The data imply that in both cases the th of I’ in the models has swung round so that the ring system points across to the opposite strand rather than penetrating so deeply into the strand to which the Ru centre is attached (G’ or G”). This tendency is then likely to be stabilized by th–base interactions and C–H⋯X (X = O or N) hydrogen bonding,68 and the upfield shift induced by the intercalation effect on the protons located above or below the intercalator, due to the ring current effect of the aromatic groups. The H9,10 protons are located exactly central to the two strands of DNA, so the downfield shifts for H9,10 protons are larger (1.0 ppm).

The largest downfield shift (+1.00 ppm) is observed for H9,10 protons of ring B of I,266 | 2D NOESY and 2D NMR spectra (Table S5 and Figs. S5, S6), and the downfield shifts of NH resonances in the adducts are consistent with the presence of H-bonding to the C6 carbonyl of the coordinated G.22,23,27,28 Only one broad peak for NH(u) was detected in the 2D [1H, 13N] HSQC NMR data at 283 K; the resonance for NH(d) was too broad to observe (Fig. S3f). NH exchange was rapid for the en-NH(u) protons in the mono-intercalated duplexes II-Ru-G’(I) and II-Ru-G’(I) (Fig. 6). It is clear that H5,8 and H6,7 protons are located within the confines of the purine ring G in model II-Ru-G’(I) or G’ in II-Ru-G’(I), and the H9,10 protons are located exactly in the middle of the two strands (Fig. 6) and in the ring planes beyond the confines of the purine or pyrimidine rings.

It has been reported that proton chemical shifts may change by up to +2.1 ppm (downfield) on formation of C–H⋯X (X = O or N) hydrogen bonds.68 It is clear in the present case that the protons of rings B and C do sit near the edge of the purine or pyrimidine rings from the models II-Ru-G’(I) and II-Ru-G’(I), and the protons are not directed to the centres of these rings. Other than the fact that protons of ring B are within the arene ring plane in Ru-bip, the H5,8 and H9,10 protons of I’ are located above or below the ring plane in the models. Such orientation makes the distances between H5,8 or H9,10 and N or O atoms of purine or pyrimidine rings of G or C bases nearly 0.9 Å shorter than those of Ru-bip. For model II-Ru-G’(I): short H9,10⋯N1 of G (−2.68 Å), H9,10⋯N3 of C (−2.51 Å), H5,8⋯N1 or N7 of G (−2.70 or 2.75 Å, respectively), H5,8⋯OC2 of C (−2.60 Å), H6,7⋯sugar O of G (−2.58 or 2.80 Å, respectively) distances are observed (Fig. 6a–c). For model III-Ru-G’(I): short H9,10⋯N9 of G (−2.66 Å), H9,10⋯N3 of C (−2.54 Å), H5,8⋯N3 of G, C and G (−2.60, 2.61 or 2.61 Å, respectively), H6,7⋯OC2 of C (−2.59 Å) distances are observed (Fig. 6d–f). Thus, in the present case, the shifts of the protons of rings B and C of I’ reflect both the downfield shift induced by the formation of C–H⋯X (X = O or N) hydrogen bonding,68 and the upfield shift induced by the intercalation effect on the protons located above or below the intercalator, due to the ring current effect of the aromatic groups. The H9,10 protons are located exactly central to the two strands of DNA, so the downfield shifts for H9,10 protons are larger (1.0 ppm).

The data imply that in both cases the th of I’ in the models has swung round so that the ring system points across to the opposite strand rather than penetrating so deeply into the strand to which the Ru centre is attached (G’ or G”). This tendency is then likely to be stabilized by th–base interactions and C–H⋯X (X = O or N) hydrogen bonding within the G’-th ring-C’ and the G’-th ring-C’ ‘sandwich’ as shown in Fig. 6a–c, or within the G’-th ring-C’ and the G’-th ring-C’ ‘sandwich’ as shown in Fig. 6d–f. This kind of intercalation distorts the DNA more than that of aromatic intercalators, such as Ru-bip, and reduces the strength of H-bonding between en-NH and G’O6. The intercalation of the non-coordinated phenyl ring of Ru-bip,22 of actinomycin D (ActD) and daunomycin65 between the GC step in previous work suggested that steric crowding at the GpC step is less than that at the GpG step, which thereby allows accommodation of the bulky non-aromatic rings of th. A further driving force for GpC rather GpG intercalation is the weaker purine–pyrimidine π–π stacking interaction for GpC compared with purine–purine GpG steps.65 It is interesting in the present work that all of the intercalation of Ru-bound tetrahydroanthracene occurs between GpC base steps, and there is no evidence for intercalation at the GpG base steps.

