

1
2 *Keywords:* antitumor; Ru(III)-bis(aminophenolate) complexes; Ru(III)-salan complexes; DNA binding
3 constants; cell cycle arrest; apoptosis; cell morphology
4

5 **1. Introduction**

6

7 Cancer is the second largest cause of death in developed countries. Recent statistics (2007-2009)
8 indicate that two in every five people born today will be diagnosed with cancer at some time during
9 their life.¹ According to the World Health Organization, cancer mortality is projected to rise to over
10 13.1 million people in 2030.²

11 Rosenberg's breakthrough of the antitumor activity of cisplatin in the sixties paved the way for the
12 research and development of metal-based drugs in cancer chemotherapy.³ Notwithstanding the
13 remarkable advances in the field, three platinum-based compounds (cisplatin, carboplatin and
14 oxaliplatin) remain to date the only approved metallodrugs for worldwide use in the chemotherapy of
15 cancer given their outstanding efficiency.⁴ However, numerous severe side-effects, their limited
16 spectra of action and tumour resistance to treatment, either intrinsic or acquired during cycles of
17 therapy, limit their clinical value. New and efficient alternative therapies are urgently and intensely
18 sought for.

19 Ruthenium complexes have consistently shown excellent anti-cancer properties both *in vitro* and *in*
20 *vivo*, and are quite promising as effective alternatives to platinum-based agents, typically offering a
21 wider spectrum of activity and the potential to overcome platinum-resistance, as well as different
22 mechanisms of action and, in general, lower toxicity.⁵⁻⁷ The intense research on this field in the last
23 few years has led to the development of several families of compounds, with a significant number of
24 Ru(II) and Ru(III) complexes showing antitumor activity in a variety of cancer cell lines (including
25 cisplatin resistant cells), ranging from inorganic-based agents to organometallic complexes.^{5,8-11} Two
26 Ru(III) lead compounds (NKP-1339 and NAMI-A) have indeed progressed successfully through phase
27 I and preliminary phase II clinical trials in patients.^{5,6}

28
29 The biological targets of antitumor ruthenium complexes have not been unequivocally identified and
30 remain the subject of intense research. Serum protein binding is crucial for the *in vivo* behavior of
31 drugs in general and in that of ruthenium-based agents in particular, since albumin and transferrin have
32 been proposed to play a role in their mechanism of action.^{8,9,12,13} It is proposed that albumin is
33 involved in the transport of ruthenium compounds in the blood stream and can act as a Ru-compound
34 warehouse^{8,13}. Transferrin has been considered to play an important role in the cellular accumulation
35 of ruthenium complexes (Ru(III) in particular) *via* transferrin-receptor mediated endocytosis.^{8,13}

36 The set of co-ligands around the metal centre is in fact crucial in modulating biological behaviour of
37 the overall complex in what concerns the cytotoxic activity, protein binding, subcellular distribution

1 and the interaction with DNA.^{8,9,13} DNA is a classical target for metallodrugs in general. Although it
2 may not be their primary target, it is very often involved in their mode of action.^{6,7} Irreversible
3 modifications in the control of cell division resulting in unrestrained cell proliferation are one of the
4 hallmarks of cancer. The first genetic alterations identified in cancer cells were seen in master genes
5 responsible for cell cycle regulation that resulted in high proliferation rate and resistance to
6 apoptosis.¹⁴ Thus, a highly desirable feature for a drug candidate is the ability to promote cell death in
7 a controlled process such as apoptosis.

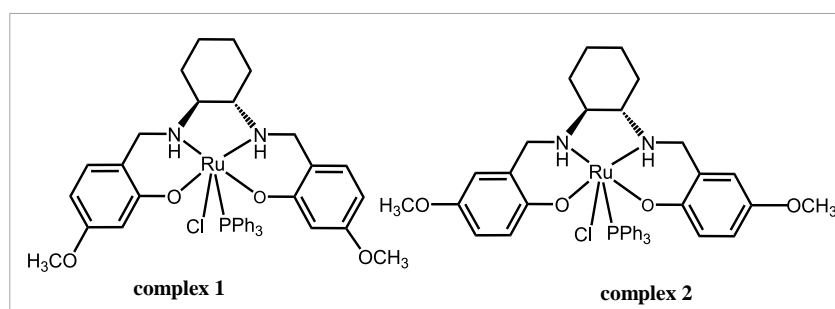
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9 Although Ru(III) complexes NKP-1339 and NAMI-A undergoing clinical development show
10 remarkable potential as anticancer drugs despite their low half-life in aqueous media, chemical
11 stability is a common requirement for a successful pharmaceutical which it can be achieved by
12 increasing the denticity of the ligand(s). Tetradentate N₂O₂ ligands combining phenolate oxygen hard
13 donors suitable for stabilizing ruthenium in the +3 oxidation state with strong nitrogen donors in
14 chelating positions are quite attractive in this context.

15 We recently reported our first work on two new ruthenium octahedral complexes with the (N₂O₂)
16 binding motif in tetradentate bis(aminophenolate) ligands (also known as ‘salan’-type). These new
17 Ru(III)-salan complexes were the first compounds of this type reported in the literature for antitumour
18 purposes.¹⁵ Both salan ligands **L1** ((6,6'-(1S,2S)-cyclohexane-1,2-diylbis(azanediyl) bis(methylene)
19 bis(3-methoxyphenol) and **L2** (2,2'-(1S,2S)-cyclohexane-1,2-diylbis(azanediyl)bis(methylene)bis(4-
20 methoxyphenol)) and their Ru(III) complexes formulated as [Ru(III)(L)(PPh₃)Cl] (see **Scheme 1**) were
21 evaluated in several human tumor cell lines.

22
23 All compounds exhibited an antiproliferative effect to different extent depending on the line tested,
24 and coordination of **L1** and of **L2** to the Ru(III) metal center resulted in enhanced activity in all cell
25 lines.¹⁵ Despite the structural resemblance of these ligands (the only difference being the *para*- or
26 *meta*-position of the methoxy group relative to the O donor on phenolate rings) complexes **1** and **2**
27 exhibited a somehow different profile in what concerned their activity: although similar and
28 comparable to cisplatin against A2780 cells (cisplatin sensitive ovarian adenocarcinoma), in breast
29 adenocarcinoma (MCF7 and MDAMB231) cells complex **2** was clearly more active than **1**, both
30 compounds being more active than cisplatin against the MCF7 ER α + (non-invasive) breast
31 adenocarcinoma line.¹⁵

32
33 The role of serum proteins on the action of these complexes was assessed as well. Binding of both
34 compounds to human serum albumin was strong enough to infer that their transport in the blood
35 plasma by this protein is quite likely, although it was observed that **2** interacted with albumin more
36 strongly than **1**.¹⁵ Prior incubation of Ru(III)-salan complexes with albumin or transferrin seemed to

1 deactivate **2** to some extent in A2780 cells, the activity of **1** being fairly unaffected by the formation of
2 the (protein-complex) adduct(s). Our results thus indicated that transferrin is likely to play a minor role
3 (if any) in the uptake of these Ru(III) agents, and that alternative pathways should be proposed when
4 clarifying their mode of action.¹⁵ These differences in their behaviour stimulated our interest on
5 understanding whether this structure-activity relationship would transpose into their pharmacological
6 properties.



9 **Scheme 1.** Structures of the two [Ru(III)(salan)(PPh₃)Cl] complexes used in this study (PPh₃ is
10 triphenylphosphane, P(C₆H₅)₃). Salan ligands **L1** and **L2**, and their corresponding [Ru(III)(L)(PPh₃)Cl]
11 complexes **1** and **2** were synthesized as previously reported.¹⁵

12
13 In this work, we proceeded with further tests for the two Ru(III)-salan complexes and used the A2780
14 line as the cell model to clarify mechanisms behind their biological activity and to understand whether
15 the small structural difference would transpose into their pharmacological properties, and with what
16 impact. In particular, we focused on the role of DNA as a possible target, assessing this interaction
17 through a novel quantitative approach to DNA-[metal complex] binding. We also evaluated the
18 contribution of programmed cell death for their cytotoxicity, as well as their effect on the cell cycle of
19 tumour cells and on the cell ultra-structural morphology.

