Luminescent Properties of Ruthenium(II) Complexes with Sterically Expansive Ligands Bound to DNA Defects

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Supporting Information

ABSTRACT: A new family of ruthenium(II) complexes with sterically expansive ligands for targeting DNA defects was prepared, and their luminescent responses to base pair mismatches and/or abasic sites were investigated. Design of the complexes sought to combine the mismatch specificity of sterically expansive metalloinsertors, such as [Rh(bpy)2(chrysi)]3+ (chrysi = chrysene-5,6-quinone diimine), and the light switch behavior of [Ru(bpy)2(dppz)]2+ (dppz = dipyrido[3,2-a:2′,3′-c]phenazine). In one approach, complexes bearing analogues of chrysi incorporating hydrogen-bonding functionality similar to dppz were synthesized. While the complexes show luminescence only at low temperatures (77 K), competition experiments with [Ru(bpy)2(dppz)]2+ at ambient temperatures reveal that the chrysi derivatives preferentially bind DNA mismatches. In another approach, various substituents were introduced onto the dppz ligand to increase its steric bulk for mismatch binding while maintaining planarity. Steady state luminescence and luminescence lifetime measurements reveal that these dppz derivative complexes behave as DNA “light switches” but that the selectivity in binding and luminescence with mismatched/abasic versus well-matched DNA is not high. In all cases, luminescence depends sensitively upon structural perturbations to the dppz ligand.

INTRODUCTION

Deficiencies in the DNA mismatch repair pathway, which corrects DNA defects including base pair mismatches and single-base bulges, are associated with several forms of human cancer.1–3 Our laboratory has been interested in developing octahedral metal complexes that target DNA defects.4–8 Since deficiencies in mismatch repair necessarily lead to an increased frequency of DNA defects within the cell, targeting these defects with small metal complexes provides a new strategy in designing new cancer diagnostics and chemotherapeutics.

We reported octahedral rhodium complexes that specifically bind single-base mismatch sites in DNA and, upon photoactivation, cleave the DNA backbone.4–6 The Rh complex [Rh(bpy)2(chrysi)]3+ (Figure 1) recognizes 80% of mismatched sites in all sequence contexts, and selectively targets a single-base mismatch in a 2725 base pair DNA plasmid.5 Structural studies of the rhodium complex bound to mismatched DNA reveal that it inserts into the destabilized mismatch site from the minor groove with complete ejection of the mismatched base pair into the major groove and no change in base pair rise.9 Thus, the [Rh(bpy)2(chrysi)]3+ complex is a metalloinsertor rather than a classical metallointercalator. Furthermore, we demonstrated that abasic sites and single-base bulges are also recognized by the [Rh(bpy)2(chrysi)]3+ complex.10 The source of defect-specific binding is the chrysi-inserting ligand. Structural properties of the chrysi ligand (e.g., steric bulk and planarity) make it too wide to intercalate into well-matched duplex DNA, but it is well suited to bind DNA mismatches in the minor groove. The binding affinity of this rhodium complex to each mismatch is correlated with the thermodynamic stability of the mismatch.

Another effort in our laboratory has been constructing metal complexes as luminescent probes for DNA.8 Complex [Ru(bpy)2(dppz)]2+ (Figure 1, bpy = 2,2′-bipyridine) exhibits light switch behavior in the presence of well-matched duplex DNA, but it is well suited to bind to destabilized sites in DNA. The binding affinity of this complex to each mismatch is correlated with the thermodynamic stability of the mismatch.

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Figure 1. Chemical structures of [Rh(bpy)2(chrysi)]3+ and [Ru(bpy)2(dppz)]2+.
that the \([\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+}\) complex intercalates into DNA by one of two binding modes: side-on or perpendicular.\(^8\) In the perpendicular mode, the Ru–dppz axis lies along the DNA dyad axis, protecting the phenazine fragment from water. In the side-on mode, the Ru–dppz axis lies along the long axis of the base stack, partially exposing the ligand to solvent. Thus, complexes bound in the better-protected perpendicular intercalation mode show greater luminescent enhancement upon binding to DNA and have longer excited state lifetimes than those in the side-on mode. Recent crystal structures have shown intercalation of dppz complexes into the DNA duplex from the minor groove,\(^14\) though NMR studies have pointed to intercalation from the major groove;\(^13\) the energetic differences between major and minor groove orientations must be small.

