

Reduced and Excited States of (bpym)[PtCl₂]_n (bpym = 2,2'-Bipyrimidine; n = 1, 2): Experiments and DFT Calculations

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The complexes (bpym)PtCl₂ (**1**) and the new (μ-bpym)[PtCl₂]₂ (**2**), bpym = 2,2'-bipyrimidine, were synthesized and, in the case of **1**, crystallized in solvent-free form for X-ray diffraction. The molecules **1** exhibit two different kinds of stacking motifs in the crystal with an interstack CH–N interaction. Complexes **1** and **2** were found to be sufficiently soluble for cyclic voltammetry, spectroscopy (absorption and emission), and spectroelectrochemical studies (UV–vis, EPR). As a result of single or double coordination of the strongly σ-accepting [PtCl₂] fragment to bpym, the paramagnetic anions **1**^{•−} and **2**^{•−} and the dianions **1**^{2−} and **2**^{2−} could be reversibly generated, despite the presence of metal–halide bonds. DFT calculations of A(195Pt) and g tensor components confirm that the singly occupied MOs of the monoanionic species have mainly π*(bpym) character with nonnegligible platinum d orbital participation. The assignments of the electronic absorption and emission and resonance Raman spectra for both complexes are supported by DFT calculations.

Introduction

Although dichloroplatinum(II) complexes of α-diimine ligands such as 2,2'-bipyridine (bpy) or 2,2'-bipyrimidine (bpym) have been known now for some time,^{1,2} these and related compounds have received increasing attention recently because of various special features:

(i) The complex photophysical behavior³ of such materials such as variable temperature emission in the solid state or in solution is related to their ability to form aggregates (via stacking) between the largely planar molecules.^{4,5} Differences in crystallization are also responsible for distinctly different color forms of e.g. yellow and red (bpy)PtCl₂.⁵

(ii) Chemical or electrochemical reduction produces paramagnetic complexes which are better formulated as Pt^{II}/radical anion compounds and not as Pt^I species.^{6,7} EPR and spectroelectrochemical techniques were employed to characterize such one-electron reduced systems.^{6,7}

(iii) Chloroplatinum compounds with ancillary nitrogen ligands are generally interesting for their physiological

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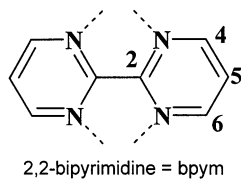
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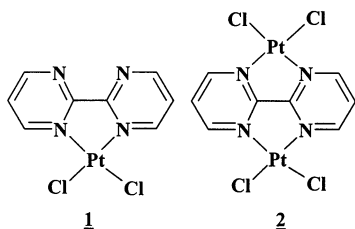
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behavior inside cells and toward the DNA in continuing efforts for the advancement of metal-based tumor therapy.^{8,9}

(iv) Platinum(II) compounds have a potential as catalysts and catalyst precursors for C–H bond activation.^{10,11} It was reported that (bpym)PtCl₂ (**1**), bpym = 2,2'-bipyrimidine, is a robust catalyst precursor for efficient methane oxidation to methanol.^{11a}



Whereas **1** has been studied since 1978² and two crystal structure analyses of solvated (bpym)PtCl₂ have appeared more recently,^{4,12} it was reported that the obvious extension, the dinuclear compound (μ -bpym)[PtCl₂]₂, could not be obtained.² To our knowledge, there has not yet been any report of solution properties of a bis(dichloroplatinum(II)) complex with a π -conjugated bridging ligand. Drawing on previous experience with organometallic analogues (μ -bpym)[PtR_n]₂ (R = Me, $n = 2, 4$;^{13,14} R = Mes,¹³ p -Tol, Ad¹⁵ $n = 2$), we have now succeeded in preparing (μ -bpym)-[PtCl₂]₂ (**2**). This report contains information on the reduced and excited states of **1** and **2**, based on electrochemical, EPR, UV–vis–spectroelectrochemical, emission, and resonance Raman (rR) studies. In addition, solvent-free **1** could be crystallized for structure analysis; the structural and spectroscopic data were interpreted using results from DFT calculations.



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Further reasons for synthesizing dinuclear **2** and for studying its electrochemical and spectroscopic properties include a comparison with other dinuclear compounds of bridging bpym with complex fragments containing a platinum metal halide function such as (η^6 -C₆R₆)ClO_s⁺¹⁶ or (η^5 -C₅R₅)CIM⁺ (M = Ir, Rh).¹⁷ A related 5d metal halide fragment is (OC)₃ClRe, which has also been studied in bpym-bridged dinuclear form.¹⁸ Additionally, the combination between the conjugated bridging ligand bpym with its relatively high-lying π^* MO¹⁹ and two electron-poor d⁸ metal complex fragments [PtCl₂] was expected to produce on reduction a singly occupied MO of rather mixed metal/ligand character. For a symmetrical dinuclear compound a mixed-valent^{13,20,21} (d⁸/d⁹) formulation could thus contribute to the description of the paramagnetic state as evident from EPR²² or spectroelectrochemical studies.²³ Finally, the electronic structures of related bpym-bridged dinuclear compounds such as (μ -bpym)[M(CO)₄]₂¹⁹ (M = Mo⁰, W⁰; d⁶ configuration) have been studied by resonance Raman and emission spectroscopy.²⁴