Ru-en-NH⋯G06 H-bonding

Two distinct NHu-NHd cross-peaks were identifiable in the 283 K 2D NOESY and 2D 15N-edited NOESY NMR spectra (Table S5 and Figs. S5, S6), and the downfield shifts of NH resonances in the adducts are consistent with the presence of H-bonding to the C6 carbonyl of the coordinated G.22,23,27,28 Only one broad peak for NH(u) was detected in the 2D [1H, 13N] HSQC NMR data at 283 K; the resonance for NH(d) was too broad to observe (Fig. S3f). NH exchange was rapid for the en-NH(u) protons in the mono-intercalated duplexes II-Ru-G’(I) and II-Ru-G’(I) (no 1H/15N cross-peaks being detected for a D2O solution of II-Ru-G’(I) and II-Ru-G’(I), Fig. S3f). Such rapid NH exchange of the en-NH(u) protons was also observed in the double-intercalated duplex II-Ru2-G’G’(Ru-bip),22 suggesting that mono-intercalation of I’ into duplex II gives rise to similar effects to that of the di-intercalation of Ru-bip.

Sequence specificity of G metallation

Adducts of duplex II eluted as single strands from the reverse-phase HPLC column22,23 (Fig. 2d and 2e) due to the denaturing character of the HPLC solvent. Three mono-ruthenated products (I-Ru-G’; I-Ru-G’ and I-Ru-G’), and three di-ruthenated products (I-Ru2-G’G’; I-Ru2-G’G’ and I-Ru2-G’G’) were detected in the 1 : 1 I/I reaction mixture, showing that all three guanine bases can be ruthenated readily in the single-stranded
hexamer DNA d(CGGCCG) (Table S1†). Only the two mono-ruthenated products II-Ru-G‡ and II-Ru-G‡ were detected in the reaction of duplex II with I at a Ru : II molar ratio of 1 : 1; no G† ruthenated adduct was detected (Table S1†). When the above mono-ruthenated products II-Ru-G‡ and II-Ru-G‡ were reacted with a second mol equiv. of I at the same temperature, only two mono-ruthenated single strand adducts (I-Ru-G and I-Ru-G) and two di-ruthenated single strand adducts (likely to be I-Ru₂-G-G† and I-Ru₂-G-G†) eluted from the reverse-phase HPLC column (Table S1†). The products may therefore involve ruthenation on the same strand: II-Ru₂-G-G† and II-Ru₂-G-G†, or on different strands: II-Ru₂-G-G†, II-Ru₂-G-G*, II-Ru₂-G-G†, II-Ru₂-G-G‡ and II-Ru₂-G-G‡. No G† ruthenated duplex adducts such as II-Ru₂-G-G† or II-Ru₂-G-G‡ were detected. These results are consistent with 2D TOCSY experiments (Fig. 3): the cross-peaks intensities of C-H/5-H6 (C‘, C‘ and C‘) of ruthenated species increased in the 2 : 1 I/II reaction mixture. The cross-peaks for H5/H6 resonances of C5/C4 and C4/C5 of ruthenated species increased in the 2 : 1 I/II reaction mixture.

The cross-peaks for H5/H6 resonances of C5/C4 and C4/C5 of ruthenated species increased in the 2 : 1 I/II reaction mixture.