How might the luminescent properties of the \([\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+}\) complex change in the presence of DNA containing a defect? Previously, we have shown that \([\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+}\), while showing luminescence bound to duplex DNA, shows still greater luminescence in the presence of a mismatch.\(^16\) More recent efforts to understand how mismatches affect \([\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+}\) binding have led to determination of an atomic-resolution crystal structure of \(\Delta([\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+}\) bound to a mismatch-containing oligonucleotide.\(^14\) The structure reveals that, like its rhodium counterpart, the ruthenium complex also recognizes the destabilized mismatches through metalloinsertion: the dppz ligand inserts into the base stack from the minor groove, extruding the mispaired bases out of the helix. Thus, it is conceivable that by combining the structural features of \([\text{Rh}(\text{bpy})_2(\text{chrysi})]^{3+}\) (for mismatch-specific recognition) and \([\text{Ru}(\text{bpy})_2(\text{dppz})]^{2+}\) (for luminescence) it might be possible to develop an improved luminescent reporter for DNA defects. Here, we investigated two approaches for enhancing the mismatch specificity: (i) replacement of the dppz ligand with sterically expansive inserting ligands, such as chrysi, and (ii) incorporation of various functionalities onto the dppz ligand to increase the steric bulk. We report the preparation of a new family of \([\text{Ru}(\text{bpy})_2(\text{L})]^{2+}\) derivatives (Figure 2) and the luminescent behavior of each derivative in the presence of duplex DNA with and without defects.

**EXPERIMENTAL PROCEDURES**

**Materials and Methods.** All reagents and solvents were purchased from commercial suppliers and used without further purification. Chrysene-5,6-quinone and benzo[a]phenazine-5,6-quinone were synthesized according to literature procedures.\(^17\) Ru complexes \([\text{Ru}(\text{bpy})_2\text{Cl}_2]\) \([\text{Ru}(\text{bpy})_2(\text{dppz})]\) \([X]=\text{PF}_6 \) or \(\text{Cl}\) were prepared by previously
described procedures. UV–vis spectra were recorded on a Beckman DU 7400 UV–visible spectrophotometer (Beckman Coulter). Oligonucleotides 3′-GAC CAG CTT ACC CCT AGA TAA GCCG-3′ and 3′-CTG GTC GAA TAG TXG GGA TGT ATT CGC-3′ [X = G (M), C (MM), or R (AB, R denotes a tetrahydrofurany1 albacite)] were synthesized on an ABI 3400 DNA synthesizer (Applied Biosystems) or purchased from Integrated DNA Technologies and purified as previously reported.20

Synthesis. 1,2-Dihydrobenz[c]acridine (1). This compound was synthesized according to the literature procedure21 and purified by SiO2 column chromatography (9:1 hexane/EtOAc). Yield: 57%. 1H NMR (CDCl3, 300 MHz, δ (ppm)): 8.60 (d, J = 7.4 Hz, 1H), 8.17 (d, J = 8.3 Hz, 1H), 7.95 (s, 1H), 7.77 (d, J = 8.1 Hz, 1H), 7.67 (t, J = 7.6 Hz, 1H), 7.59–7.33 (m, 3H), 7.33–7.25 (m, 1H), 3.15 (t, J = 6.5 Hz, 2H), 3.03 (t, J = 6.5 Hz, 2H). ESI+(MS): [M+H]+ calc 232.1, found 232.4.

5,6-Dihydronaphthalene-1,2-diamine (2). A suspension of α-tetralone (0.39 g, 2.7 mmol), (2-aminopyridin-3-yl)methanol22 or 5,6-Naphtho[1,2-b][1,8]naphthyridine (naphthi) Quinone (2.0 mmol) in 1:1 acetic acid/acetic anhydride (10 mL) was added dropwise, and the reaction mixture was stirred at room temperature for 8 days, during which time a yellow solid precipitated from the reaction mixture. Yellow precipitate was filtered, washed with H2O, and dried. Inorganic Chemistry

A suspension of Na2Cr2O7 (0.90 g, 3.0 mmol) in 1:1 acetic acid/acetic anhydride (10 mL) was heated at reflux under an Ar atmosphere for 2 h. The red suspension was cooled to room temperature, filtered through Celite, and poured into a saturated NH4Cl solution (20 mL). This was extracted with EtOAc (3 × 20 mL), and the combined organic layers were dried over Na2SO4 and filtered, and solvent was removed. Crude material was purified by SiO2 column chromatography (1:1 hexane/EtOAc) to give 2 as a pale red solid (0.37 g, 61%). 1H NMR (CDCl3, 300 MHz, δ (ppm)): 9.06 (d, J = 4.2 Hz, 1H), 1.84–8.73 (m, 3H), 8.20–8.05 (m, 1H), 7.95 (s, 1H), 7.50–7.35 (m, 3H), 7.33–7.23 (m, 1H), 3.29–3.09 (m, 2H), 3.10–2.96 (m, 2H). ESI+(MS): [M+H]+ calc 233.1, found 233.4.