Experimental Section

Instrumentation. EPR spectra were recorded in the X band on a Bruker System ESP 300 equipped with a Bruker ER035M gaussmeter and a HP 5350B microwave counter. Spectra simulation was performed using in-house software.²⁵ ¹H NMR spectra were taken on a Bruker AC 250 spectrometer; infrared spectra were obtained using a Perkin-Elmer Paragon 1000 PC instrument. Resonance Raman spectra were obtained for KNO₃ pellets by excitation with the lines of an SP 2016 Ar⁺ laser or by a CR-590 dye laser employing Coumarine 6 or Rhodamine 6G dyes pumped by the Ar⁺ laser. The spectra were recorded by a Dilor XY spectrometer. UV/vis/NIR absorption spectra were recorded on Shimadzu UV160 and Bruins Instruments Omega 10 spectrophotometers. A Perkin-Elmer LS 50 B fluorescence spectrometer served to record emission spectra. Cyclic voltammetry was carried out at 200 mV/s scan rate in dmf/0.1 M Bu₄NPF₆ using a three-electrode configuration (glassy carbon working electrode, Pt counter electrode, Ag/AgCl reference) and a PAR 273 potentiostat and function generator. The ferrocene/ferrocenium couple served as internal reference. Spectroelectrochemical measurements were performed using an optically transparent thin-layer electrode (OTTLE) cell²⁶ for UV/vis spectra and a two-electrode capillary for EPR studies.²⁷

(bpym)PtCl₂ (**1**). This compound has been prepared in analogy to previous reports.^{2,4} Elemental analysis, IR, and ¹H NMR data

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Table 1. Crystallographic Data for **1**

chem formula	C ₈ H ₆ Cl ₂ N ₄ Pt
fw	424.15
space group	monoclinic, P2 ₁ /c (No. 14)
T (K)	293(2)
a (Å)	15.201(1)
b (Å)	20.763(2)
c (Å)	6.7942(5)
β (deg)	102.906(8)
V (Å ³)	2090.28(7)
Z	8
ρ _{calcd} (g cm ⁻³)	2.696
μ (mm ⁻¹)	13.90
λ (Å)	0.710 69
final R [I > 2σ(I)]	R ₁ = 0.0346
final R (all data) ^a	R _w = 0.0503

$$^a R_1 = (\sum ||F_o| - |F_c||) / \sum |F_o|, R_w = \{ \sum [w(|F_o|^2 - |F_c|^2)] / \sum [w(F_o^2)] \}^{1/2}.$$

and diffraction studies of the orange-red dmf solvate¹² and of the red solvent-free form confirmed the identity of the product. Solvent-free **1** was obtained in single crystal form through slow evaporation from a dmf solution at elevated temperature (320 K).

(μ-bpym)[PtCl₂]₂ (2). A suspension of 500 mg (1.18 mmol) of (bpym)PtCl₂ in 100 mL of CH₃NO₂ was treated with 3.98 g (9.43 mmol) of (DMSO)₂PtCl₂ and heated to 80 °C for 72 h. The orange-brown solid was collected through filtration, washed with 100 mL of CH₂Cl₂, and dried under vacuum to yield 706 mg (86%) of the poorly soluble product. IR (KBr): ν = 3094 (m), 3062 (m), 1587 (m), 1414 (vs), 1383 (w), 1136 (m), 1046 (w), 810 (m), 715 (s), 373 (m), 341 (m) cm⁻¹. Anal. Calcd for C₈H₆Cl₄N₄Pt₂ (M_r = 690.14): C, 13.92; H, 0.88; N, 8.12. Found: C, 14.17; H, 0.88; N, 8.12.

Crystallography. Solvent-free single crystals of **1** were grown from a solution in dmf through slow evaporation at 320 K. The crystals gave the appearance of an orthorhombic unit cell with a base-centered Bravais lattice (e.g. *a* = 6.7942(5), *b* = 20.7634(16), and *c* = 29.635(2) Å for the *B*-centered case), obviously due to merohedral twinning. However, the transformation matrix [1/2 0 1/2, 0 1 0, -1 0 0] readily generates the correct monoclinic unit cell (see Table 1). Data were collected of a red specimen (0.1 × 0.05 × 0.05 mm) with a STOE IPDS diffractometer. The structure was solved using direct methods with refinement by full-matrix least squares of *F*², employing the programs SHELXS 86 and SHELXL 93.^{28,29} All non-hydrogen atoms were refined anisotropically; hydrogen atoms were introduced at appropriate positions.

DFT Calculations. Ground-state electronic structure calculations of **1**, **2**, and their radical anions have been performed through density-functional theory (DFT) methods using the ADF2000^{30,31} and Gaussian98³² program packages. The lowest excited states of the neutral complexes were calculated by the time-dependent DFT (TD-DFT) method. The Gaussian 98 program was also used for the calculations of vibration frequencies and Raman intensities.

Within Gaussian 98, quasirelativistic effective core pseudopotentials and corresponding optimized sets of basis functions³³ were used for the Pt and Cl centers. Dunning's³⁴ valence double-ζ functions with polarization were used for the H, C, and N atoms. The "Becke-3-LYP" (B3LYP) potential has been employed.³⁵

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Within the ADF program Slater type orbital (STO) basis sets of double-ζ for H and triple-ζ quality with polarization functions for C, N, Cl, and Pt were employed. The inner shells were represented by a frozen core approximation; viz., 1s for C and N, 1s–2p for Cl, and 1s–4d for Pt were kept frozen. The following density functionals were used within ADF: a local density approximation (LDA) with VWN parametrization of electron gas data and a functional including Becke's gradient correction³⁶ to the local exchange expression in conjunction with Perdew's gradient correction³⁷ to the LDA expression (BP). The scalar relativistic (SR) zero-order regular approximation (ZORA) was used within this study. The **g** matrix was obtained from a spin-nonpolarized wave function after incorporating spin-orbit (SO) coupling, using a triple-ζ basis within the frozen core approximation. **A** matrices were obtained from spin-unrestricted wave functions using a quadruple-ζ basis including core electrons.³⁸ **A** and **g** matrices were obtained by first-order perturbation theory from the ZORA Hamiltonian in the presence of a time-independent magnetic field.³⁹