20

21 **2. Results and discussion**

22

23 **2.1. Interaction with DNA**

24

25 **2.1.1. Steady-state spectrofluorometric measurements**

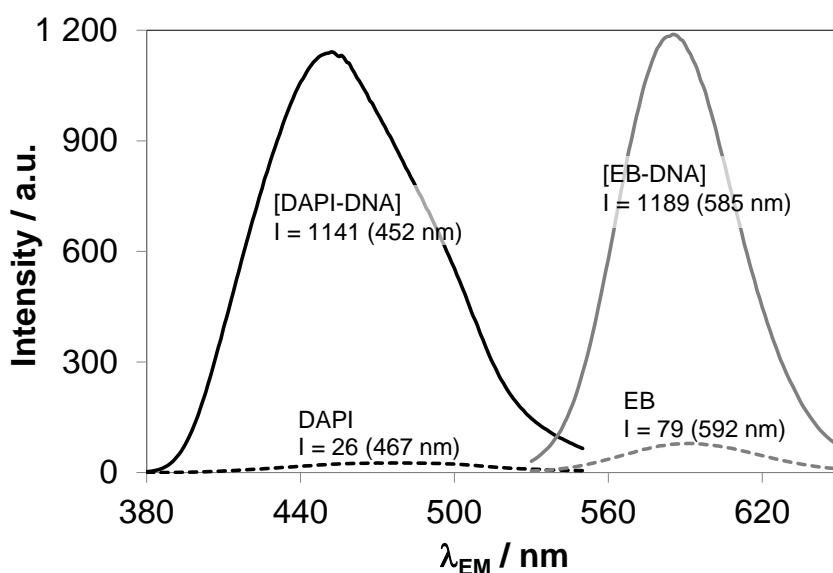
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27 Fluorescent DNA probes ethidium bromide (EB) and 4',6-diamidino-2-phenylindole (DAPI) were used
28 to characterize the interaction of the two Ru(III)-salan complexes (**Scheme 1**) with calf thymus DNA
29 (ctDNA or DNA) since they exhibit negligible fluorescence at pH 7.4 in 10 mM 4-(2-hydroxyethyl)-1-
30 piperazineethanesulfonic acid (HEPES)/2% (v/v) DMSO medium, and the interaction between ctDNA
31 and either of the two Ru-salan complexes does not lead to any measurable fluorescence. Probe

1 displacement experiments are in fact ternary systems (consisting of the probe, DNA and the ruthenium
2 complex), and the binary systems (probe and DNA) must first be characterized in the experimental
3 conditions used.

4 EB and DAPI (**Scheme S1** in **Supporting Information**) have weak intrinsic fluorescence emission at
5 the appropriate excitation wavelengths, and the binding of these probes to DNA induces an increase of
6 fluorescence quantum yield. EB is known as an intercalative agent while DAPI is a minor groove
7 binder of DNA.^{16,17} Adduct formation of both probes with DNA enhances the fluorescence intensity
8 (I_F) up to 44- and 15-fold for DAPI and EB, respectively (**Fig. 1**). At the same time a blue shift of the
9 emission can be observed (15 nm and 7 nm for DAPI and EB, respectively) due to the insertion of the
10 probe into a less polar environment as compared to the aqueous medium surrounding the unbound free
11 molecule.¹⁸

12



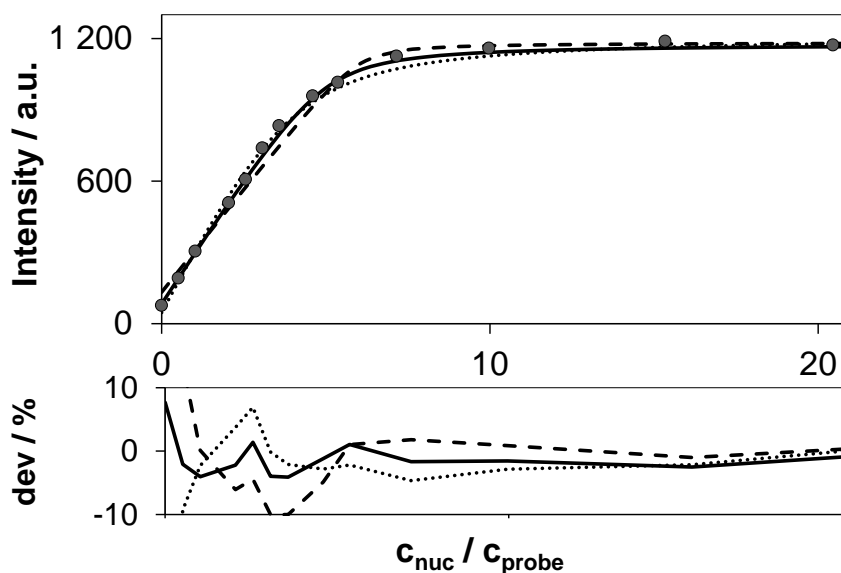
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14 **Fig. 1.** Fluorescence emission spectra of DAPI (black dotted line), DAPI-ctDNA (black solid line), EB (grey
15 dotted line) and EB-ctDNA (grey solid lines). ($C_{\text{DAPI}} = 3 \mu\text{M}$; $C_{\text{EB}} = 5 \mu\text{M}$; $\lambda_{\text{EX}} = 365 \text{ nm}$ (DAPI) or 510 nm
16 (EB); [probe-DNA] spectra were obtained by probe-DNA titrations when the saturation of the probe with DNA
17 was achieved, $c_{\text{ct-DNA}} = 51 \mu\text{M}$ (DAPI), $15 \mu\text{M}$ (EB); $t = 25 \text{ }^\circ\text{C}$; $\text{pH} = 7.40$ 10 mM HEPES/ 2% (v/v) DMSO).

18

19 Prior to the probe displacement experiments, the probe-DNA binding constants and the size of the
20 binding site were calculated on the basis of independent measurements. In order to evaluate the
21 binding site size of a probe on DNA two series of experiments are needed: measurements at constant
22 DNA concentration with varying probe concentrations and *vice versa*, where in both cases saturation
23 should be attained. Two “virtual” molar intensities (I_{molar}) can be calculated from the measured
24 intensities (corrected by the emission intensity of non-bound probe). The ratio of these values (I_{molar} (at
25 constant DNA concentration) / I_{molar} (at constant probe concentration)) gives an estimation of the
26 average binding site size represented in the number of the nucleotides (nuc).

1 To calculate the binding constants between DNA and the probes, complex formation between the
 2 binding site of DNA and the probe is assumed to be a one-step reaction; hence it can be characterized
 3 by one conditional binding constant ($\log K'$) (see details in **Supporting Information, Section S1**).
 4 The binding constants were calculated with the computer program PSEQUAD.¹⁹ The PSEQUAD
 5 program was originally developed for determination of stability/binding constants from experimental
 6 equilibrium data collected by various methods such as pH-potentiometry, and/or spectroscopy
 7 (UV/Vis, NMR), and this software can also be used on fluorometric data evaluation as reported
 8 previously for protein-ligand (metal complex) interactions.^{20,21} After a set of iterative steps, the
 9 program provides the binding constants of the selected adducts with standard deviations and the molar
 10 emission spectra of the emitting components/adducts. Calculations were performed for different sizes
 11 of the binding site (*i.e.* different number of nucleotides), and the best fit for EB and DAPI was found
 12 for 5 and 23 nucleotides, respectively. In **Fig. 2** the measured and calculated fluorescence intensities
 13 are shown for the EB – DNA system considering binding site sizes of 4, 5 and 6 nucleotides, as well as
 14 the experimental data. The calculated values (average binding site size) were based on the results of
 15 PSEQUAD calculations, namely the binding constant and molar intensities. Although the fit for any of
 16 the assumed binding site sizes seems to be very similar at a first glance, a deeper statistical analysis
 17 revealed that the best fit is obtained for an average binding site size of 5, which is in good agreement
 18 with literature data^{16,22,23} and the $\log K'$ value obtained is collected in **Table 1**. Further refinements did
 19 not result in any appreciable improvements in fitting to the experimental data.



20
 21 **Fig. 2.** Total fluorescence intensity of EB samples titrated by ctDNA: *up*: measured data (●) and simulated
 22 curves for binding site sizes $n = 4$ (dotted line), 5 (black line) or 6 (dashed line); *below*: deviation (dev.) of
 23 simulated curves from measured intensities in %; ($c_{EB} = 5 \mu\text{M}$; $c_{nuc} = 0\text{-}100 \mu\text{M}$; $\lambda_{EX} = 510 \text{ nm}$; $\lambda_{EM} = 585 \text{ nm}$;
 24 $t = 25 \text{ }^\circ\text{C}$; $\text{pH} = 7.40$ (10 mM HEPES/2% (v/v) DMSO); simulated values based on PSEQUAD calculations –
 25 see **Table 1**).
 26

