Molecular Recognition of Metal Complexes by DNA: A Comparative Study of the Interactions of the Parent Complexes \([\text{PtCl(TERPY)}]\)Cl and \([\text{AuCl(TERPY)}]\)Cl₂ with Double Stranded DNA.

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ABSTRACT

The interactions of the parent complexes \([\text{AuCl(Terpy)}]\)Cl₂ and \([\text{PtCl(Terpy)}]\)Cl with DNA were analysed by various physicochemical methods. Surprisingly, these metal complexes produce different interaction patterns with DNA in spite of their profound structural similarity. Indeed, important modifications are detected in the characteristic UV-Vis bands of \([\text{PtCl(Terpy)}]\)Cl upon addition of ct-DNA, while the spectrum of \([\text{AuCl(Terpy)}]\)Cl₂ is almost unaffected. Gel electrophoresis studies confirm these findings: \([\text{PtCl(Terpy)}]\)Cl – but not \([\text{AuCl(Terpy)}]\)Cl₂ – retards significantly the mobility of the supercoiled form of the pHV14 plasmid after a short incubation time. Ultrafiltration studies indicate that the affinity of \([\text{PtCl(Terpy)}]\)Cl for ct-DNA is significantly greater than that of \([\text{AuCl(Terpy)}]\)Cl₂. On the other hand, both \([\text{AuCl(Terpy)}]\)Cl₂ and \([\text{PtCl(Terpy)}]\)Cl induce important changes in the CD spectrum of ct-DNA, at high concentration, and increase its Tₘ value. Remarkably, the analysed metal-complex/DNA interaction patterns depend critically on the incubation times. We propose that \([\text{PtCl(Terpy)}]\)Cl quickly intercalates DNA; then, formation of coordinative bonds progressively takes place with time. At variance, \([\text{AuCl(Terpy)}]\)Cl₂ first interacts electrostatically with the DNA surface, with subsequent slow formation of some coordinative bonds.

Keywords: Gold(III) complexes, platinum(II) complexes, calf thymus DNA (ct-DNA), pHV14 plasmid, circular dichroism, absorption spectroscopy, DNA melting, ultrafiltration

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INTRODUCTION

There is today great interest in understanding how DNA recognizes metal complexes, particular attention being devoted to non-covalent binding modes /1/. As previously suggested, metal complexes interact non-covalently with double helix DNA in three basic ways: external binding, groove binding and intercalation /2/. External binding refers to those interactions that occur on the DNA surface, mainly governed by electrostatic effects; groove binding is a stronger type of interaction that takes place when a molecule of proper size enters one of the grooves of DNA. Intercalation consists of the insertion of a flat aromatic molecule between two adjacent bases.

The interactions of various platinum terpyridine complexes with DNA have been extensively investigated in the last three decades, mainly by the groups of Lippard /3-7/, McMillin /8-10/, Romeo /11-13/ and Lowe /14,15/. Interest in these compounds is related, at least in part, to their promising pharmacological properties, specifically as potential antitumor and anti-tripanosomial agents. On the ground of the large body of experimental evidence collected so far it has been argued that square planar platinum terpyridine compounds bind DNA predominantly by intercalation /3-7/. The resulting duplex unwinding angle is comparable to that measured following DNA intercalation by ethidium bromide /16/. The [PtCl(Terpy)]Cl shows a binding constant for ct-DNA of about 1x10^5; remarkably this complex first intercalates DNA, then slowly forms covalent links to the nucleobases /10, 17/.

[AuCl(Terpy)]Cl₂, the gold(III) analogue of [PtCl(Terpy)]Cl, originally characterized by Lippard /18/, was recently reported by us to possess relevant cytotoxic properties /19/. By analogy with the case of [PtCl(Terpy)]Cl, [AuCl(Terpy)]Cl₂ may be believed to act as an intercalating agent. However the structural features of the binding of this complex to DNA are still controversial and crystallographic information is lacking due to intrinsic difficulties in obtaining crystals suitable for X-ray diffraction.

In the present paper we have further addressed this issue by investigating, comparatively, the interactions of the structurally related [PtCl(Terpy)]Cl and [AuCl(Terpy)]Cl₂ complexes (see Figure 1) with double stranded DNA.

The metal complexes/DNA interactions were analysed in solution, at physiological pH, by a variety of physicochemical techniques including spectrophotometry, circular dichroism, ultrafiltration, DNA melting and gel electrophoresis. The joint application of these methods offers a rather complete description of the reactions of either complex with the DNA double helix. To our surprise, despite their strict structural similarity, these two complexes turned out to produce largely different DNA interaction patterns.