To examine the possible deviation from planarity the geometry optimization and vibrational analysis of **1** and **2** were done without any symmetry constraints. Frequency calculations were done for G98/B3LYP-optimized geometries which involve essentially planar molecules **1** and **2**. The scaling factor for vibrations calculated with the B3LYP functional and double-ζ basis was set at 0.963.⁴⁰ Since the deviations of the optimized structures from ideal higher symmetry are small, single point calculations for **1** and **2** were performed in *C*_{2v} or *D*_{2h} constrained optimized structures, respectively. Within both systems, the bpym ligand defines the *xy* plane and the Pt atoms lie on the *x* axis. To estimate the possible response of electronic structure due to the solvation, the solvent was modeled in ADF/BP calculations by the conductor-like screening model (COSMO) or by the polarized continuum model (PCM) implemented in G98.⁴¹

Results and Discussion

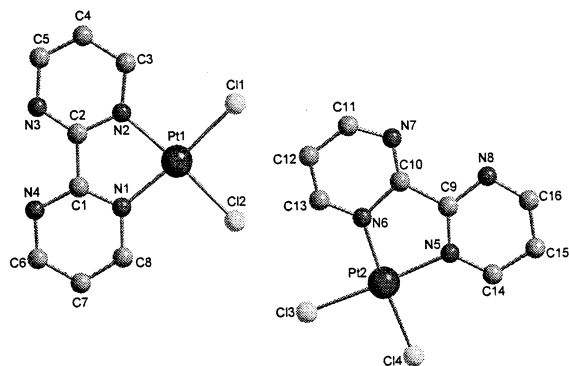
Synthesis. The new dinuclear (μ-bpym)[PtCl₂]₂ (**2**) was obtained as a sparingly soluble yellow material from the

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Table 2. Selected Bond Lengths (Å) and Angles (deg) for the Molecules in the Crystal of **1**^a

Pt1–N1	1.983(10)	Pt2–N5	1.984(9)
Pt1–N2	2.015(9)	Pt2–N6	1.995(10)
Pt1–Cl1	2.286(3)	Pt2–Cl3	2.278(3)
Pt1–Cl2	2.291(4)	Pt2–Cl4	2.289(4)
C1–C2	1.48(2)	C9–C10	1.48(2)
N1–Pt1–N2	78.7(5)	N5–Pt2–N6	80.9(5)
N1–Pt1–Cl1	174.4(4)	N5–Pt2–Cl3	175.1(4)
N2–Pt1–Cl1	95.9(3)	N6–Pt2–Cl3	94.3(3)
N1–Pt1–Cl2	96.1(4)	N5–Pt2–Cl4	95.4(4)
N2–Pt1–Cl2	173.9(3)	N6–Pt2–Cl4	176.2(3)
Cl1–Pt1–Cl2	89.43(13)	Cl3–Pt2–Cl4	89.45(13)

^a For numbering, compare Figure 1.

**Figure 1.** Molecular structures and atomic numbering of the two independent molecules in the unit cell of solvent-free **1**.

prolonged reaction of mononuclear (bpy)PtCl₂ (**1**)^{2,4,11a,12} with (DMSO)₂PtCl₂ in nitromethane. Analytical, spectroscopic, and electrochemical data (cf. below) confirm its identity in relation to **1**. Although it certainly requires some activation to add a second metal complex fragment to the mononuclear (bpy)PtCl₂ molecule^{4,12} with its typically^{2,17} larger “backside” N---N bite in comparison to N–Pt–N, there is no intrinsic problem in obtaining stable dinuclear complexes of platinum metal or other transition metal fragments with the bpy bridging ligand.^{16,17,42} Any loss in basicity at the free nitrogen atoms of the bpy ligand in **1** through the σ acceptor effect of PtCl₂ may be offset by back-donation from occupied metal d orbitals. The slow reactivity of platinum precursor compounds may have prevented the synthesis of **2** in the past.²

Structural Aspects. In addition to two previous crystal structure reports on **1**·0.5nmp (nmp = 1-methyl-2-pyrrolidinone)⁴ and **1**·dmf (dmf = *N,N*-dimethylformamide)¹² we have now obtained crystals of the solvent-free mononuclear complex (bpy)PtCl₂ (Table 1). Whereas the intramolecular structure parameters (Table 2) of the two independent molecules (Figure 1) in the solvent-free crystals are unsurprising and essentially as observed before^{4,12} and calculated by DFT (see Table 3), the arrangement of molecules exhibits an interesting variation to a much studied theme:^{1,2,4,5,12,43,44}

Table 3. Selected ADF/BP Calculated Bond Lengths (Å) and Angles (deg) of [Pt(bpy)Cl₂] (**1**), [Cl₂Pt(bpy)PtCl₂] (**2**) and Their Radical Anions^a

	1	1 ^{•-}	2	2 ^{•-}
Pt–N	2.017	2.011	2.047	2.040
Pt–Cl	2.289	2.330	2.297	2.305
N ₁ –C ₆	1.354	1.341	1.354	1.339
N ₁ –C ₂	1.372	1.409	1.357	1.383
N ₃ –C ₂	1.330	1.361	1.357	1.383
N ₃ –C ₄	1.339	1.324	1.354	1.339
C ₄ –C ₅	1.386	1.392	1.394	1.403
C ₅ –C ₆	1.398	1.412	1.394	1.403
C ₂ –C ₂ '	1.472	1.420	1.427	1.386
N ₁ –Pt–N ₂	80.0	80.6	80.9	81.9
Cl–Pt–Cl	90.7	90.0	92.7	91.9
Pt–N ₁ –C ₆	126.7	126.9	130.5	131.5
Pt–N ₁ –C ₂	115.9	114.9	112.6	111.2
N ₁ –C ₂ –C ₂ '	114.1	114.8	117.0	117.9
N ₁ –C ₂ –N ₃	125.1	122.7	126.1	124.3
N ₃ –C ₂ –C ₅	120.8	122.5	117.0	117.9