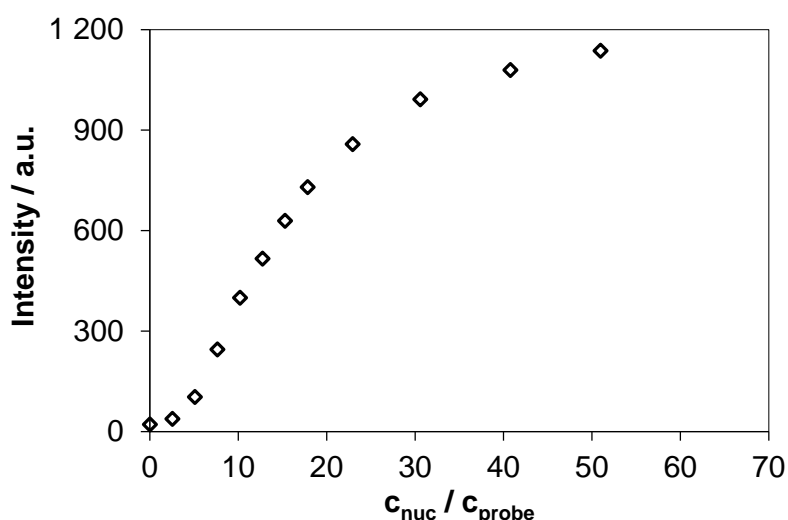


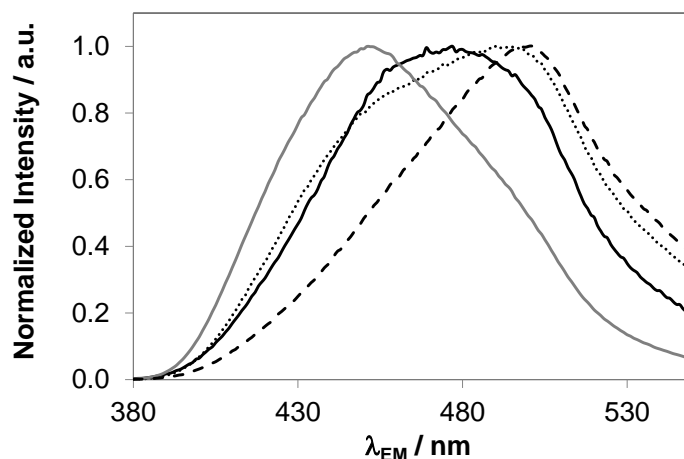
Fig. 3. Change in fluorescence emission intensity at 450 nm with increasing ratio of ctDNA nucleotides (nuc)-to-DAPI ($C_{\text{nuc}}/C_{\text{probe}}$) { $c_{\text{DAPI}} = 3.06 \mu\text{M}$; $c_{\text{nuc}} = 0\text{-}160 \mu\text{M}$; $\lambda_{\text{EX}} = 365 \text{ nm}$; $t = 25 \text{ }^\circ\text{C}$; $\text{pH} = 7.40$ (10 mM HEPES / 2% (v/v) DMSO)}.

Table 1

The interaction of DAPI with DNA is much more complicated, the binding mode presumably varying with the increasing ratio of DAPI-to-nucleotides. The appearance of a sigmoidal shape in the measured fluorescence emission intensity (**Fig. 3.**) confirms the existence of at least two different binding modes of DAPI to DNA that has an effect on the quantum yield of the DNA-DAPI adduct. Other works based on various techniques reported the existence of two distinct DNA-DAPI adducts depending on the molar ratio.^{17,24,25} This behaviour is also observed for samples where the DNA concentration was kept constant and DAPI concentrations were varied (**Fig. 4.** and **Fig. S2.**). Normalized spectra undoubtedly show a significant red shift of the emission maxima at 1:2 and 1:5.6 DAPI-to-DNA ratios compared with the spectrum of DAPI without DNA (**Fig. 4.**). This effect is usually explained by the intercalative interaction between DNA and the probe.^{17,25} The spectrum recorded at high excess of nucleotides (DAPI: nucleotide = 1:14) represents the expected typical blue shift associated with groove binding. Under these conditions, a binding site size of 23-26 nucleotides and a log K' value of (6.42 ± 0.22) could be estimated (**Table 1**). The nature of the interaction at higher DAPI/nucleotide ratios is however not relevant at the conditions used here for ternary systems.

Eriksson *et al.* published a value of 10 nucleotides (5 base pairs) for the interaction of DAPI with synthetic poly{d(A-T)₂} DNA.²⁶ The difference between our findings and this data can be explained by the use of different DNA source, and reflects the base sequence within the nucleotide chain. DAPI prefers A-T rich sequences and ctDNA has roughly 40% G-C (42 mole % G-C, 58 mole % A-T). Hence, when using spectroscopic methods, the estimated value for the binding site size also expresses the frequency of the preferred base sequence within the nucleotide chain.

1



2

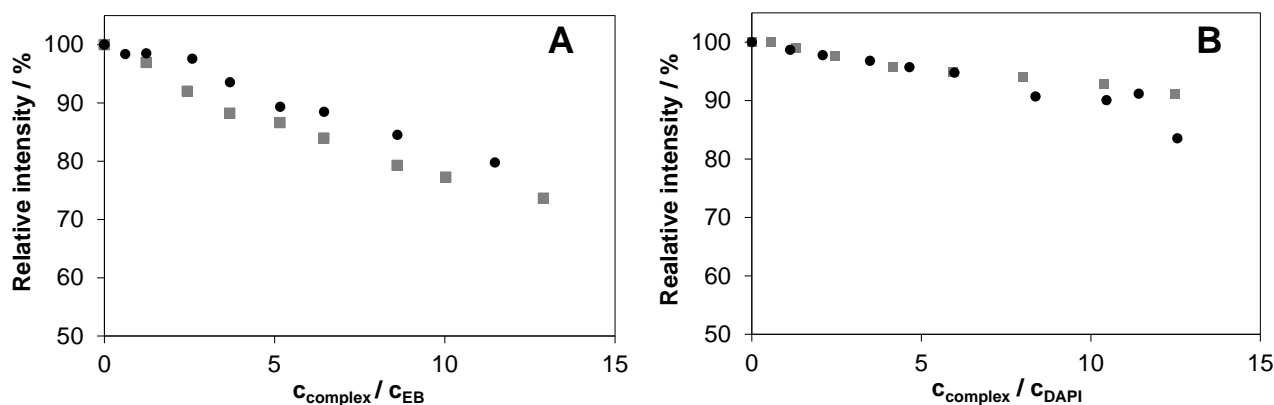
3 **Fig. 4.** Normalized emission spectra for the DAPI-DNA system at various DAPI:nucleotide ratios: 1:0 (black
 4 line), 1:2 (dashed line), 1:5.6 (dotted line), 1:14 (grey line); $\{c_{\text{DAPI}} = 0.5\text{-}13 \mu\text{M}; c_{\text{nuc}} = 26 \mu\text{M}; \lambda_{\text{EX}} = 365 \text{ nm}; t =$
 5 $25 \text{ }^\circ\text{C}; \text{pH} = 7.40 (10 \text{ mM HEPES}/ 2 \% (v/v) \text{ DMSO})\}$.

6

7 An important factor to consider when comparing literature data, in addition to the specific
 8 experimental conditions, is the calculation method used. Binding constants found in the literature are
 9 typically obtained by graphical methods such as Lineweaver-Burk, Stern-Volmer or Hildebrand-
 10 Benesi (the former two approaches are used for fluorometry data, the latter one for UV-Vis
 11 measurements), and are not directly comparable with our results since the methods mentioned do not
 12 take into consideration the actual size of the binding site.^{18,27-29} These approaches are somewhat
 13 limited since the analytical (total) concentration of DNA and probe are used instead of their
 14 equilibrium (free) concentrations, and only one emitting species is usually proposed. They should be in
 15 most cases considered as an empirical description of the results allowing only for comparison of the
 16 binding affinities of different compounds measured under the same experimental settings.³⁰ The use of
 17 analytical instead of equilibrium concentrations leads to only a rough estimate of the binding constants
 18 (see **SI** for details). It is also noteworthy that authors often do not mention whether the binding
 19 constants are calculated on the basis of the concentration of DNA in nucleotides or in base pairs,
 20 which results in different values that again are not comparable with each other. Our approach using
 21 PSEQUAD circumvents these limitations by calculating the binding constant(s) involved from actual
 22 equilibrium concentrations of all species present in each system, as well as the binding site size (in
 23 probe-DNA binary systems) and the stoichiometry of the adduct(s) formed, affording a more reliable
 24 quantitative evaluation of the systems being studied.