Ligands were synthesized according to the literature procedures.8,18,19 UV-vis spectra were recorded on a Varian Cary 50 Bio spectrophotometer. The reaction mixture was stirred for 10 min at 0 °C and allowed to warm slowly to room temperature. The reaction was quenched by addition of water after stirring for 4 h at room temperature, and the solution was filtered. Filtrate was extracted with CH2Cl2 (three times). After removal of CH2Cl2, the crude product was purified by SiO2 column chromatography with a solvent gradient (50%–50% H2O/CH2Cl2 to 100% CH2Cl2) to obtain the desired product 7 (153 mg, 0.98 mmol, 89%). 1H NMR (CDCl3, 300 MHz, δ (ppm)): 6.82 (s, 2H), 3.98 (br, s, 4H), 3.44 (s, 2H). HREI (m/z) for M+ calc 156.0687, found 156.0687.

5,6-Dihydronaphthalene-1,2-diamine (2) was deoxygenated and stirred at room temperature for 4 h before quenching the reaction with a couple of drops of 1 M HCl. The reaction mixture was diluted with water (10 mL), excess NH4PF6 (s) was added, and the solution was concentrated in vacuo. Precipitates were collected, washed with cold ethanol (three times, 20 mL), dried under vacuum, and used for preparation of the Ru complexes without further purification.


Br2dppe. Yield: 78%. ESI+(MS): [M + H]+ calc 490.9, found 491.0.


[Ru(bpy)2(NH2)2(PF6)2]. A solution of Ru(bpy)2Cl2 (0.47 g, 0.9 mmol) and NH4OH/MeOH (10 mL) was heated at 60 °C for 4 h. The reaction mixture was cooled to room temperature, and the solvent was removed in vacuo. Residue was redissolved in MeOH and precipitated with Et2O. Solid was collected and redissolved in water (10 mL), and excess NH4PF6 (s) was added. The suspension was stirred in the absence of light for 1 h and filtered, and the red precipitate was washed with cold water (15 mL) and Et2O (3 × 10 mL) and dried under vacuum to give [Ru(bpy)2(NH2)2(PF6)2] (0.58 g, 0.8 mmol, 88%). 1H NMR (CD3OD, 300 MHz, δ (ppm)): 9.18 (d, J = 5.6 Hz, 2H), 8.64 (d, J = 8.1 Hz, 2H), 8.46 (d, J = 8.1 Hz, 2H), 8.20 (t, J = 7.9 Hz, 2H), 7.82 (t, J = 7.7 Hz, 4H), 7.65 (d, J = 5.7 Hz, 2H), 7.27–7.08 (m, 8H), 2.89 (s, 6H).

[Ru(bpy)(L)Cl] Complexes. Method 1. A suspension of [Ru(bpy)(L)Cl](PF6) (0.043 mmol) and NaH (0.08 mmol) in dry MeCN (10 mL) was deoxygenated and stirred under an Ar atmosphere for 30 min. Ligand (0.048 mmol) was added; the mixture was deoxygenated and stirred at room temperature for 4 h before quenching the reaction with a couple of drops of 1 M HCl. The reaction mixture was diluted with water (10 mL), excess NH4PF6 (s) was added, and the solution was concentrated in vacuo. Precipitates were collected and washed with water (2 × 10 mL) and then converted to the soluble chloride salt by anion exchange chromatography (Sephadex QAE). Resides were purified on a Waters C18 sep-pak cartridge and by preparative HPLC using a gradient of H2O (with 0.1% TFA) to CH2CN over 60 min. Complex was converted to the chloride salt by anion exchange chromatography (Sephadex QAE).