^a Numbering (subscripts) according to heterocyclic nomenclature.

the two independent molecules in the unit cell undergo two different kinds of stacking aggregation (Figure 2). Molecule A (with Pt1) exhibits some similarity with the red form of (bpy)Pt(NCS)₂.⁴ One six-membered ring of each molecule A appears to engage in partial π/π interaction with another such ring from the complex A in the next layer. As a result, the vectors bisecting the individual molecules A form an approximately 120° angle φ (ca. 137° rotation in (bpy)Pt(NCS)₂).⁴ The Pt1–Pt1 distance is 3.505 Å, and the Pt1–Pt1–Pt1 zigzag angle 151.5°. A more linear arrangement (170.0°) with closer metal–metal contacts of 3.299 Å was observed for (bpy)Pt(NCS)₂.⁴ Molecules B (with Pt2), on the other hand, are related to the red forms of (bpy)PtCl₂,^{1,5} (bpy)Pt(CN)₂·dmf,⁴⁴ and (bpy)PtCl₂·solvent.^{4,12} The aforementioned vectors are antiparallel ($\varphi = 180^\circ$), and there is no overlap of the six-membered rings. The Pt2–Pt2 distances of 3.434 and 3.466 Å and the Pt2–Pt2–Pt2 angle of 159.9° in the stack are in the range reported previously for chloride-containing “red forms”.^{1,4,5,12}

The availability of catalytically relevant^{11c} free basic nitrogen atoms in **1** allows for weak intermolecular (inter-stack) C–H–N interactions in the crystal as illustrated by a relatively short distance of 3.357 Å between C4 (from molecule A) and N8 (from molecule B) with an approximate C–H–N angle of 176.5°. This additional interaction may be responsible for the occurrence and coexistence of two kinds of stacks within the same crystal structure. Interstack Pt1–Pt2 distances are larger than 7.9 Å.

The structurally observed intermolecular C–H–N interaction suggests that the free nitrogen atoms are still rather basic, probably because of back-donation from the platinum(II) atom. In the catalytic cycle suggested for methane activation by **1** as a precursor in hot sulfuric acid,¹¹ the proposed loss of a chloride ligand would leave a cationic platinum(II) center which could not provide as much electron density for back-donation to bpy. Even then, however, such a mononuclear

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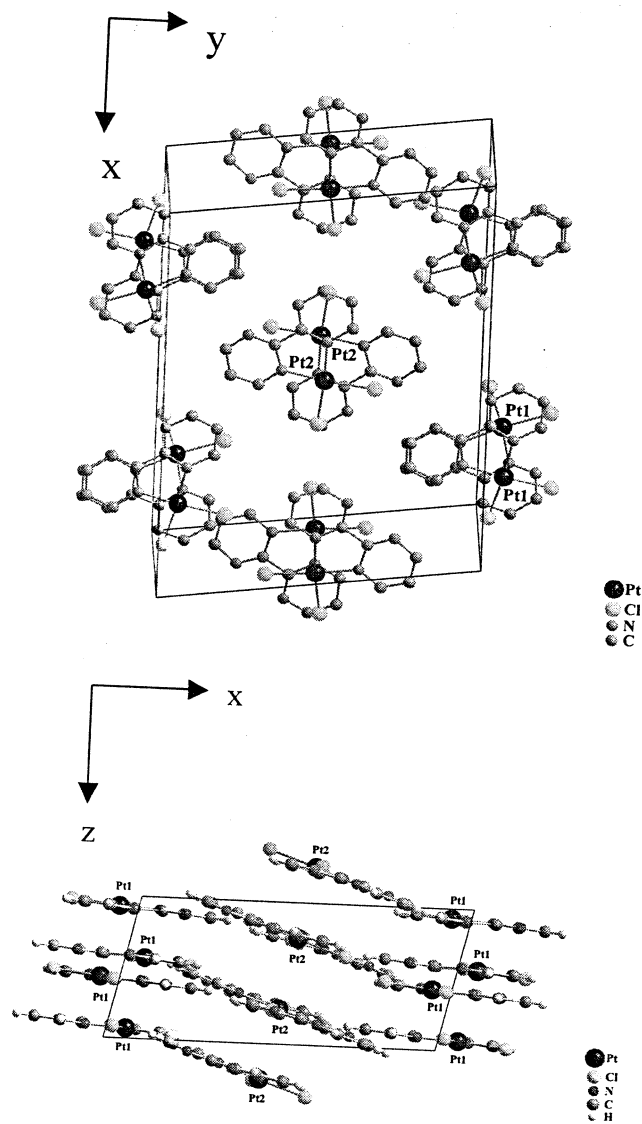


Figure 2. Crystal packing in **1**: Views along the *z* axis (top) and along the *y* axis (bottom).

bpym complex would offer additional binding sites for electrophiles such as H⁺ in concentrated H₂SO₄.^{11c}

The new dinuclear complex **2** has so far failed to yield single crystals of X-ray-diffraction quality. The DFT calculation (Table 3) predicts an essentially planar structure with very similar bond parameters as for **1**. Significant differences lie in the slightly longer Pt–N bonds for **2** due to the absence of bending along the bpym molecular axis^{1,17} (cf. the increase in the angle Pt–N¹–C⁶) and in the shortening of the interring C²–C^{2'} bond.