25 Considering the ternary systems, probe displacement reactions with both Ru(III) complexes were
 26 carried out using constant DNA and probe concentrations, and varying the concentration of the metal
 27 complex. The ratio of DNA-to-EB was chosen as 1:4, and at this EB excess we could calculate that
 28 83% of the probe is bound to DNA. **Fig. 5A** shows the effect of complex **1** and **2** on the emission

1 intensity measured for the DNA-EB adduct. The simulated curves are in good agreement with the
 2 experimental data (see **Fig. S3**). This suggests that the complex can displace the intercalated EB from
 3 its binding site with a concomitant decrease in the measured intensities, however the formation of a
 4 ternary DNA adduct containing both the probe and the bound metal complex is also feasible. The fact
 5 that no shift of the emission maximum is observed together with the results from our time resolved
 6 experiments (*vide infra*) confirms the simple displacement hypothesis. The binding constants
 7 calculated for the complexes (**Table 1**) is only valid for EB "binding sites" since the size of binding
 8 site (or binding frequency) of complex **1** and **2** on DNA is most likely not the same as in the case of
 9 EB. In order to solve this question, it would be helpful to see the saturation section of the displacement
 10 reaction, which was not attainable due to solubility limitations at higher concentrations.
 11 A similar behaviour was found for both complexes in the displacement reactions with EB, although the
 12 estimated binding constant was somewhat lower for complex **2** (see **Table 1**).
 13 To assess the possible interactions of the metal complexes and DNA at its minor groove, DAPI was
 14 used at fixed DAPI:DNA ratio of 1:25 and the fluorescence emission intensity recorded is depicted in
 15 **Fig. 5B**. In this case, even for a 12-fold excess of metal complex **1** or **2** the extent of quenching is only
 16 *ca.* 10%, *i.e.*, no remarkable changes in the measured emission intensities were observed (see **Fig S4**
 17 for fitted curve). The probe-DNA displacement reaction for both complexes is less extensive in the
 18 case of DAPI as compared to EB, and thus our results suggest that binding at the minor groove is not
 19 the preferred mode of interaction with ctDNA for the two Ru(III) complexes studied.
 20



21
 22
 23 **Fig. 5.** Relative fluorescence intensity measured for the EB – DNA (A) and DAPI – DNA (B) system titrated by
 24 complexes **1** (■) and **2** (●) { $c_{EB} = 5.0 \mu\text{M}$; $c_{DAPI} = 3.1 \mu\text{M}$; $c_{nuc} = 19.2 \mu\text{M}$ (A) or $77.1 \mu\text{M}$ (B); $\lambda_{EX} = 510$ (A),
 25 365 (B) nm; $\lambda_{EM} = 595$ (A), 450 (B) nm; $t = 25 \text{ }^\circ\text{C}$; pH = 7.40 (10 mM mM HEPES)/ 2 (v/v) % DMSO}.

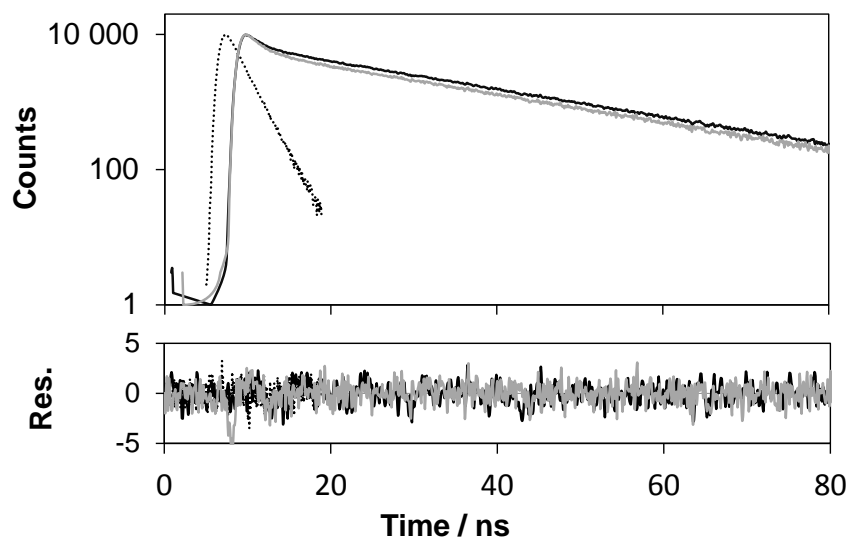
27 2.1.2. Time-resolved fluorescence measurements

28
 29 Fluorescence lifetime measurements were carried out in order to have a deeper insight into the nature
 30 of the interaction of the metal complexes **1** and **2** with DNA. The lifetime of the probe alone (EB and

1 DAPI) and of probe-DNA adducts (EB-DNA and DAPI-DNA) were obtained in independent
2 experiments prior to the competition studies. The fluorescence intensity decays of free EB (assessed
3 using samples without DNA) were very well described by a single exponential, with an associated
4 lifetime of 1.8 ns (**Fig. 6**). In the case of the samples containing EB and DNA, the intensity decay was
5 always described by a sum of two exponentials because there was always a fraction of free probe in
6 solution. Indeed, the fluorescence lifetime of one component of the exponential fit was similar to that
7 obtained in the absence of DNA (1.86 ns), and the other one was 21.2 ns (**Table 2**). Both these values
8 are in very good agreement with literature data for free and DNA-bound EB, respectively. The large
9 difference between these two lifetime values allows to separate very easily the contributions of DNA-
10 bound and unbound EB to the total fluorescence decay in a sample where both species are present.
11 Moreover, it is consistent with the marked increase in fluorescence intensity previously described for
12 the emission spectra of the probe (**Fig. 1**).

13
14 In the EB-DNA system (**Fig. 6**) the fluorescence intensity decay is very well described as a sum of two
15 exponentials in both the presence or absence of the studied metal complexes (**Table 2**), indicating the
16 presence of only two emitting species in the system. Additionally, the lifetime retrieved for the adduct
17 [EB-DNA] is not significantly changed at a 13-fold excess of the metal complexes, which can exclude
18 the possibility of collisional quenching, or any other mechanism of interaction where *e.g.* both probe
19 and metal complex would be interacting at the same DNA site. The same fluorescence lifetime values
20 for the free and for the bound probe are retrieved in the ternary systems, which indicates not only the
21 presence of only two emitting species in solution, but also that these are the same species as in the
22 probe – DNA binary systems. This shows that the reaction taking place in solution is a true
23 displacement reaction and that neither of the Ru(III) complexes are damaging the DNA structure, thus
24 preventing EB from binding in the same way as in the absence of the compounds. In **Table 2**, results
25 for EB – DNA system in the presence of increasing amounts of complex **1** and **2** are shown.

26



1
 2 **Fig. 6.** Intensity decay of the emission fluorescence of EB alone (black dotted line), of EB – DNA (black line),
 3 and of EB – DNA – (complex **2**) (grey line) systems { $c_{\text{EB}} = 5.0 \mu\text{M}$; $c_{\text{nuc}} = 19.5 \mu\text{M}$; $c_{\text{complex } 2} = 63.9 \mu\text{M}$; $\lambda_{\text{EX}} =$
 4 460 nm ; $\lambda_{\text{EM}} = 590 \text{ nm}$; $t = 25 \text{ }^\circ\text{C}$; $\text{pH} = 7.40$ (10 mM HEPES/2% DMSO)} and (bottom) residuals plot for the
 5 fitting of each decay in the ternary systems with a sum of two exponentials ($\tau_1 = 1.86 \text{ ns}$, $\tau_2 = 21.2 \text{ ns}$ – see
 6 **Table 2**).

7
 8 Time-resolved fluorescence is a sensitive enough technique to differentiate the contribution from two
 9 emissive species even when one of them is a minor contributor to the overall fluorescent signal. In this
 10 way, if the changes in average fluorescence lifetime calculated from the two-exponential fit to the
 11 fluorescence intensity decay of EB are in quantitative agreement with the binding model proposed
 12 with log K' values calculated with PSEQUAD (*vide supra*), it further supports the suitability of the
 13 model.

14
 15 **Table 2**

16
 17 The pre-exponential of the short lifetime component (α_1) increases as the concentration of Ru(III)
 18 complex is increased (for both **1** and **2**), which is the expected trend for an EB displacement from
 19 DNA. The fraction of light emitted by free probe (f_1) is always below 11%, but it is still possible to
 20 appreciate its increase upon complex addition to the system, highlighting the fact that it is possible to
 21 separate the contribution from both species using time-resolved fluorometry. Moreover, it justifies the
 22 trend observed for the steady-state fluorescence intensity decrease observed in the competition
 23 experiment with EB (**Fig. 5**) where the extent of quenching never attained even 50%. This does not
 24 mean that the compound under test has a weak interaction with DNA, but reflects merely the fact that
 25 even a very small fraction of DNA bound probe will contribute a very significant fraction of light to
 26 total emission.³¹ The amplitude-averaged mean fluorescence lifetime (τ_{av}) is a measure of the

1 fluorescence quantum yield in the absence of static quenching. It decreases by 22% for a complex **1**:
2 EB ratio of 1:12.0. This is in excellent agreement with the simulated curve based on PSEQUAD
3 calculations shown in **Fig. S3**. Such result not only rules out that a static quenching process could
4 occur, *e.g.* by an interaction of probe with the complex in solution, but also it quantitatively validates
5 the binding model proposed (**Table 1**). For complex **2**, τ_{av} decreases by 15% for a complex: EB ratio
6 of 10.5, which is also in agreement with the binding constant present in **Table 1**.

7 Similar measurements were performed with DAPI. Again, a single exponential function yielded a
8 good fit to the experimental intensity decay for the probe in solution (without any improvement by
9 adding further exponentials). The value retrieved for the fluorescence lifetime of DAPI (in the absence
10 of DNA) was 1.57 ns, in close agreement with the literature.¹⁸ The fluorescence intensity decay of
11 DAPI was also measured in the presence of DNA and complexes **1** and **2** (**Table 2**). As for EB, the
12 fluorescence decay became a bi-exponential function, with one component coincident with the one
13 observed for free DAPI, and a longer component of 3.6 ns assigned to DNA-bound probe. In contrast
14 to the observations with EB but in agreement with the steady-state intensity results (**Fig. 5B**), the
15 average fluorescence lifetime for DAPI and the adduct [DAPI-DNA] are barely changed upon addition
16 of metal complexes. Noteworthy is the fact that although the noted changes are rather small for both
17 complex **1** and **2**, it is slightly stronger for the latter, a behaviour also suggested by the fluorescence
18 intensity results of **Fig. 5B**. This might denote that this compound may have a very mild minor groove
19 binding ability, or because it is not such a strong intercalating agent as **1** it may be more available for
20 non-specific interactions with DNA eventually displacing a small fraction of the minor groove binder
21 DAPI.