[Ru(bpy)(L)Cl][X] (12). X = TFA or Cl. Yield: 18%. 1H NMR (CD3CN, 300 MHz, δ (ppm)): 12.91 (s, 1H), 12.42 (s, 1H), 8.58–8.44 (m, 4H), 8.39–8.30 (m, 1H), 8.24 (d, J = 8.8 Hz, 1H), 8.19–8.03 (m, 5H), 7.94 (d, J = 5.6 Hz, 2H), 7.82 (t, J = 8.3 Hz, 1H), 7.73–7.59 (m, 5H), 7.54–7.39 (m, 4H). ESI+(MS): [M−H]+ calc 669.1, found 669.1; [M+H]+ calc 335.1, found 334.6. UV–vis in H2O, λmax/nm (ε × 104 M−1 cm−1): 225 (11.5), 282 (14.5), 550 (6.4).

[Ru(bpy)(L)Cl][X] (12). X = TFA or Cl. Yield: 22%. 1H NMR (CD3CN, 300 MHz, δ (ppm)): 14.06 (s, 1H), 13.07 (s, 1H), 9.75 (s, 1H), 9.12 (d, J = 8.1 Hz, J = 1.4 Hz, 1H), 8.60–8.44 (m, 5H), 8.19 (d, J = 8.6 Hz, 1H), 8.13–8.00 (m, 7H), 7.97–7.84 (m, 1H), 7.83 (t, J = 7.6 Hz, 1H), 7.76–7.59 (m, 4H), 7.54–7.37 (m, 4H). ESI+(MS): [M−H]+ calc 670.1, found 670.3; [M+H]+ calc 335.6, found 346.4.
Method 2. Ethylene glycol (7 mL) was added to a mixture of Ru(bpy)$_2$Cl$_2$ (0.069 mmol) and the dppz derivatives (0.069 mmol), and the solution was heated to 130 °C for 12 h (Scheme 3). After cooling the reaction mixture to room temperature, the solution was diluted with water (7 mL) followed by addition of excess NH$_4$PF$_6$ (s). Orange precipitates were collected, washed with water (3 × 5 mL), and dried under vacuum. Complexes were reccrystallized by addition of Et$_2$O into their CH$_3$CN solutions at room temperature and afterward converted to the soluble Cl salt by anion exchange chromatography (Sephadex QAE). They were further purified by preparative HPLC using a gradient of H$_2$O (with 0.1% TFA) to CH$_3$CN (with 0.1% TFA) over 30 min.

[Ru(bpy)$_2$(dppz)]Cl$_2$ (16, X = PF$_6$ or Cl). Yield: 63%. $^1$H NMR (CD$_3$CN, 500 MHz, δ (ppm)): 9.76–9.68 (m, 2H), 9.06 (m, 1H), 8.70 (m, 1H), 8.59–8.54 (m, 4H), 8.49 (1H, m), 8.22 (s, br, 2H), 8.14 (m, 2H), 8.05 (m, 2H), 7.94 (m, 2H), 7.79 (2H, m), 7.49 (m, 2H), 7.30 (m, 2H), ESI(+)MS (m/z): [M – 2(PF$_6$)]$^{2+}$ calcd 370.1, found 370.1. [M – PF$_6$]$^{+}$ calcd 885.1, found 885.2. UV–vis in H$_2$O, $\lambda_{\text{abs}}$/nm ($\varepsilon \times 10^4$/M$^{-1}$ cm$^{-1}$): 284 (10), 362 (1.9), 378 (1.9), 444 (1.6).

Scheme 3
Ruthenium complexes were synthesized in two steps from \( \alpha \)-tetralone by a Friedländer synthesis and subsequent oxidation with sodium dichromate (Scheme 1).\(^{21,23}\) Ruthenium complexes 12–14 were synthesized by reacting [Ru(bpy)\(_2\)(NH\(_3\))\(_2\)]PF\(_6\) with the appropriate quinone ligand in the presence of sodium hydride at ambient temperature (Scheme 3). Complexes 12–14 were isolated in 4–20% yield following purification and anion exchange to the chloride salt. Unexpectedly, [Ru(bpy)\(_2\)(phenzhi)]\(^{2+}\) (Figure 2) was not the major product from the reaction with benzo[\(\alpha\)]phenazine-5,6-quinone and was only isolated in trace amounts, as characterized by HPLC and mass spectrometry. We propose that [Ru(bpy)\(_2\)(iqi)]\(^{2+}\) (Figure 2) is the major product of the reaction as the \( ^1H \) NMR spectrum of the isolated complex only contains one imine proton signal, and the mass spectrum is consistent with loss of carbon monoxide. It has been reported that a related ligand, 1,10-phenanthroline-5,6-dione, loses carbon monoxide under basic conditions.\(^{27}\) Similar ligand decomposition in the presence of sodium hydride during the course of the reaction could account for the low yields of complexes 12–14\(^{28}\) and formation of byproduct 22. Following the loss of carbon monoxide from benzo[\(\alpha\)]phenazine-5,6-quinone, unlike the other ligands, it is still possible to coordinate to the metal center in a bidentate fashion through the phenazine nitrogen and imine formed from the remaining quinone.