Reduced State: Cyclic Voltammetry, EPR, and DFT Calculations. Both complexes **1** and **2** are reduced in two well-separated reversible one-electron steps as evident from the cyclic voltammograms obtained in dmf solution (Figure 3). Oxidation processes were not observed up to 1.0 V vs ferrocene. The potentials are listed in Table 4 in comparison to values reported for (μ-bpym)[PtMes₂]₂ (Mes = mesityl = 2,4,6-trimethylphenyl).¹³

Differences of about 0.7 V between the first and second one-electron addition are typical for bpym and its com-

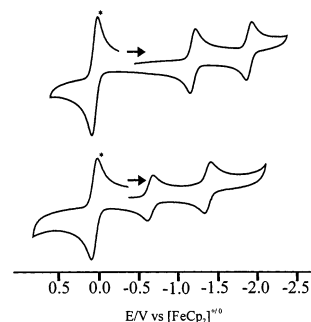


Figure 3. Cyclic voltammograms of **1** (top) and **2** (bottom) in dmf/0.1 M Bu₄NPF₆ (ferrocene standard indicated by an asterisk).

plexes;¹⁹ the ligand itself has $E_{1/2}(\text{red1}) = -2.24$ V and $E_{1/2}(\text{red2}) = -2.95$ V.^{18,19,45} The strong anodic shift of the reduction potential on single and especially double coordination illustrates the σ -acceptor capacity of [PtCl₂]. In contrast to the -1.18 V for (μ-bpym)[PtMes₂]₂, -1.32 V for (μ-bpym)[PtMe₂]₂, and -1.34 V for (μ-bpym)[PtMe₄]₂,¹³ the -0.65 V value for **2** further supports this notion of [PtCl₂] as being a superb σ -acceptor.

A comparison with other 5d metal halide dimers bridged by bpym as shown in Table 5 reveals similarities and differences.

Whereas the potentials for the reversible first electron uptake to a bpym^{•-}-bridged compound are rather similar (Table 5), the chloro complexes containing rhenium(I),^{18,23} osmium(II),¹⁶ and iridium(III)¹⁷ but not platinum(II) exhibit an irreversible second reduction due to chloride dissociation.^{16,17} In addition to an intrinsic stability of the platinum(II)–chloride bond, we attribute this difference to the planar situation at the Pt^{II}N₂Cl₂ center with its 5d⁸ configuration; the other systems mentioned above have a 5d⁶ configuration with the metal–halide bond not in the π plane but largely perpendicular to it. According to simple orbital models^{46,47} this arrangement would favor dissociation through π^*/σ^* conjugative interaction. The controlled reductive dissociation of transition metal-to-chloride bonds is essential for catalytic processes such as the rhenium-catalyzed CO₂ reduction⁴⁸ or the rhodium- or iridium-centered hydride generation.^{49,50}

In situ generation of the radical anion complexes **1**^{•-} and **2**^{•-} allowed us to study the EPR properties of these intermediates in fluid and glassy frozen solution (Figures 4 and 5). X band measurements provided sufficient resolution of the hyperfine-split *g* components, in contrast to other platinum metal radical compounds where high-frequency EPR is necessary.⁵¹ The data as obtained from simulations are summarized in Table 4, again in comparison with those

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Table 4. Electrochemical^a and Spectroscopic Data^{b,c} of the Complexes

	[(bpym)PtX ₂] ^a		{(μ-bpym)[PtX ₂] ₂ } ^a	
	X = Cl	X = Mes ^d	X = Cl	X = Mes ^e
<i>E</i> _{1/2} (red1) ^a	-1.28(66)	-1.67(61)	-0.65(58)	-1.18(84)
<i>E</i> _{1/2} (red2) ^a	-2.01(70)	-2.38(102)	-1.31(76)	-1.92(105)
Δ <i>E</i> _{1/2} (red)	0.73	0.71	0.66	0.74
	EPR ^b			
<i>g</i> ₁	2.034	2.0291	2.049	2.0435
<i>g</i> ₂	2.005	2.0051	2.008	2.0091
<i>g</i> ₃	1.938	1.9453	1.902	1.916
<i>g</i> ₁ - <i>g</i> ₃	0.096	0.0838	0.147	0.1275
<i>g</i> _{iso}	1.9910	1.9927	1.9862	1.9915
<i>A</i> ₁	5.0	3.6	3.9	<2.5
<i>A</i> ₂	6.9 ^f	≈3.0	5.1 ^g	1.6
<i>A</i> ₃	≤3.0	<2.5	≤3.0	<2.0
<i>A</i> _{iso}	4.6	3.0	3.7	<2.5
	UV-Vis ^c			
λ _{max} /ε (<i>n</i> = 0)	470 sh, 402/2200, 390 sh, 365 sh, 300/7200	491/1500, 423 sh	550 br, 465/3300, 435/3200, 400 sh, 345 sh, 315/12400	627/3000, 432/6300, 375/10000, 362
λ _{max} (<i>n</i> = 1)	1020 sh, 910; 483, 452, 425; 350 sh	1150 sh, 950, 828; 506, 477, 453 sh; 356 sh	450, 425 sh; 340 sh, 315 ^h	899, 610 sh, 488
λ _{max} (<i>n</i> = 2)	360 sh, 320	377	1030, 870; 360 sh, 305	n.a.