22
23 To summarize, the interaction of both Ru(III)-salan complexes with DNA is assessed by steady-state
24 and time-resolved fluorescence spectroscopy using the probes EB and DAPI. In this work we use a
25 method for evaluating fluorescence results that provides the conditional stability binding constants for
26 the interaction by an iterative procedure based on the true equilibrium concentrations for all species
27 present in solution, in contrast with other methods reported in the literature where total concentrations
28 are used instead. Our approach also affords the binding site size involved in DNA-probe interactions
29 without any *a priori* assumptions. Time-resolved fluorescence spectroscopy enabled to identify the
30 emitting species in these systems. In both DNA-probe binary systems and DNA-probe-Ru(III)
31 complex ternary systems, the only two fluorescent species observed are the bound and unbound probe,
32 the concentration of the latter reflecting the increase in the concentration of the Ru(III) complex as a
33 consequence of the competition reaction. EB is displaced from its DNA adduct in the presence of both
34 Ru(III) complexes, and the competition experiments afford stability constants for the interaction of
35 complexes **1** and **2** with DNA with higher accuracy than in the case of DAPI. It was possible to
36 estimate stability constants from the effect of **1** and **2** on the displacement reaction of DAPI from the

1 DNA-DAPI adduct, which yielded the same value for both complexes but since the effect observed in
2 these systems is quite small, these K'_{binding} values must be taken as rough estimates. Log K'_{binding} values
3 obtained indicate that the interaction is moderately strong, and suggest that the interaction of **1** with
4 DNA is somewhat stronger than that of **2**.

5
6 These results are not reflecting the relative cytotoxic activity (IC_{50} values) observed for these
7 complexes, with **2** being more active than **1** in all cell studies done (at either 24 h or 72 h challenges).
8 As such, DNA is a possible molecular target for these complexes, but other targets are most probably
9 involved in their action as well.

11 **2.2. Studies with ovarian adenocarcinoma cells**

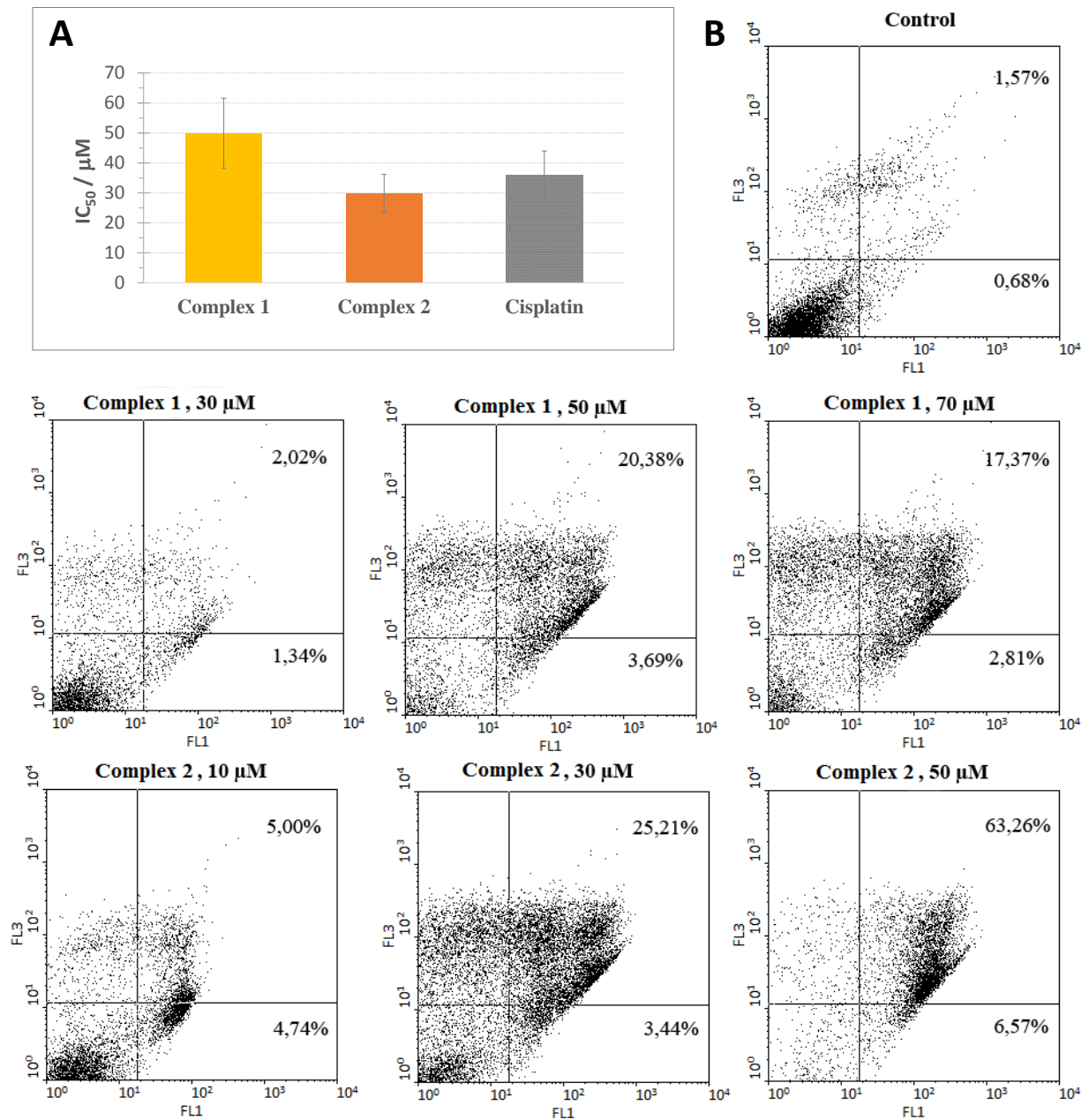
13 *2.2.1. Cytotoxic activity and mechanism of cell death*

14
15 The cytotoxic activity of complexes **1** and **2** was evaluated in human cisplatin sensitive ovarian
16 adenocarcinoma A2780 cells for a 24 h incubation to allow a suitable choice of the concentrations to
17 use in the flow cytometry assays which involved a 24 h incubation period. The IC_{50} values determined
18 in these conditions were 50 μM for complex **1** and 30 μM for complex **2**. These values are comparable
19 to the cytotoxic activity of cisplatin for A2780 cells in the same conditions (after a 24 h incubation
20 period), 36 μM (**Fig. 7A**), and the quantitative relation among the IC_{50} values for these compounds is
21 in general the same as observed for a 72 h challenge.¹⁵

22
23 The mechanism of cytotoxicity induced by complexes **1** and **2** was assessed by flow cytometry
24 analysis of cells stained for Annexin-V, a Ca(II)-dependent phospholipid-binding protein with high
25 affinity for phosphatidylserine (FL1), and propidium iodide (PI, that stains DNA of permeabilized
26 cells and shows unviable cells). Fluorescence of PI (FL3) was plotted versus Annexin-V fluorescence
27 (FL1) as shown in **Fig. 7B**.

28 Viable cells are both Annexin-V and PI negative (Annexin-V/PI⁻), while cells that are entering
29 apoptosis are Annexin-V positive (Annexin-V⁺) and progressively more positive for PI. **Fig. 7B** shows
30 the majority of control cells stained negative for both PI and Annexin-V, corresponding to viable cells.
31 Cells treated with low doses of the compounds showed a small population of Annexin-V⁺ PI⁻ cells,
32 indicating apoptosis at an early stage. As expected, with higher complex concentrations, the population
33 of double positive cells increased indicating an increase in apoptosis. Taken together, results show that
34 both complexes **1** and **2** induce cell death mainly through an apoptotic process, and that complex **2**
35 promotes this more efficiently than **1**, an interpretation validated by the good correlation with the
36 cytotoxic activity data (**Scheme 1** and **Fig. 7A**).¹⁵

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Fig. 7. Effect of complexes 1 and 2 in A2780 cells in a 24 h challenge: A) Cytotoxic activity (measured as the IC₅₀) for both complexes and for cisplatin tested in the same conditions, and B) Quantification of apoptosis in cells treated with complexes 1 and 2 using the Annexin-V/PI double staining assay. A2780 cells (control) were treated with 30, 50 and 70 μM of complex 1 and 10, 30 and 50 μM of complex 2 for 24 h, and stained with FITC-conjugated Annexin-V and PI. Fluorescence was analyzed by flow cytometry. The dual parametric dot plots combining Annexin-V (FL1) and PI (FL3) fluorescence show the viable cell population in the lower left quadrant Annexin-V⁻/PI⁻, the early apoptotic cells in the lower right quadrant Annexin-V⁺/PI⁻, and the late apoptotic cells in the upper right quadrant Annexin-V⁺/PI⁺. Data shown are a representative experiment of two independent experiments with similar findings.