Bulky lesion and penetrative intercalation

Reported data show that the intercalative binding of a bulky intercalator cholesterol group on a modified base of a duplex DNA is a classical intercalation, and the lesion site and the distortions in the structure of the DNA produced by these cholesterol derivatives are somehow similar to those induced by other adducts containing polycyclic aromatic groups. However, the base opposite the modified nucleotide is displaced and the local structure of the double helix is highly distorted. These observations are quite different from those in the present work involving penetrative intercalation of the intercalator cholesterol group on a modified base of a duplex DNA when Ru-tha is coordinated at G(N7). It is notable that there is no displacement of the base opposite the modified nucleotide, but C–H⋯X (X = O or N) hydrogen bonding between protons of ring C of tha and O or N atoms of bases opposite the ruthenated nucleotides. The local structure of the ruthenated double helix is highly distorted. Such a highly distorted double helix is not observed in the DNA adducts of platinum complexes with an acridine side arm intercalator, e.g. PT-ACRAMTU, where the threading intercalation occurs at the central base-pair step but does not cause helical bending.

Experimental section

Materials

Organometallic ruthenium(II) complex [(η⁴-tha)Ru(en)Cl][PF₆] (IPF₆) (th = 1,4,9,10-tetrahydroanthracene, en = ethylenediamine) and 15N-labeled I (15N-I) were synthesised as described previously. The sodium salt of FPLC-purified oligonucleotide d(CGGCCG) I was purchased from Oswel (Southampton, UK) and was further purified by HPLC. Sodium perchlorate and acetonitrile (HPLC grade) were obtained from Fisher, and triethylammonium acetate buffer (TEAA) from Fluka.

High performance liquid chromatography (HPLC)

This was carried out on reversed-phase columns with TEAA and TEAA/acetonitrile as mobile phases.

HPLC-electrospray ionisation mass spectrometry (HPLC-ESIMS)

Negative-ion electrospray ionisation mass spectra were obtained on a mass spectrometer interfaced with a reversed phase HPLC column eluted with TEAA/acetonitrile gradients as above.
NMR spectroscopy

NMR data were acquired on an 800 MHz or 600 MHz Bruker Avance NMR spectrometer equipped with a multiple resonance TXI (1H, 13C, 15N, 31P) xyz-gradient probe.

Molecular modelling

A structure for canonical B-form duplex d(CGGCCG)2 was generated within the biopolymer module of Sybyl (version 6.3, Tripos Inc.). Crystal coordinates from the X-ray crystal structures of I allowed accurate representation of the Ru complex to be incorporated into the model Ru-DNA constructs. Docking of the Ru-complex onto G•N7 (x = 3 or 6) of the DNA structure was achieved by manual independent manipulation of both DNA and Ru-complex molecules. The Ru-N7 inter-atomic distance was based on reported crystal structures of Ru-GMP complexes. A pseudo-atom at the centre of the nπ-six-membered aromatic ring (ring A) of tha was attached to the Ru centre to provide a rotatable bond about which the tha moiety could be manipulated. In a similar way, the Ru–GN7 bond was activated to form a rotatable bond, about which the entire Ru ligand could be rotated independently of the DNA structure. Ru–G•N7 models were prepared in such a way as to reduce steric contact as far as possible. Constraints were applied where deemed plausible and structures were energy minimized to remove the effects of steric clash.

Details of reactions of II with 15N-I, HPLC, HPLC-ESI-MS, NMR and pH measurements are in the ESI.