Ru complexes [Ru(bpy)\(_2\)(dpzp)]\(^{2+}\) (15) and [Ru(bpy)\(_2\)(tactp)]\(^{2+}\) (18) were prepared by previously published methods.\(^{8,15}\) For the ligands dpzp, dpae, dpzn, and Br\(_2\)dpzn (Scheme 2), the amine moieties were condensed with 1,10-phenanthroline-5,6-dione and 5,6-diamino-1,10-phenanthroline\(^{26}\) in ethanol.\(^{80}\) Complexes 16, 17, and 19–21 were obtained by refluxing a solution containing Ru(bpy)\(_2\)Cl\(_2\) and 1 equiv of the corresponding dpzp derivative in ethylene glycol at 130 °C for 8 h, as depicted in Scheme 3. Addition of excess NH\(_4\)PF\(_6\) (s) into the reaction solution of 1:1 ethylene glycol:water allowed isolation of the Ru complexes as PF\(_6\) salts, followed by purification via recrystallization or column chromatography. All complexes show the ligand emissive properties of the parent complex [Ru(bpy)\(_2\)(dppz)]\(^{2+}\) when compared to the complex [Ru(bpy)\(_2\)(dpzp)]\(^{2+}\) (15) and [Ru(bpy)\(_2\)(tactp)]\(^{2+}\) (18).
characteristic MLCT transition absorption in the visible region at \( \sim 440 \) nm.

**Luminescent Characteristics of the Ru Complexes with and without DNA.** While complexes 12−14 (Scheme 3 and Figure 2) are not luminescent at ambient temperature in various solvents (dry MeCN, Tris buffer, and 10 M LiCl), \([\text{Ru(bpy)}_2(\text{chrysi})]^2^+\) (12) and \([\text{Ru(bpy)}_2(\text{naphthi})]^2^+\) (14) are luminescent in the absence of DNA at 77 K in a 10 M LiCl glass (Figure 3). In the presence of DNA, there is no luminescence differential between mismatched and matched DNA for both complexes. \([\text{Ru(bpy)}_2(\text{acri})]^2^+\) (13) is not luminescent at 77 K in dry MeCN nor in the presence or absence of DNA in 10 M LiCl upon excitation at 440 nm. The DNA “light switch” complex \([\text{Ru(bpy)}_2(\text{dpdz})]^2^+\) (15) is also luminescent at 77 K in the absence of DNA (Figure S2, Supporting Information). Therefore, it is not possible to determine the mismatch specificity or light switch behavior of the complexes from low-temperature experiments.

At ambient temperature, \([\text{Ru(bpy)}_2(\text{chrysi})]^2^+\) (12), \([\text{Ru(bpy)}_2(\text{acri})]^2^+\) (13), and \([\text{Ru(bpy)}_2(\text{naphthi})]^2^+\) (14) are not luminescent in the absence or presence of DNA. Therefore, competition experiments were carried out with the “light switch” complex \([\text{Ru(bpy)}_2(\text{dpdz})]^2^+\) (15) in order to investigate the room-temperature mismatch specificity of the complexes. Control experiments reveal that complexes 12−14 do not quench the luminescence of 15 in the absence of DNA in dry MeCN (Figure S3, Supporting Information). Luminescence of 1:1 DNA/15 mixtures in aqueous buffer was measured upon addition of up to 10 equiv of complexes 12−14. A decrease in the luminescence is expected if the complex displaces 15, as 15 is luminescent only when bound to DNA. For all three complexes 12−14, there was a small decrease in the luminescence of 15 in the presence of matched DNA with the largest decrease observed with 12 (33% after 1 equiv). In contrast, there was a much more significant luminescence decrease, following the order 12 (55% after 1 equiv) > 13 (35%) > 14 (21%), in the presence of mismatched DNA than for matched DNA (Figure 4). While it was not possible to directly detect the luminescence of the complexes at room temperature, competition studies provide evidence that the