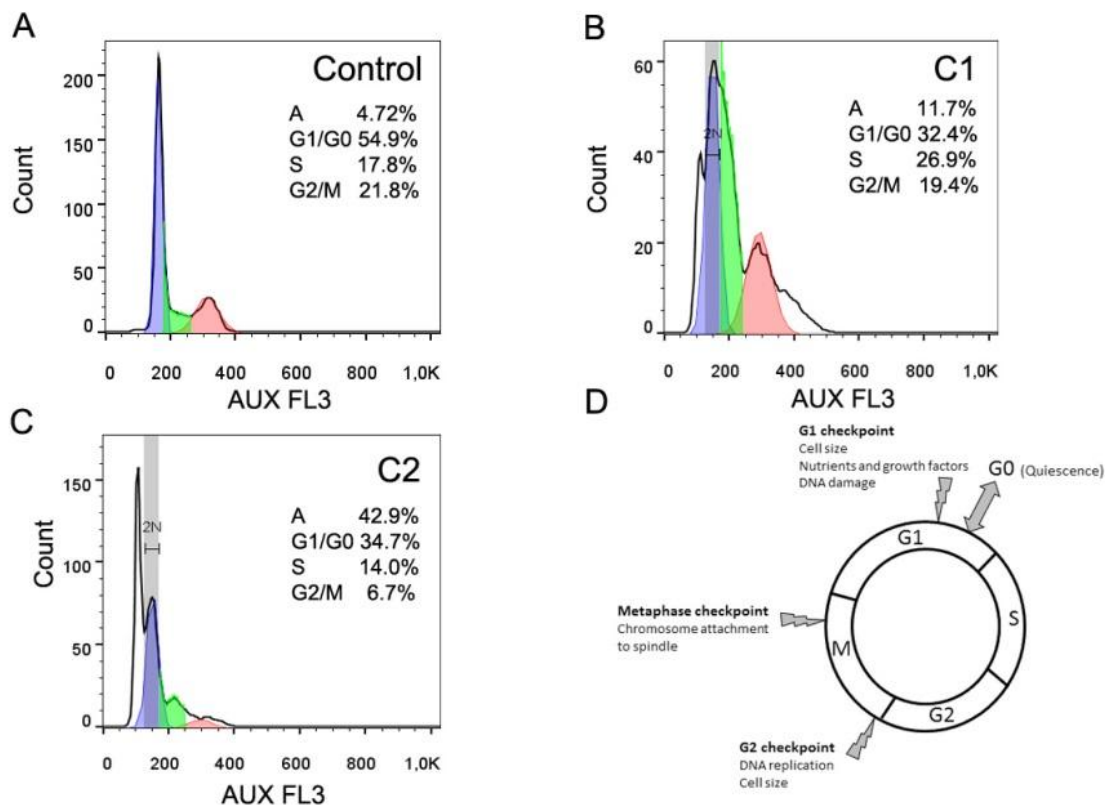
1 2.2.2 Effect of Ru(III)-salan complexes on the cell cycle

2

3 To investigate whether **1** and **2** affect cell cycle progression, cell cycle analysis was assessed by flow
4 cytometry in A2780 cells treated with the Ru(III) complexes for 24 h at the IC₅₀ concentration. **Fig. 8**
5 shows that the effect of complexes **1** and **2** in the cell cycle of A2780 cells is different. While control
6 cells were mainly in G1/G0 and G2/M phases of cell cycle (**Fig. 8A**), the cycle progression was altered
7 after treatment with complex **1** (**C1**), leading to a decrease in the percentage of cells in the G1/G0
8 phases and an increase in the proportion of cells in the S phase of the cell cycle compared to the
9 control (**Fig.8B**). This result suggests that complex **1** might induce cell cycle arrest at G2 or M phases
10 indicating that it might induce incomplete DNA synthesis (G2 checkpoint) and/or disruption of the
11 mitosis process including alterations in the chromosome assembly and/or in the spindle formation
12 (**Fig. 8D**). These alterations are not repaired by cellular mechanisms resulting in apoptosis. By
13 contrast, cells treated with complex **2** (or **C2**) presented a decrease in the percentage of cells in S and
14 G2/M phases of the cell cycle compared to control (**Fig. 8C**), which suggests that complex **2**
15 efficiently induces apoptosis with a minor effect on cell cycle arrest, and that **2** might induce DNA
16 damage or dysfunctional metabolic state that prevents cell cycle progression to the synthesis phase (S).

17

18 The dissimilar effect observed on the A2780 cell cycle promoted by these complexes could be related
19 to the different importance that DNA has as a molecular target for each compound (**Table 1, Fig. 6**).
20 In fact, although both complexes could efficiently displace EB from ctDNA, the one that was able to
21 do it more efficiently is the compound with higher IC₅₀, suggesting that other cellular targets
22 contribute to the cytotoxic activity. The different position of the methoxy group in the benzene ring
23 may influence the hydrophilicity of the compound. In fact, complex **2** has a more hydrophilic character
24 than complex **1** (see **Supporting Information, Section S2**), which might influence their relative
25 affinity for different sub-cellular structures, *e.g.* cellular membranes or cytoskeleton.



1
2 **Fig. 8.** Cell cycle analysis of A2780 cells in control (A, cells with no treatment), treated with complex 1 (C1)
3 (B) and treated with complex 2 (C2) (C), and schematic overview of cell cycle regulation including the cell cycle
4 checkpoints (D). Cells were treated with the complexes (30 μ M for 1, 10 μ M for 2) for 24 h and were analyzed
5 by flow cytometry; values indicated are the percentage of cells in the different phases of the cell cycle; *insets* -
6 A: apoptosis; G1/G0: Gap 1 and Gap 0 S: DNA synthesis G2: Gap 2; M: mitosis.

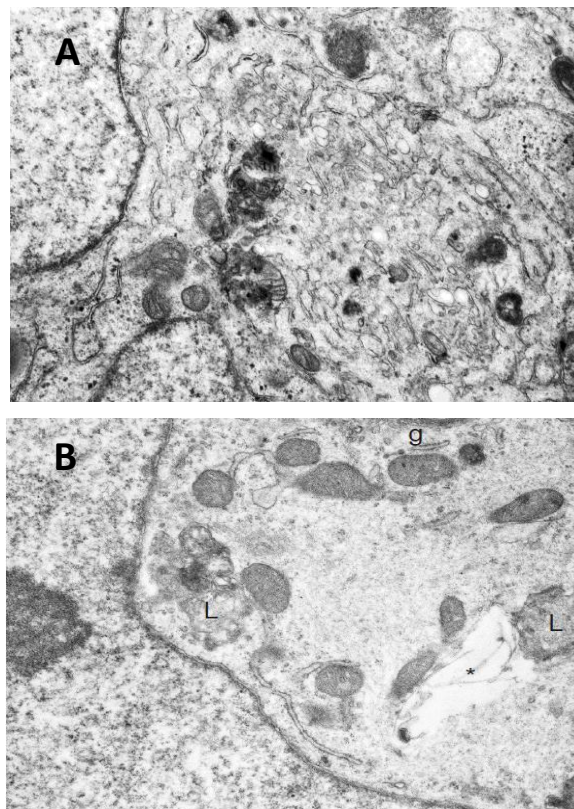
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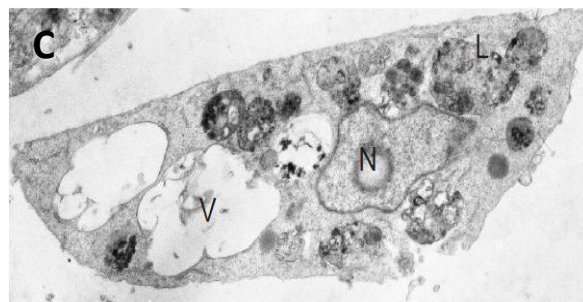
9 **2.2.3. Ultrastructural analysis and effect on cell morphology**

10

11 Transmission Electron Microscopy (TEM) was used to investigate the effects of both ruthenium
12 complexes on the morphology of ovarian cancer cells. The ultrastructure analysis by TEM can give an
13 indication of the cell damage brought upon by the action of prospective anticancer agents. By
14 affording ultra-resolution images, it allows detecting morphological changes associated with the action
15 of the agents being tested, points out potential intracellular targets, and makes apparent alterations in
16 cell physiology, thus providing information on the underlying mechanisms of action and potential
17 targets in the cell. Both complexes 1 and 2 were analyzed for their effect on A2780 cells after a 24 h
18 incubation period, and images of treated and untreated cells were compared. **Fig. 9** depicts
19 representative TEM images recorded in each case.