EXPERIMENTAL

Materials

[AuCl(Terpy)]Cl₂ was synthesized as reported /20/. The purity of the compound was checked through elemental analysis and ¹H-NMR spectroscopy. Ct-DNA and [PtCl(Terpy)]Cl•2H₂O (98% purity) were purchased from the Sigma Chemical Company. All other reagents used in this study were of analytical grade.
Spectrophotometric measurements.

Electronic absorption spectra were measured by means of a Perkin Elmer Lambda 20 Bio instrument. The reaction of the gold(III) complexes with ct-DNA was first analysed through absorption spectroscopy, since each complex has a characteristic spectrum in the UV-Visible region.

Electronic spectra of [AuCl(Terpy)]Cl₂ 5 × 10⁻⁵ M and [PtCl(Terpy)]Cl 2 × 10⁻⁵ M were recorded before and after addition of ct-DNA (at r = 0.1, where r is the molar ratio between the added metal and DNA, expressed as base-pair) in the 50 mM phosphate, 4 mM NaCl, pH 7.4 buffer, at room temperature. The further evolution of the two systems over time was analysed until the spectral changes reached completion (data not shown).

In another set of experiments, visible absorption spectra were recorded at increasing complex/ct-DNA ratio (r = 0.0, 0.1, 0.2, 0.5, 1.0, 2.0 and 4.0) for [AuCl(Terpy)]Cl₂ and [PtCl(Terpy)]Cl immediately after mixing, in the previous buffer, at room temperature.

CD spectra

ct-DNA is characterized by two bands in the UV region: a positive peak at 260 nm and a negative one at 230 nm. We analysed the behaviour of the metal/DNA adducts at a fixed r during time, working in the 200-400 nm range. A ct-DNA solution in the reference buffer (50 mM phosphate and 4 mM NaCl, pH=7.4) was added to solutions containing [AuCl(Terpy)]Cl₂ 5 × 10⁻⁵ M (to obtain r = 0.2) and [PtCl(Terpy)]Cl 2 × 10⁻⁵ M (r = 0.08).

To establish in more detail whether binding of the complexes brings about any significant conformational change of the DNA double helix, CD spectra of ct-DNA were recorded at increasing complex/ct-DNA ratios.
The CD technique is indeed very sensitive to detect minor conformational changes of the DNA conformation produced by ligand binding.\(^{22}\)

CD spectra were recorded at increasing complex/ct-DNA ratio (r = 0.0, 0.1, 0.2, 0.5, and 1.0) for \([\text{AuCl(Terpy)}]\text{Cl}_2\) and \([\text{PtCl(Terpy)}]\text{Cl}\) immediately after mixing. A Jasco J600 dichrograph operating at room temperature was used, interfaced with a PC, and spectra (in the 200-400 nm range) were analysed through the standard Jasco software package. ct-DNA concentration was 1.25x10^4 M, 66 µg/ml in the reference buffer.

A second titration was carried out, with higher DNA concentration (2.5x10^4 M), to better analyse the "induced CD" bands, which appear in the presence of the metal complex.

**Ultrafiltration experiments**

1.25x10^4 M DNA was dissolved in the physiological buffer and \([\text{AuCl(Terpy)}]\text{Cl}_2\) and \([\text{PtCl(Terpy)}]\text{Cl}\) were added to obtain a r = 0.1 complex/ct-DNA ratio. The obtained samples were ultracentrifugated till half volume immediately after mixing at room temperature.

Ultrafiltration measurements were performed using tubes for concentration and purification of biological samples “Centricon Centrifugal Filters YM-10” by Amicon Bio-separation, Millipore Corporation, USA, with a Centurion centrifuge working at 3500 rpm for 30’ - 45’.

The spectra of the upper solution containing ct-DNA and the bound metal, and of the lower solution, containing only the metal complex, were recorded using electronic absorption spectroscopy.

Afterwards, the amount of free and of DNA-bound metal was determined, by the electronic spectra, and the affinities of the individual complexes for DNA were qualitatively estimated.

**DNA melting studies**

The effects of binding of the complexes on the stability of DNA double helix were investigated through analysis of the helix-to-coil transition. Melting experiments were carried out according to the reported procedure.\(^{23}\) Renaturation profiles, recorded by progressive cooling of the sample, allow establishment of the formation of interstrand crosslinks, which favour the re-association process.