^a From cyclic voltammetry in dmf/0.1 M Bu₄NPF₆ (X = Cl) or THF/0.1 M Bu₄NPF₆ (X = Mes). Potentials in V vs ferrocenium/ferrocene; peak potential differences in mV (in parentheses). ^b EPR spectra from in situ reduction in dmf/0.1 M Bu₄NPF₆ (X = Cl) or DCE/0.1 M Bu₄NPF₆ (X = Mes). ¹⁹⁵Pt hyperfine coupling constants *A* in mT. ^c Absorption maxima in nm from OTTLE spectroelectrochemistry measurements in dmf/0.1 M Bu₄NPF₆ (X = Cl), THF/0.1 M Bu₄NPF₆ (X = Mes, mononuclear complex) or DCE/0.1 M Bu₄NPF₆ (X = Mes, dinuclear complex). Molar extinction coefficients ε in M⁻¹ cm⁻¹. ^d Reference 7c. ^e Reference 13. ^f A(¹H) = 0.7 mT (2H). ^g A(¹H) = 0.75 mT (2H). ^h Long-wavelength band not observable due to low signal-to-noise ratio (poor solubility).

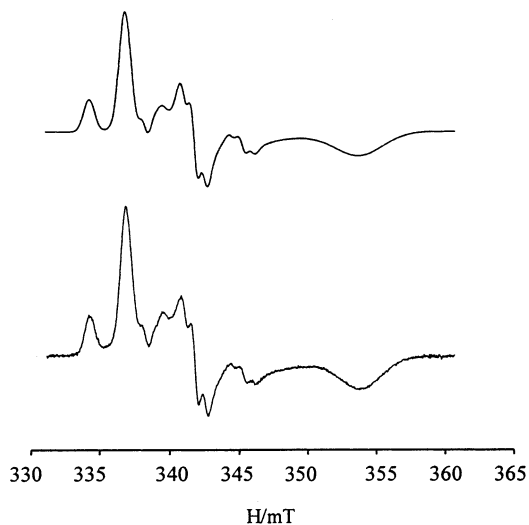
Table 5. Electrochemical,^a EPR,^b and UV-Vis Absorption Data^c for Radical Complexes [(μ-bpym^{•-})[L_nCIM]₂]^k

	L _n CIM			
	(OC) ₃ ClRe	(η ⁶ - <i>p</i> -Cym)ClO _s ⁺	(η ⁵ -C ₅ Me ₅)ClIr ⁺	Cl ₂ Pt
<i>E</i> _{1/2} (ox) ^a	-1.01 ^d	-0.66	-0.74 ^e	-0.65
<i>E</i> _{1/2} (red) ^a	irr ^f	irr ^f	irr ^f	-1.31
	EPR ^b			
<i>g</i> ₁	n.a.	2.0076	2.0029	2.049
<i>g</i> ₂	n.a.	1.9962	1.9849	2.008
<i>g</i> ₃	n.a.	1.9739	1.9451	1.902
<i>g</i> ₁ - <i>g</i> ₃	n.a.	0.0337	0.0578	0.147
<i>g</i> _{iso}	2.0005 ^g	1.9926	1.9778	1.9862
<i>A</i> _{iso} ^h	≈1.2 ^g	1.2	n.o.	3.7
<i>A</i> _{iso} / <i>a</i> _o ⁱ	1 × 10 ⁻³	2.5 × 10 ⁻³	n.o.	3.0 × 10 ⁻³
	UV-Vis ^c			
λ _{max} ^j	444	466	510	450
ref	18, 23	16	17	this work

^a From cyclic voltammetry in dmf/0.1 M Bu₄NPF₆. Potentials in V vs ferrocenium/ferrocene. ^b EPR spectra from in situ reduction in dmf/0.1 M Bu₄NPF₆. ¹⁹⁵Pt hyperfine coupling constants *A* in mT. ^c Absorption maxima in nm from OTTLE spectroelectrochemistry measurements in dmf/0.1 M Bu₄NPF₆. Extinction coefficients not determined for X = Cl because of low solubility. ^d Converted from -0.51 V vs SCE: Juris, A.; Campagna, S.; Bidd, I.; Lehn, J.-M.; Ziessel, R. *Inorg. Chem.* **1988**, *27*, 4007. ^e Low-temperature measurements (-15 °C). ^f Dissociation of chloride. ^g Measurement in acetone. ^h ^{185,187}Re (*I* = 5/2, *a*_o ≈ 1260 mT), ¹⁸⁹Os (*I* = 3/2, *a*_o = 471 mT), ¹⁹⁵Pt (*I* = 1/2, *a*_o = 1228 mT). ⁱ Isotropic hyperfine constant *a*_o from ref 37. ^j Main band in the visible region (π(7) → π(10)).²³

of the mononuclear and dinuclear dimesitylplatinum analogues.

The dichloroplatinum(II) radical species exhibit higher *g* anisotropies *g*₁-*g*₃, lower *g*_{iso} values, and larger ¹⁹⁵Pt hyperfine coupling constants than the [PtMes₂] analogues. While the EPR results for mononuclear **1**^{•-} are expectedly

**Figure 4.** EPR spectrum of electrochemically generated **1**^{•-} in dmf/0.1 M Bu₄NPF₆ at 110 K (bottom) and computer simulation (top).

close to those of [(bpy)PtCl₂]^{•-},⁶ the dinuclear **2**^{•-} is distinguished by a *g* anisotropy which is remarkably large for a radical complex. Both the *g* and the *A* values signify considerable metal participation at the singly occupied MO; however, the alternative mixed-valent resonance structure [Cl₂Pt^{II}(μ-bpym)Pt^{II}Cl₂]^{•-} is still considered to have only a minor contribution in comparison to [Cl₂Pt^{II}(μ-bpym⁻¹)-Pt^{II}Cl₂]^{•-}. The hyperfine coupling in **2**^{•-} indicates equivalence of the two Pt centers. For a delocalized mixed-valent situation (Pt^{1.5}/Pt^{1.5}), one would expect A(**1**^{•-}) = 2A(**2**^{•-}).⁵² In fact, the ¹⁹⁵Pt coupling is only about 30% higher in the mononuclear compound (Table 4), confirming the still predominant