1 As can be seen in **Fig. 9B**, cells treated with complex **1** exhibit an underdeveloped/shrunken Golgi
2 apparatus and the lysosomes present a heterogeneous content. Golgi fragmentation is recognized as an
3 early apoptotic event.³² Although mitochondria seemed normal, breaches in the cytoplasm could be
4 observed suggesting disorganized membranes. These observations are in contrast with those of
5 controls (**Fig. 9A**) confirming the effects of complex **1** on the endomembrane system. Cells treated
6 with complex **2** (**Fig. 9C**) show enlarged lysosomes, but most noticeably, large vacuolar structure,
7 with the prototypical morphology of autophagic vacuoles, suggesting that (macro)autophagy might be
8 occurring. Annealed nucleoli were also observed, suggesting that complex **2** induces the inhibition of
9 ribosome synthesis, may be as a result of DNA dysfunction. Overall, cell alterations are more marked
10 with compound **2** that display signs of profound and probably irreversible cellular damage. This could
11 be consistent with increased late apoptosis induced by compound **2** found by flow cytometry.
12 However, until recently apoptotic cell death and autophagy (that could also lead to cell death) have
13 been considered in general mutually exclusively processes. Nonetheless, it is currently known that in
14 certain cases, a mixed apoptotic-autophagic route can be followed as a different type of programmed
15 cell death.³³ Therefore, it is tempting to speculate that compound **2** could be inducing this sort of
16 process, a hypothesis that we will test with more detailed studies on the cell death mechanism induced
17 by complex **2** in the near future.
18





1
2 **Fig. 9.** Representative TEM images showing the morphology and ultrastructure of A2780 cells (g - Golgi
3 apparatus; L - lysosomes; N – nucleolus; V- vacuoles). (A) Controls (untreated cells) showing normal well
4 developed Golgi apparatus; (B) cells treated with complex 1 (50 μ M, 24h) showing underdeveloped Golgi
5 System; Lysosomes with heterogeneous content, and breaches in the cytoplasm suggesting disorganized
6 membranes; (C) cells after treatment with complex 2 (50 μ M, 24h) showing annealed nucleolus indicating the
7 inhibition of ribosome synthesis, large lysosomes suggesting autophagy and large vacuoles.

10 3. Conclusions

11
12 Two Ru(III)-salan complexes with promising antitumor activity were evaluated to outline the basis of
13 their biological action. DNA, typically involved in the mode of action of metallodrugs, is usually
14 proposed as a possible target and the interaction of both complexes with the biopolymer was assessed
15 by steady-state and time-resolved fluorescence spectroscopy using competition experiments with the
16 well-established probes EB and DAPI. Although the displacement effect from its DNA adduct
17 observed with DAPI in the presence of both Ru(III) complexes is quite small, EB displacement
18 afforded accurate stability conditional binding constants for **1** and **2**. Log K'_{binding} values obtained
19 indicate that the interaction is moderately strong, and suggest that the interaction of **1** with DNA is
20 somewhat stronger than that of **2**. These findings indicate that DNA is a possible molecular target for
21 these complexes, but other targets are most likely involved in their mode of action. In fact, this data is
22 not reflecting the relative cytotoxic activity observed, with **2** being more active than **1** in all cell
23 studies done, and results on the morphology and cycle progression of A2780 cells treated with the two
24 compounds seem to support this hypothesis.

25 In what concerns the mechanism of cell death, both complexes were found to induce apoptosis in
26 A2780 cells with complex **2** being the most efficient. This finding is in agreement with the cytotoxic
27 activity found for the complexes considering apoptosis as the main cell death mechanism.

28 In the cell cycle of A2780 cells complex **2** induces apoptosis without a relevant effect of cell cycle
29 arrest. This suggests that **2** might induce DNA damage or a dysfunctional metabolic state that
30 precludes cell cycle progression to the synthesis phase, which is in agreement with the autophagic

1 process suggested by the ultra-structural analysis observed in cells treated with **2**. In contrast, results
2 for complex **1** suggest that it may induce cell cycle arrest at G2 or M phases, resulting in promoting
3 apoptosis.

4 The distinct effect induced by these Ru(III) complexes on the A2780 cells is probably related to
5 differences in their hydrophilicity. In fact, complex **2** has a more hydrophilic character than **1**, which
6 indicates differences in how they interact with plasma and intracellular membranes *in vivo* are to be
7 expected, and would have an impact in their cellular uptake.

8 The fact that cell morphology is affected in a different manner upon incubation with **1** and **2** further
9 supports the hypothesis that several targets are involved in their action. This is in agreement with the
10 belief that Ru-based metallodrugs exert their effect through multiple targets.¹³ This might be a valuable
11 advantage when compared with other anti-cancer agents, either by allowing the use of lower doses *in*
12 *vivo* (thus reducing side effects) or by rendering the development of chemoresistance by tumor cells
13 more difficult. The results presented in this work add further support for the high potential of these
14 Ru(III)-salan complexes as prospective metallodrugs for cancer therapy and highlights the striking
15 impact that small differences in the structure of the ligands coordinated to the metal centre may have
16 on the biological response of the overall compound.

19 **4. Materials and Methods**

21 *4.1. Chemicals*

23 Deoxyribonucleic acid sodium salt from calf thymus (ctDNA), ethidium bromide (EB), 4',6-
24 diamidino-2-phenylindole (DAPI), 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES),
25 dimethylsulfoxide (DMSO) and ethanol were purchased from Sigma-Aldrich and used without further
26 purification. RPMI 1640 with L-Glutamine, Tripsin-EDTA, Penicilin/Streptomycin, phosphate buffer
27 saline (PBS) and fetal bovine serum (FBS) were purchased from Invitrogen. Propidium iodide (PI) and
28 Ribonuclease (RNase) were purchased from Sigma-Aldrich, and Annexin-V from BD Biosciences.
29 Complexes **1** and **2** were synthesized and characterized as previously reported.¹⁵ Millipore® water was
30 used for the preparation of all aqueous solutions.

32 *4.2 Sample preparation for spectrofluorometric measurements*

34 Samples were prepared in 10 mM HEPES buffer (pH 7.40). Concentrated DNA stock solutions (*ca.* 1
35 mg/mL) were prepared by gently hydrating the solid DNA in pH 7.40 HEPES buffer, leaving it
36 overnight at 4 °C on an orbital stirrer with slow gentle shaking. DNA concentrations, expressed in

1 nucleotide units, were determined by UV spectrophotometry using the molar absorption coefficient
2 $6600 \text{ M}^{-1}\text{cm}^{-1}$ at 260 nm.³⁸ The ratio of UV absorbance at 260 and 280 nm was checked for each new
3 stock solution, and the ratio $A_{260\text{nm}}/A_{280\text{nm}} \geq 1.8$ indicating that the DNA was sufficiently free of
4 proteins.³⁴ DNA stock solutions were stored at 4 °C and used within a maximum of 4 days. Stock
5 solutions of EB and DAPI were prepared by dissolving a known amount of the probe in Millipore®
6 water and were kept in dark for sample preparation.
7 Spectroscopic measurements were carried out on individually prepared samples to ensure the same
8 pre-incubation time at $(25 \pm 1)^\circ\text{C}$ for all samples in each assay. Due to the limited solubility of
9 complexes **1** and **2** in aqueous media, DMSO was used to prepare concentrated stock solutions of each
10 complex, followed by appropriate dilution with DMSO (if needed) and 10 mM HEPES buffer (pH
11 7.40) to obtain the desired complex concentration and DMSO content, kept constant as 2% (v/v) in the
12 final samples. Dilutions were carried out immediately prior to sample preparation. Data presented are
13 the mean value \pm standard deviation of at least 3 independent assays.

14

15 *4.3. Steady-state spectrofluorometric measurements*

16

17 Fluorescence measurements were carried out on a Spex Fluoro-log 3-22/Tau 3 spectrofluorometer
18 equipped with double grating monochromators in both excitation and emission light paths from Horiba
19 Jobin Yvon at room temperature. For measurements with the probes EB and DAPI at 25.0 ± 0.1 °C
20 semi-micro quartz cells (1 cm \times 0.41 cm) from Hellma were used. Samples usually contained 3-5 μM
21 probe, and various probe:nucleotide ratios (from 1:0 to 1:50) were used. In the probe displacement
22 experiments the probe-to-nucleotide ratio was 1:4 and 1:25 for EB and DAPI, respectively, and the
23 concentrations of complexes **1** and **2** were varied at 0-64 μM after 60 min incubation in the dark. The
24 excitation wavelengths were 510 nm for EB (with 4 nm/4 nm bandwidths for excitation and emission,
25 respectively) and 365 nm for DAPI (2 nm/2 nm slit widths) and the fluorescence emission was
26 measured in the range of 380-650 nm depending on the experiment. The conditional binding constants
27 were calculated with the computer program PSEQUAD.²³ Data were also analysed by Lineweaver-
28 Burk linearization method for comparison (see **Supporting Information, section S1**).²⁷
29 Corrections for self-absorbance and inner filter effect were necessary in the steady-state fluorometric
30 experiments since the emitted light is partly absorbed by the probes and by the Ru(III) complexes.
31 Corrections were carried out according to the equation (1),

$$32 \quad I_{corr} = I_{meas} \times 10^{(A_{EX}+A_{EM})/2} \quad (1)$$

33 where I_{corr} and I_{meas} are the corrected and measured fluorescence intensities, A_{EX} and A_{EM} are the
34 absorbance values at the excitation and emission wavelengths in the samples, respectively.¹⁸ The
35 geometry of the cell was considered when executing the corrections. The spectrophotometric

1 absorption measurements were recorded on a Jasco V-560 (Tokyo, Japan) double beam
2 spectrophotometer.