All melting measurements were carried out in the 10^{-2} M NaClO_4 and 10^{-3} M NaCl buffer. Ct-DNA was dissolved in the buffer and DNA concentration determined by absorption measurements at 260 nm. DNA concentration was equal to 4.9x10^{-5} M [base pairs].

We recorded the melting curves (denaturation and renaturation) of the samples with r = 0.1 metal complex/DNA ratio.

Thermal denaturation experiments were performed in quartz cuvettes with a Perkin Elmer Lambda 20 Bio UV/Vis spectrophotometer equipped with a thermostated cell and a Peltier for temperature-control. Samples were continuously heated with a scan rate of 0.5°C/min while monitoring the absorbance changes at 260 nm. The investigated interval of temperature ranged from 45°C to 90°C. Upon reaching 90°C, samples were cooled back to 40°C in order to follow the renaturation process. Values for the melting temperatures (Tm) and the melting interval (ΔT) were determined according to the reported procedures.\(^{24}\).
Adducts with pHV14 plasmid DNA were prepared by adding the required volume of a freshly prepared solution of either [AuCl(Terpy)]Cl₂ or [PtCl(Terpy)]Cl in 50 mM sodium phosphate and 4 mM sodium chloride (pH 7.4) buffer. The concentration of pHV14 DNA in the reaction mixture was 75 ng/μl (6x10⁴M), while the concentration of the metal complexes varied to give different metal-to-bp stoichiometries (0.1, 0.2, 0.4 and 0.8). The mobility of the [AuCl(Terpy)]Cl₂ and [PtCl(Terpy)]Cl treated pHV14 samples was analyzed by gel electrophoresis /25/ on 0.8% (w/v) agarose gel (Boehringer-Mannheim, Germany) at 10 V/cm at 4 °C for 8 h in Tris-acetate-EDTA buffer (TEA1X, pH=8), and then the gel was stained for 1 h in 0.5 µg/ml (w/v) of ethidium bromide. Later on, the gel was washed twice with the TEA1X buffer. The bands were photographed and analyzed with an UVP gel scanner (GDS2000; Ultra Violet Product Ltd., Cambridge, UK).

RESULTS

The absorption spectra

The reactions of [AuCl(Terpy)]Cl₂ and [PtCl(Terpy)]Cl with ct-DNA were first analysed by absorption spectroscopy in the visible spectra. Remarkably, both complexes are characterized by a number of intense transitions in the 300-400 nm region. It was previously shown that the visible spectra of both complexes, dissolved in a physiological buffer, are stable with time implying that the species existing in solution do not undergo further transformations /19/. Thus, samples of the two complexes were dissolved in the buffer and titrated with increasing amounts of ct-DNA. The spectral profiles of these titrations are shown in Figure 2. Addition of ct-DNA just induces a moderate hyperchromic effect on the visible spectrum of [AuCl(Terpy)]Cl₂, probably arising from the interference of ct-DNA with the intrinsic stacking processes of this gold(III) complex. In contrast, marked changes of the visible bands of [PtCl(Terpy)]Cl are observed mainly consisting of a large hypochromic effect and a red shift of the main transition, in line with previous observations on similar systems /10,26,27/.

Circular dichroism

Addition of either [PtCl(Terpy)]Cl or [AuCl(Terpy)]Cl₂ to calf thymus DNA, at r=0.1 produces modest changes of its characteristic CD spectrum; however, if the concentration of these metal complexes is increased to r=1, large changes of the CD spectra are observed (see Figure 3). Pairwise, addition of increasing amounts of either complex to ct-DNA solutions results in the appearance of characteristic "induced CD" bands in the 300-400 nm region (see Figure 4).
Fig. 2: The visible spectra of the two complexes before and after addition of ct-DNA at variable r in the reference buffer (50 mM Na₂HPO₄ and 4 mM NaCl, pH 7.4). [AuCl(Terpy)]Cl₂ concentration was 50 μM; [PtCl(Terpy)]Cl concentration was 20 μM.
Fig. 3: CD titration of calf thymus DNA with increasing amounts of the two complexes; [AuCl(Terpy)]Cl₂ and [PtCl(Terpy)]Cl were added at the ratios: r=0.1, and 1.0 (from top to bottom on the positive band at 260 nm and the reverse on the negative band at 240 nm). DNA concentration was 1.25 × 10⁻⁴M in base pairs.
Fig. 4: CD titration of calf thymus DNA with increasing amounts of the two complexes; [AuCl(Terpy)]Cl₂ and [PtCl(Terpy)]Cl were added at the ratios: r=0.1, and 1. DNA concentration was $1.25 \times 10^{-4}$M (base pairs).
Ultrafiltration studies