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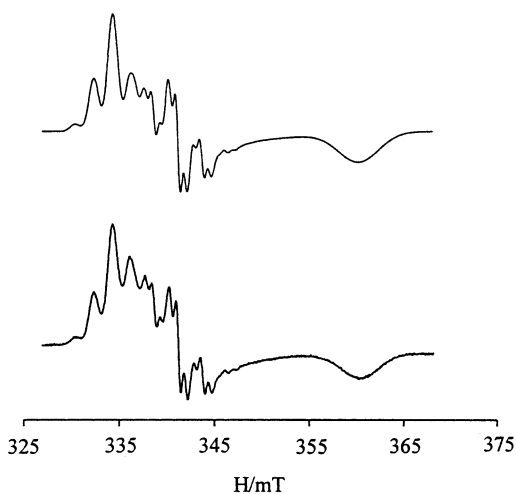


Figure 5. EPR spectrum of electrochemically generated $2^{\bullet-}$ in dmf/0.1 M Bu₄NPF₆ at 110 K (bottom) and computer simulation (top).

radical character of the species. This notion is also supported by the observation of a ¹H coupling of about 0.7 mT for the central *g* components in $1^{\bullet-}$ and $2^{\bullet-}$; in agreement with previous HMO calculations²² we attribute this splitting to the H^{5,5'} protons of the bpym^{•-} ligand.

The particularly large *g* anisotropy and metal hyperfine coupling constant for $2^{\bullet-}$ are also evident (Table 5) from a comparison with other 5d metal halide dimers bridged by the 2,2'-bipyrimidine radical anion.^{16–18} Besides better orbital mixing for increased metal contribution, the higher spin–orbit coupling factor ζ^{53} of platinum(II) in comparison to osmium(II) and rhenium(I) will play an important role. Concerning the deviation of *g* from *g*(electron) = 2.0023, the high ζ value for trivalent iridium is apparent from *g*_{iso} = 1.9778,¹⁷ which lies even lower than that of $2^{\bullet-}$ (Table 5).

For a confirmation and quantification of the above arguments we have used DFT methodology to calculate the structures and EPR properties of $1^{\bullet-}$ and $2^{\bullet-}$. The structural parameters are listed in Table 3 together with those of the nonreduced molecules. Changes in electron acquisition affect predominantly the bpym ligand, especially through the characteristic shortening of the inter-ring bond C²–C^{2'}.

The composition of the ADF/BP-calculated frontier molecular orbitals of [Pt(bpym)Cl₂] is given in Table 6. The close-lying HOMOs 8b₁, 21a₁, and 6a₂ are mainly formed by combinations of Cl orbitals with Pt d orbitals. The MOs 20a₁ and 17b₂ below this set of orbitals are Pt or Cl localized, respectively. The contribution of bpym π MOs to these orbitals is 18% or lower. The LUMO, 9b₁, has the same character as the lowest unoccupied π^* orbital of the free bpym ligand,²⁴ slightly perturbed by a small contribution (6%) from the d_{xz} Pt orbital. The next two higher lying unoccupied molecular orbitals correspond mainly to higher π^* orbitals of bpym.²⁴ An analogous sequence of frontier MOs is obtained for 2 (Table 7). Here the metal-localized orbitals are characterized as pairs of orbitals that correspond to the symmetrical or antisymmetrical combination of PtCl₂

Table 6. ADF/BP Calculated One-Electron Energies and Percentage Composition of Selected Frontier MOs of 1 , Expressed in Terms of Composing Fragments

MO	<i>E</i> (eV)	predominant character	Pt	bpym	Cl (tot.)
Unoccupied					
18b ₂	−2.35	Pt + Cl	46 (d _{yz})	20	33
7a ₂	−3.32	bpym (π_3^*)	3 (d _{xy})	96 (π^*)	1
10b ₁	−3.52	bpym (π_2^*)	4 (d _{xz})	95 (π^*)	1
9b ₁	−4.16	bpym (π_1^*)	6 (d _{xz})	90 (π^*)	3
Occupied					
6a ₂	−5.54	Cl + Pt	41 (d _{xy})	4	54
21a ₁	−5.70	Cl + Pt	25 (d _{z²}); 11 (d _{x²−y²})	13	51
8b ₁	−5.93	Cl + Pt	35 (d _{xz})	18	46
20a ₁	−6.20	Pt	18 (s); 23 (d _{z²}); 53 (d _{x²−y²})	2	3
17b ₂	−6.25	Cl			99
16b ₂	−7.32	bpym	2 (p _y)	9	89

Table 7. ADF/BP Calculated One-Electron Energies and Percentage Composition of Selected Frontier MOs of 2 , Expressed in Terms of Composing Fragments