Conclusions

In conclusion, the results presented here provide a rare example of coordinative binding and the penetrative intercalation of a bulky intercalator into DNA, and may help to explain why ruthenium arene complexes have a different mechanism of antitumour activity (perhaps related to recognition by nucleotide repair enzymes) compared to cisplatin. Firstly, the NMR results are indicative of the penetrative intercalation of the tha rings B and C of I, selectively between two base pairs GII/CIII:GIV/GV or GVI/CII:GIII/GIV, which contrasts with that observed between one base pair GII/CIV and or GII/CV for the classic aromatic intercalator Ru-bip. The two slightly different intercalation models for the extended non-aromatic rings of Ru-tha, indicate that the distortion of the DNA duplex is sequence-related. Secondly, large low-field shifts for proton resonances of the intercalated non-coordinated rings B and C of tha reflect both the downfield shift induced by the formation of short C–H···X (X = O or N) hydrogen bonds and upfield shift induced by the intercalation effect on the protons located above or below the intercalator, due to the ring currents of aromatic groups. The downfield shifts of H9,10 protons of I are larger (+1.0 ppm) for they are located exactly in the middle of the two strands. Such deshielding of intercalator NMR resonances is rare, indicating that the intercalative interactions between this bulky tha intercalator and classical aromatic DNA intercalators are somewhat different. Thirdly, the DNA structural perturbations induced by Ru-tha are larger than those observed for Ru-bip and Ru-cym complexes; distortions of base-pair planes are observed around the sites at which the tha has penetrated, and the dynamics of the terminal base ruthenated adduct II–Ru-G2(I') are significantly different from those of internal base ruthenated adduct II–Ru-G1(I'). These findings agree with the fact that the precipitation of DNA duplex adducts of Ru-tha is observed only at very high concentrations compared with Ru-bip, suggesting that the intercalation of sterically bulky tha into a DNA duplex makes the duplex DNA behave differently from intercalation by the aromatic bip. Fourthly, selective ruthenation at N7 of GII and GV in the hexamer DNA duplex is similar to that of Ru-cym and Ru-bip, but the mono-intercalation of tha reduced the strength of H-bonding between en-NH and GO6 as much as that for the di-intercalated di-ruthenated Ru-bip duplex. Intercalation at GpC by tha appears to have a lower energy penalty when compared with intercalation at GpG base steps, thereby allowing accommodation of the non-aromatic, bulky rings of tha. Although all 3 G's were readily ruthenated at N7 in the single-stranded DNA hexamer, only GII (or GIV) and GV, and not GIII (GVI) were ruthenated in the free DNA duplex which is attributed to unfavorable steric interactions between the duplex and arene for binding to GII (GV). The different ratios of II–Ru–GII: II–Ru–GIV adducts in the reaction mixtures with Ru:II ratio of 1:1, indicates that there are differences in specificity from binding to internal bases or terminal nucleotides for the non-intercalator Ru-cym, the aromatic intercalator Ru-bip and non-aromatic intercalator Ru-tha. Little ruthenation of GII was observed in the mono-ruthenated duplexes, but the favorable binding sites were GIV and GVII when di-ruthenated duplexes were reacted with (II–tha)Ru(en)2. These results also demonstrate that the combination of HPLC, ESI-MS together with 2D 1H, 1H TOCSY NMR experiments is powerful for elucidating the selectivity of G-base ruthenation of the free duplex II, mono-ruthenated duplexes and di-ruthenated duplexes. Such knowledge of DNA interactions may be incorporated into design concepts for this class of anticancer agents and assist the exploration of structure–activity relationships.

The coordinative and penetrative intercalative interactions between Ru-tha and duplex DNA are different from that of DNA modified covalently by aromatic or bulky intercalators, in which the displacement or flip-out of bases near the modified sites may occur. Although both involve the modification of a DNA base via coordinative bonding, the penetrative intercalative interactions between Ru-tha and duplex DNA are also different from that of platinum complexes with an acridine side arm intercalator, where the threading intercalation does not cause helical bending. The C–H···X (X = O or N) hydrogen bonds between protons of ring C of tha and O or N atoms of bases opposite the ruthenated nucleotides may contribute significantly to the intercalative interaction between Ru-tha and duplex DNA. The fact that penetrative intercalation has rarely been reported for mono-metallointercalators, implies that the direct Ru–N bonding may also assist with penetrative intercalation for the bulky tha ligand. Unwinding and distortion, while still maintaining the basic duplex structure, could contribute to the toxicity of the Ru-tha complex by hindering DNA repair. A bulky lesion is one of the six main DNA lesions that may invoke NER, for example, the first and rate-determining step in NER is the recognition of the bulky lesions by the XPC/HHR23B protein heterodimer complex. However, mutations and potentially cancer may result if the bulky lesions are resistant to NER. The
high anticancer activity both in vitro and in vivo and the high potency of the tha complex may arise in part from the lack of repair of the lesions formed on DNA by this complex, and assist with elucidation of structure–activity relationships for this class of complexes.

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Notes and references

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