Solutions containing either [PtCl(Terpy)]Cl or [AuCl(Terpy)]Cl₂ in the presence of excess ct-DNA (r = 0.1), were ultrafiltered with a Centricon device and the content of the upper and lower solutions analysed by spectrophotometry. [PtCl(Terpy)]Cl is retained in the upper solution by ct-DNA far more strongly than [AuCl(Terpy)]Cl₂, indicating that the affinity of the former complex for ct-DNA is significantly greater with respect to [AuCl(Terpy)]Cl₂ (see Figure 5).

![Graph of Ultrafiltration Experiments](image)

**Fig. 5:** The ultrafiltration experiments at half volume of the two metal compounds/calf thymus DNA adducts (r=0.1). Visible absorption spectra of the lower (down) and upper (up) solution were obtained after ultrafiltration (reducing the volume to half).
Melting studies.

The consequences of adduct formation on the thermal stability of the double helix in ct-DNA were assayed by recording the DNA melting profiles. The effects of the two complexes on the DNA helix-to-coil transition were analysed at fixed metal to bp ratio (0.1) immediately after mixing and after 24 hours of incubation (see Table 1 and Table 2). Addition of either compound produces a large thermal stabilization of the double helix. The measured increases in $T_m$ are $+6.2^\circ C$ ($8.6^\circ C$ after 24 hours) for $[AuCl(Terpy)]Cl_2$ and $+3.2^\circ C$ ($2.7^\circ C$ after 24 hours) for $[PtCl(Terpy)]Cl$.

\begin{table}
\centering
\begin{tabular}{|c|c|c|c|c|}
\hline
 & $r$ & $T_m(\circ C)$ & $\Delta T(\circ C)$ & $\Delta H$ & $\Delta T_m(\circ C)$ \\
\hline
ct-DNA & 0.0 & 60.1 & 18.4 & 0.9 & - \\
DNA+ $[AuCl(Terpy)]Cl_2$ & 0.1 & 76.0 & 12.7 & 0.3 & 6.2 \\
DNA+ $[PtCl(Terpy)]Cl$ & 0.1 & 73.1 & 10.0 & 0.3 & 3.2 \\
\hline
\end{tabular}
\caption{Table 1}
\end{table}

\begin{table}
\centering
\begin{tabular}{|c|c|c|c|c|c|}
\hline
 & $r$ & $T_m(\circ C)$ & $\Delta T(\circ C)$ & $\Delta H$ & $\Delta T_m(\circ C)$ \\
\hline
ct-DNA & 0.0 & 69.9 & 18.4 & 0.9 & - \\
DNA+ $[AuCl(Terpy)]Cl_2$ & 0.1 & 78.5 & 12.5 & 0.3 & 8.6 \\
DNA+ $[PtCl(Terpy)]Cl$ & 0.1 & 72.7 & 11.8 & 0.3 & 2.7 \\
\hline
\end{tabular}
\caption{Table 2}
\end{table}

Gel electrophoresis experiments.

Representative gels are shown in Figure 6. Treatment of plasmidic DNA with $[PtCl(Terpy)]Cl$, using a short incubation time (1h), results in a marked decrease in the mobility of the supercoiled form, while $[AuCl(Terpy)]Cl_2$ is virtually ineffective on the electrophoretic profiles.
The effects of the two complexes on the electrophoretic behaviour are strongly dependent on the incubation times. This prompted us to analyse gels of DNA-metal adducts at increasing time intervals after mixing. Results are shown in Figure 7. Electrophoretic retardation caused by [PtCl(Terpy)]Cl increased for increasing reaction times. The maximum effect, obtained after 10 hours of interaction, consisted of a remarkable retardation of plasmid mobility in the gel electrophoresis, similarly to what previously observed for cisplatin /28,29/. This behaviour is most likely attributed to formation of coordinative bonds. Notably, also in the case of [AuCl(Terpy)]Cl, some retardation in plasmidic mobility, though smaller than in the case of [PtCl(Terpy)]Cl, was observed for incubation times of 3 hours or larger (24 hours), again suggestive of formation of some coordinative bonds.
**Molecular Recognition of Metal Complexes by DNA: A Comparative Study of the Interactions**