Figure 3. Emission spectra of \([\text{Ru(bpy)}_2(\text{chrysi})]^2^+\) (12) and \([\text{Ru(bpy)}_2(\text{naphthi})]^2^+\) (14) in the presence and absence of DNA at 77 K and room temperature (10 M LiCl, \( \lambda_{ex} = 440 \) nm). [Ru] = 5 \( \mu M \), [DNA] = 0 or 5 \( \mu M \). DNA sequences are shown at the top of the figure.

Figure 4. Plots of the integrated emission intensity of 0.1 \( \mu M \) \([\text{Ru(bpy)}_2(\text{dpdz})]^2^+\) (15) and 0.1 \( \mu M \) matched (M) or mismatched (MM) DNA with up to 10 equiv of \([\text{Ru(bpy)}_2(\text{L})]^2^+\), where L is chrysi (12), acri (13), or naphthi (14) (5 mM Tris, pH 7.5, 50 mM NaCl, \( \lambda_{ex} = 440 \) nm). See Figure 3 for DNA sequences.
complexes preferentially bind to DNA mismatches through their sterically expansive inserting ligands.

Restoration of luminescence at low temperature for [Ru(bpy)_2(chrysi)]^{2+} (12) and [Ru(bpy)_2(naphthi)]^{2+} (14) suggests that nonradiative solvent relaxation, most likely through the exchangeable imino protons, is responsible for the loss of luminescence at ambient temperature. Preliminary studies have revealed that methylation of the imino protons of 12 restores luminescence at ambient temperature. However, synthetic difficulties and decomposition of the methylated complex limited further studies.

Unlike complexes 12−14, the [Ru(bpy)_2(dppz)]^{2+} derivatives (15−21, Figure 2) all show enhanced luminescence with matched (M), mismatched (MM), and abasic (AB) DNA versus in the absence of DNA in aqueous buffer, indicating that the conserved phenazine portion retains some of the photophysical properties of the parent complex 15 (Figure 5). However, complexes having extensively \( \pi \)-conjugated ligands (18−21) emit luminescence with detectable intensity in the absence of DNA, possibly due to partial shielding of one of the phenazine nitrogens from the solvent by the added steric bulk.

Similar to 15, the [Ru(bpy)_2(dppa)]^{2+} complex (16) exhibits luminescence enhancement in the presence of MM and AB DNA, relative to well-matched DNA, although its binding affinity is lower than that of 15 (Figure 5 and Table S1, Supporting Information). Attachment of a benzo group to the end of the phenazine moiety increases the length of the dppz-type ligand (dppn, Scheme 2). We assume binding to the DNAs resembles that of the dppz complex. Significantly, [Ru(bpy)_2(dppn)]^{2+} (19) exhibits an increase in luminescence with MM over M and AB DNA (Figure 5). This may be because the extended dppn ligand of 19, when bound to well-matched DNA from the major groove, offers less protection to the phenazine nitrogens from solvent water molecules than does dppz. However, for dppn, binding to mismatches likely occurs from the minor groove as it does for dppz. The minor groove is deep and narrow, allowing deeper intercalation of the Ru complex and consequently better protection of the ligand from solvent water.

Introduction of the expansive dppz derivatives dppae and tactp, employing acetylene and chrysene functionalities, respectively, to increase the width of the ligand, eliminates the luminescence differential with DNA defects (Figure 5).
observed for these dppz derivatives might be due to di-
binding to DNA mismatches and the lack of speci-
ferences. In the [Ru(bpy)2(dppz)]2+ derivatives
in the proximity of the steric bulk to the metal center in the
bromine.29 The increased luminescence of
20
MM, and AB DNA, however (Figure 5). The weak
conformational
well removed from the metal center, allowing considerable
previous study comparing the binding of [Rh(bpy)2(chrysi)]3+
octahedral center. This notion echoes the conclusions from a
its binding geometry is constrained to that imposed by the
complex, the chrysi ligand is much closer to the metal center, so
12
changes are due to displacement of [Ru-
15
from the mismatch
DNA.30
Greater protection of the phenazine moiety from water via
intercalation leads to longer emission lifetimes for the two
binding modes (side-on and perpendicular), a higher relative
population of the longer-lived species, or both.