4 4.4. Time-resolved fluorescence measurements

5
6 Fluorescence lifetime was measured using nanoLED N-460 for EB ($\lambda_{EM} = 590$ nm, 15 nm/ bandwidth)
7 and N-370 for DAPI ($\lambda_{EM} = 450$ nm, 2 nm bandwidth) light sources (Horiba Jobin Yvon) using time
8 correlated single-photon counting (TCSPC) mode. The resolution of the detection system was 50 ps.
9 The number of counts on the peak channel was 20000 (for DAPI samples) or 10000 (for EB samples).
10 The number of channels per curve used for the analysis was *ca.* 1000 for EB-DNA, *ca.* 400-600 for
11 DAPI and *ca.* 300 for EB alone, with a time scale of 0.05552 ns/channel for samples containing DAPI
12 and EB alone and 0.1114 ns/channel for EB plus DNA containing samples. Ludox[®] (from Sigma-
13 Aldrich) was used as the scatter to obtain the instrumental response function. The background
14 (obtained with the blank sample) was subtracted from the decay. The program TRFA data processor
15 version 1.4 (Minsk, Belarus) was used for the analysis of the experimental fluorescence decays. The
16 fluorescence intensity decay over time is described by a sum of exponentials, *i.e*

$$17 \quad I(t) = \sum_{i=1}^n \alpha_i \exp\left(\frac{-t}{\tau_i}\right) \quad (2)$$

18 where α_i and τ_i are the normalized amplitude and lifetime of component i respectively. From these
19 parameters, the fraction of emitted light by each component i can be calculated through

$$20 \quad f_i = \frac{\alpha_i \tau_i}{\sum(\alpha_i \tau_i)} \quad (3)$$

21 and changes in the quantum yield by processes affecting fluorescence lifetime are evaluated from the
22 amplitude-weighted average fluorescence lifetime, which is given by

$$23 \quad \tau_{av} = \sum \alpha_i \tau_i \quad (4)$$

24 where α_i and τ_i have the same meaning as in (2). The quality of the fit was judged from a χ^2_R value
25 close to 1.0 and a random distribution of weighted residuals.

27 4.5. Cell studies

29 4.5.1. Cell culture and cytotoxicity

30
31 The cytotoxicity of the two complexes for a 24 h incubation time was determined against the A2780
32 (human ovarian carcinoma, cisplatin sensitive) cell line. Cells were grown in Roswell Park Memorial
33 Institute medium (RPMI-1640), supplemented with 10% fetal bovine serum and 1% antibiotics in a
34 CO₂ incubator (Heraus, Germany) at 37 °C and humidified atmosphere Cell viability was determined
35 using the MTT colorimetric assay that measures the reduction of yellow 3-(4,5-dimethylthiazol-2-yl)-

1 2,5-diphenyl tetrazolium bromide (MTT) by mitochondrial succinate dehydrogenase to purple
2 formazan product.³⁵ Since reduction of MTT can only occur in metabolically active cells the level of
3 activity is a measure of the viability of the cells. For this purpose, cells were seeded in 96-well plates
4 in growth media at 20×10^3 cells/well with 200 μL of medium. After 24 h, media were replaced and
5 cells incubated with the compounds tested in aliquots of 200 μL /well. Complexes were first solubilized
6 in DMSO and then in cell medium, and added to final concentrations ranging from 2 nM to 100 μM
7 (complexes). The final concentration of DMSO in cell culture medium was lower than 1% (v/v). After
8 continuous exposure for 24 h (37 °C/5% CO_2), the medium was discarded and cells incubated with 200
9 μL of MTT solution in PBS (1.2 mM). After 3-4 h at 37 °C/5% CO_2 , the solution was removed and the
10 purple formazan crystals formed inside the cells dissolved in 200 μL DMSO. Cell viability was
11 assessed by measuring the absorbance at 570 nm in a plate spectrophotometer (PowerWave Xs, Bio-
12 Tek, USA) and calculated for each concentration tested using
13 $\% \text{ Cell viability} = [A(\text{sample})_{570} - \text{blank} / A(\text{control})_{570} - \text{blank}] \times 100$ where A_{sample} represents the
14 absorbance reading from the cells treated with the ruthenium complexes and A_{control} that from the
15 untreated cells, *blank* represents absorbance reading from wells (DMSO only). The cytotoxicity of the
16 compounds was quantified by determining the concentration needed to inhibit tumor cell growth by
17 50% (IC_{50}), based on non-linear regression analysis of dose-response data using the GraphPad Prism
18 software (vs.4.0).

19

20 4.5.2. Mechanisms of cell death: Annexin-V/PI staining

21

22 A2780 cells were treated with complexes **1** and **2** at three different concentrations, chosen as the IC_{50}
23 concentration (24 h), one value lower and one above, namely 30 μM , 50 μM and 70 μM for complex **1**
24 and 10 μM , 30 μM and 50 μM for complex **2**.

25 After treatment with **1** and **2** for 24 h, 1.5×10^6 cells (control) and 3×10^6 cells (test samples for **1** and **2**)
26 were harvested and washed twice with ice-cold PBS. Cells were then re-suspended in 10 μL PI (50
27 $\mu\text{g}/\text{mL}$) and FITC-conjugated Annexin-V (5 μL) along with binding buffer (pH 7.4 HEPES 0.1 M/1.4
28 M NaCl/25 mM CaCl_2 , diluted 1:10 with Millipore® water). After 20 min at room temperature in the
29 dark, cell staining was analyzed by fluorescence-activated cell sorting (FACS) using a Beckman
30 Coulter EPICS XL-MCL. All data was analyzed using WinMDI 2.9 software plotting the fluorescence
31 of PI over Annexin-V fluorescence.

32 4.5.3. Cell Cycle Analysis

33

34 In order to investigate the possible effect of the ruthenium compounds on cell cycle progression by
35 flow cytometry, A2780 cells were treated with **1** and **2** for 24 h. Briefly, 1.5×10^6 (control) and 3×10^6 (**1**

1 and 2) cells were harvested washed twice with PBS, fixed with 80% (v/v) cold aqueous ethanol and
2 stored overnight at 4 °C. The fixed cells were washed twice with PBS and incubated with 10 µL of
3 RNase (stock 100 µg/mL) and 10 µL of PI (15 µg/mL). Samples were incubated at 37 °C in the dark
4 and analyzed by flow cytometry (Beckman Coulter EPICS XL-MCL). The percentage of cells in
5 G1/G0, S, G2/M and sub-G1 phases was obtained by univariate models after exclusion of duplets
6 using FlowJo V10 software.

7 8 *4.5.4. Morphological analysis by TEM*

9
10 A2780 cells at approximately 70% confluence were treated with 1 and 2 at 50 µM equimolar
11 concentration for 24 h at 37 °C, and untreated cells were used as controls. After incubation cells were
12 processed following a standard procedure used before by some of us and previously reported.³⁶
13 Briefly, the culture medium was discarded and replaced by primary fixative consisting of 3%
14 glutaraldehyde in 0.1 M sodium cacodylate buffer pH 7.3. Following primary fixation for 2 h at 4 °C
15 and a wash step in cacodylate buffer, cells were then scrapped, pelleted and embedded in 2% agar for
16 further processing. Samples were further fixed for 3 h in 1% osmium tetroxide in 0.1 M sodium
17 cacodylate buffer pH 7.3. Then, samples were washed in acetate buffer (0.1 M, pH = 5.0) and fixed in
18 0.5% uranyl acetate in the same buffer for 1 h. Dehydration was carried out with increasing
19 concentrations of ethanol. After passing through propylene oxide, samples were embedded in Epon-
20 Araldite, using SPI-Pon as an Epon 812 substitute. Glass or diamond knives were used to cut thin
21 sections which were stained with 2% aqueous uranyl acetate and Reynold's lead citrate. The stained
22 sections were analyzed and photographed in a JEOL 1200-EX electron microscope.

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34 35 **Appendix. Supplementary data**

36 Supplementary data related to this article can be found online at...

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Abbreviations

ctDNA	calf thymus DNA (Deoxyribonucleic acid)
DAPI	4',6-diamidino-2-phenylindole
EB	ethidium bromide
HEPES	4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid
PI	propidium iodide

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