**DISCUSSION**

There is today great interest in characterising the interactions of metal complexes with nucleic acids in view of their relevant implications in medicine /1/. Indeed, much work has been done, and is still being done, to understand the structural reasons through which metal complexes are recognised by DNA molecules, to define the nature of binding and to comprehend the molecular bases of binding selectivity /1,17/. Unfortunately, crystallographic data on DNA/metal complexes adducts are very scarce due to the intrinsic difficulties in obtaining and growing crystals appropriate for diffraction studies. Thus, very often, only indirect structural information on these systems is available that is derived from the application of other physico-chemical techniques.

| Incubation time | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | 13 | 14 | 15 |
|-----------------|---|---|---|---|---|---|---|---|---|----|----|----|----|----|----|
| [AuCl(Terpy)]Cl₂ | Ladder | pHV14 | 0h | 1h | 3h | 5h | 10h | 24h |
| [PtCl(Terpy)]Cl | Ladder | | 0h | 1h | 3h | 5h | 10h | 24h |

Fig. 7: Gel electrophoresis on 0.8% agar gel of the adducts of [AuCl(Terpy)]Cl₂ and [PtCl(Terpy)]Cl with plasmidic DNA (pHV14), r = 0.8, in the buffer TEA1X. A kinetic experiment.
In this paper we have considered, comparatively, the interaction modes of two strictly related complexes [PtCl(Terpy)]Cl and [AuCl(Terpy)]Cl₂ to double stranded DNA. The two complexes are structurally similar in that both of them manifest a classical square planar geometry within a d⁶ electronic configuration of the metal center /10,18/. The main formal difference is represented by the fact that the gold(III) complex possesses an additional positive charge. Moreover, other differences derive from the intrinsic differences in reactivity between platinum(II) and gold(III) complexes (i.e. gold(III) compounds are kinetically more labile, present pronounced oxidising properties and are far more acidic) /19/.

Notably, platinum terpyridine complexes have been the subject of several investigations during the last three decades, so that reliable models are available to describe their interactions with DNA /3-15/. At variance, only limited information is available on the gold(III) analogue that was first characterised by the group of Lippard /18/; then considered in our laboratory as a possible gold(III)-based cytotoxic and antitumor agent /19,30/.

Given the strict structural similarity of [AuCl(Terpy)]Cl₂ and [PtCl(Terpy)]Cl we proposed to establish whether such structural relatedness might result in a similar pattern of interaction with DNA.

An array of physicochemical methods has been utilised to gain information on the reaction of these metal terpyridine complexes with double stranded DNA, comprising visible spectrophotometry, circular dichroism, DNA melting analysis, ultrafiltration and gel electrophoresis.

Putting together the various pieces of information coming out from the different techniques the following picture emerges. Surprisingly, the two complexes show a pattern of interaction with DNA that is clearly distinct. In the case of [PtCl(Terpy)]Cl our data are in substantial agreement with previous reports from other laboratories /3-10/. The large variations of the visible absorption spectra observed at mixing, the profound effects produced by [PtCl(Terpy)]Cl on plasmidic DNA mobility and its tight binding to calf thymus DNA are fully consistent with the intercalation model proposed by Lippard /3-7/. Moreover, in line with previous observations /10/, we can postulate that the interaction of [PtCl(Terpy)]Cl with DNA progressively shifts with time from intercalation to formation of coordinative bonds. The time dependent changes of the gel electrophoresis patterns strongly support this interpretation.

Unexpectedly, [AuCl(Terpy)]Cl₂ exhibits a different interaction profile with double stranded DNA. Indeed, at mixing, there are modest changes in the visible spectra consisting of modest hyperchromic effects. Also, the binding affinity to calf thymus DNA is modest and the modifications of the gel electrophoresis patterns very small. These findings suggest that [AuCl(Terpy)]Cl₂ is not able to produce intercalative binding to DNA. This behaviour is rather surprising if one considers the tight structural similarity of the two complexes, the only major difference being the additional positive charge of the gold(III) analogue. Overall, the observed effects are consistent with external binding of the gold(III) complex to double stranded DNA mediated by electrostatic interactions between the positively charged complex and the negatively charged DNA double helix. Notably, with time, some differences appear in the interaction pattern of [AuCl(Terpy)]Cl₂ with DNA that, in analogy to the case of [PtCl(Terpy)]Cl, might be interpreted in terms of a progressive shift from external binding to formation of coordinative bonds. In line with this interpretation, the effects caused by [AuCl(Terpy)]Cl₂ on the gel electrophoresis profiles of plasmidic DNA become more marked.
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