The lifetimes do not change within experimental error in the
presence of matched DNA upon addition of up to 2 equiv of
complexes 12–14, although the relative population of the
longer-lived species decreases (Table 1). This is consistent with
a change in binding mode from perpendicular to side-on and
suggests that the small luminescence decrease observed in the
luminescence competition experiments is largely due to a
binding mode change rather than displacement of [Ru-
(bpy)2(dppz)]2+ (15) from matched DNA. In the absence of a competing complex, both lifetimes are greater for a 1:1 mixture of [Ru(bpy)2(dppz)]2+ (15) and mismatched DNA compared to well-matched DNA due to greater protection of the phenazine moiety at the mismatch (Table 1). After addition of 2 equiv of complexes 12–14, both excited state lifetimes decrease and approach each of the lifetimes of 15 observed in the presence of matched DNA. The magnitude of the decrease agrees with the steady state
luminescence competition experiments where the largest
decrease is observed with 12 followed by 13 and 14. In addition to the lifetime changes, the relative population of the perpendicular state decreases. These results contrast those from analogous experiments with matched DNA, suggesting that the changes are due to displacement of 15 from the mismatch rather than a binding mode change. This is further evidence that the complexes bind preferentially to mismatched DNA over matched DNA.

For the [Ru(bpy)2(dppz)]2+ derivatives (15–21, Figure 2), the binding preferences of the complexes in 2:1 DNA/Ru
mixtures could be probed directly at ambient temperature. As

Table 1. Luminescence Decay Parameters for Ru(bpy)2(dppz)2+ in Competition Experiments with [Ru(bpy)2(chrysi)]2+ (12), [Ru(bpy)2(acri)]2+ (13), and [Ru(bpy)2(naphthi)]2+ (14) in the Presence of Matched (M) and Mismatched (MM) DNAa

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<td>170 (48)</td>
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a$\lambda_{ex} = 470\,\text{nm}, \lambda_{em} = 610\,\text{nm}, 5\,\mu\text{M [Ru(bpy)2(dppz)]2+}, and 5\,\mu\text{M [DNA]} (5\,\text{mM Tris, pH 7.5, 50 mM NaCl}). Matched (M) and mismatched (MM) oligonucleotides shown in Figure 3 were used. Emission decays were fit to the biexponential function $Y(t) = Y_o + A_1 \exp(-t/\tau_1) + A_2 \exp(-t/\tau_2)$ to give the lifetimes $\tau_1$ and $\tau_2$. Relative contribution of each lifetime to the overall decay is indicated in parentheses. bError is estimated to be ±10%.
shown in Table 2, 15 displays a higher relative abundance bound in the perpendicular mode in the presence of DNA

<table>
<thead>
<tr>
<th>complex (ligand)</th>
<th>no DNA</th>
<th>in the presence of DNA</th>
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<tbody>
<tr>
<td>M</td>
<td>1210 (100)</td>
<td>M [Ru] 1120 (100)</td>
</tr>
<tr>
<td>MM</td>
<td>1180 (100)</td>
<td>M [Ru] 1120 (100)</td>
</tr>
<tr>
<td>AB</td>
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<td>M [Ru] 1120 (100)</td>
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<tr>
<td>19 (dppn)</td>
<td>170</td>
<td>M 62 (51)</td>
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<td></td>
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<td>MM 200 (49)</td>
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<tr>
<td></td>
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<td>AB 35 (58)</td>
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<tr>
<td>20 (Br,dppn)</td>
<td>200</td>
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<td></td>
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<tr>
<td></td>
<td></td>
<td>AB 82 (54)</td>
</tr>
<tr>
<td>21 (pyrene-phen)</td>
<td>190</td>
<td>M 930 (100)</td>
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<td>MM 880 (100)</td>
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<td></td>
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<td>AB 810 (100)</td>
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</table>

Table 2. Luminescence Decay Parameters for the [Ru(bpy)_2(dpz)]^{2+} Complexes in the Presence of Matched (M), Mismatched (MM), and Abasic (AB) DNA

Following the order AB > MM > M. This supports the observation from steady state measurements that the luminescence intensity of 15 is present in the following order: AB > MM > M (Figure 5). Likewise, in the case of [Ru(bpy)_2(dpzp)]^{2+} (16), longer lifetimes correspond to the perpendicular mode (AB > MM > M) are observed, which can explain the trend in the steady state luminescence observed for 15. Longer lifetime measurements show a lower relative yield of long-lived species for 17 and shorter lifetimes for 20 in the presence of DNA-containing defects. These results are consistent with the slight decrease in steady state luminescence observed for 17 and 20 when combined with DNA-containing defects (Figure 5).

SUMMARY AND IMPLICATIONS

Taking inspiration from the mismatch specificity of the sterically expansive chrysi ligand and light switch behavior of [Ru(bpy)_2(dpzp)]^{2+}, a family of Ru(II) complexes was prepared in an attempt to specifically target DNA defects and indicate their presence by the appearance of luminescence. In one design strategy, complexes containing sterically expansive ligands, such as chrysi and analogues with hydrogen-bonding functionality similar to dpzp, were synthesized. Incorporation of the sterically expansive ligand appears to improve the mismatch specificity; however, the complexes are nonluminescent at room temperature. The exchangeable imino protons most likely quench luminescence at room temperature through solvent relaxation as luminescence is restored at 77 K for [Ru(bpy)_2(chrysi)]^{2+} (12) and [Ru(bpy)_2(naphthi)]^{2+} (14). Despite their mismatch specificity, the lack of luminescence signals at room temperature limits the use of these complexes as probes. Light switch behavior cannot be exploited at low temperature to improve mismatch discrimination, as the known “light switch” complex [Ru(bpy)_2(dpzp)]^{2+} (15) shows luminescence at 77 K even in the absence of DNA. Ru complexes containing dpzp derivatives (15–21) were also synthesized with the goal of increasing their steric bulk through incorporation of various functionalities on the ligand framework. Their luminescence responses, however, were not substantially enhanced compared with the parent complex 15 in the presence of DNA with a mismatched or abasic site. Nevertheless, the narrow ligands (dpza and dppn) show differential luminescence behavior in the presence of a DNA defect. Widening the dpzp ligand using a simple group such as acetylene or Br does not enhance targeting to destabilized regions of DNA. Furthermore, Ru complexes with the extensively π-conjugated ligands (tactp and pyrene-phen) are likely to aggregate and are therefore unable to report mismatches via luminescence output.

These observations are consistent with a recent crystal structure in which the dpzp ligand of Δ-[Ru(bpy)_2(dpzp)]^{2+} inserts deeply from the minor groove at mismatched sites. In the crystal structure, the central ring of the phenazine portion of dpzp stacks with the DNA bases, affording the nitrogen atoms of the phenazine moiety maximum protection from solvent water. Considering the structural resemblance between dpza and dpzp, complexes of the two ligands are likely to bind DNA in the same manner. The similarity in mismatch discrimination between [Ru(bpy)_2(dpzp)]^{2+} (16) and [Ru(bpy)_2(dpzp)]^{2+} (15) is therefore not surprising. Extension of dpzp to form dppn alters the equilibrium binding geometry of the complex. The relatively high mismatch discrimination of the dppn complex may therefore indicate stronger binding and more effective protection from solvent at mismatched sites. Similarly, widening the terminal phenazine ring of dpzp to form tactp or pyrene-phen may permit resulting in stacking between the DNA bases and these wider, extended structures, decreasing the degree of protection experienced by the nitrogen atoms of the phenazine ring. Presumably, these modifications are also responsible for aggregation of the complexes. Consequently,
the phenazine nitrogens in tactp and pyrene-phen may be similarly accessible to solvent whether the complex is dimerized, intercalated, or inserted; hence, the tactp and pyrene-phen complexes exhibit similar luminescence profiles with all three types of DNA.

Of the two approaches investigated here, increasing the steric bulk of the dppz framework appeared to be more successful, as all of the derivative complexes retained “light switch” behavior. Although the structural modifications investigated here are not sufficient by themselves to enhance the luminescence differential between matched and defective DNA, future efforts to improve the mismatch specificity of [Ru(bpy)2(dpz)]2+ should perhaps shift away from appending steric bulk to the distal portion of the dppz ligand. The [Ru(bpy)2(dpz)]2+ scaffold, thus, may be able to withstand the even bolder modifications necessary to achieve mismatch-specific luminescence.

ASSOCIATED CONTENT

Supporting Information
Chemical structure and characterization information of [Ru(bpy)2(iqi)]2+ (22), Tables S1 and S2, and Figures S1–S3. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes
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REFERENCES