MO	<i>E</i> (eV)	predominant character	Pt	bpym	Cl (tot.)
Unoccupied					
12b _{3g}	−2.99	Pt + Cl	48 (d _{xz})	21	34
13b _{2u}	−3.45	Pt + Cl	48 (d _{yz})	17	35
5a _u	−4.20	bpym (π_3^*)	6 (d _{xy})	92 (π^*)	2
7b _{2g}	−4.39	bpym (π_2^*)	9 (d _{xz})	87 (π^*)	3
8b _{3u}	−5.09	bpym (π_1^*)	10 (d _{xz})	83 (π^*)	6
Occupied					
5b _{1g}	−6.14	Cl + Pt	41 (d _{xy})	2	57
4a _u	−6.20	Cl + Pt	36 (d _{xy})	9	55
14b _{1u}	−6.30	Cl + Pt	23 (d _{z²}); 12 (d _{x²−y²})	1	64
16a _{1g}	−6.31	Cl + Pt	24 (d _{z²}); 10 (d _{x²−y²})	2	63
6b _{2g}	−6.50	Cl + Pt	2 (p _y); 35 (d _{xz})	14	49
7b _{3u}	−6.70	Cl + Pt	2 (p _y); 34 (d _{xz})	14	49
13b _{1u}	−6.83	Pt	16 (s); 22 (d _{z²}); 54 (d _{x²−y²})	2	4
15a _{1g}	−6.84	Pt	18 (s); 21 (d _{z²}); 55 (d _{x²−y²})	2	3

fragment orbitals. The LUMO is also mainly formed by the π^* orbital of the free bpym ligand, each Pt center contributing by about 5% to this MO. Note that the calculated relative energies of the LUMOs of 1 and 2 are consistent with the reduction potentials. The CH₂Cl₂ solvent effect modeled by the COSMO procedure affects mainly the composition of these HOMOs with large Cl contributions. When the solvent effect is introduced, the Cl contributions within 1 diminish to 39%, 49%, and 34% in orbitals 8b₁, 21a₁, and 6a₂, respectively. In addition, the HOMO orbitals exhibit larger Pt character and the HOMO–LUMO separation increases. The LUMO character, in the other hand, does not change substantially. The corresponding frontier orbitals of 2 are influenced accordingly. Both models of solvation (COSMO within ADF and PCM within G98) enlarge the HOMO–LUMO separation by about 0.5 eV and affect the frontier orbitals comparably.

The reduction of both complexes leads to the acceptance of the added electron by the predominantly bpym-localized LUMO. Figure 6 shows the composition of the singly occupied MO (SOMO) of $2^{\bullet-}$. The SOMO compositions do not substantially differ from those of corresponding LUMOs of the neutral systems. In agreement with the previously reported electronic structure of [Pt(bpy)Cl₂]^{•-},^{6a–d} the 5d contribution to this orbital is much larger than the 6p contribution. The SOMO contains 6% 5d_{xz} and 0.7% 6p_x or

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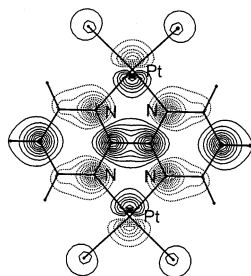


Figure 6. ADF/BP calculated contour map of the SOMO ($8b_{3u}$) of $2^{\bullet-}$, 0.5 \AA over the xy plane.

Table 8. Comparison of Experimental and Calculated g Values^a and ^{195}Pt Hyperfine Constants A (mT)^b for Complexes $1^{\bullet-}$ and $2^{\bullet-}$

	$1^{\bullet-}$		$2^{\bullet-}$	
	expt	calcd	expt	calcd
g_1	2.034	2.076	2.049	2.132
g_2	2.005	1.998	2.008	1.993
g_3	1.938	1.894	1.902	1.820
$g_1 - g_3$	0.096	0.182	0.147	0.312
g_{iso}^e	1.993	1.989	1.987	1.982
A_1	-5.0	-5.98	-3.9	-4.84
A_2	-6.9	-3.63	-5.1	-3.03
A_3	>-3.0	-2.69	>-3.0	-2.50
A_{iso}	-4.6	-4.10	-3.7	-3.46

^a Spin-restricted calculations including spin-orbit coupling. ^b Calculations using scalar relativistic UKS-ZORA approach.

5% $5d_{xz}$ and 0.2% $6p_x$ contributions/Pt atom for $[\text{Pt}(\text{bpym})\text{Cl}_2]^-$ or $[\text{PtCl}_2(\mu\text{-bpym})\text{PtCl}_2]^-$, respectively. The nonzero metal contribution is reflected by the calculated spin densities on the metal center and by the rather large separation of anisotropic g components (Table 8). The spin densities on the individual Pt atoms are 0.053 for $1^{\bullet-}$ and 0.039 for $2^{\bullet-}$, reflected by the lower isotropic A value for the dinuclear complex. The calculations of A values required the inclusion of core electrons. Basis sets with frozen core approximations gave lower A values.

The calculated ^{195}Pt coupling constants and g component values are summarized in Table 8. These calculations were performed without support by spectral parameters to evaluate current methodology. In relation to the experimental data the calculated values seem to exaggerate the metal contribution to the g anisotropy $\Delta g = g_1 - g_3$. Furthermore, the magnitudes of the calculated hyperfine coupling constants are in the order $A_3 < A_2 < A_1$ whereas the experimental observation is $A_3 < A_1 < A_2$. However, both the larger individual hyperfine coupling for $1^{\bullet-}$ vs $2^{\bullet-}$ and the higher total participation of the metals at the SOMO in $2^{\bullet-}$ are reproduced by the DFT calculations. Previous calculations based on EPR spectroscopic data for $[(\text{bpy})\text{PtCl}_2]^-$ and related systems^{6c,d} used Rieger's semiempirical approach⁵⁴ for assigning spin densities to individual centers. A similar analysis in this case gives contributions of 3.6% and 6.5% of the Pt 5d and 6p orbitals, respectively, to the SOMO of $1^{\bullet-}$. For $2^{\bullet-}$ contributions of 2.2% 5d and 4.0% 6p from each Pt center are calculated. These are in good agreement with the DFT calculations with the admixture of both the 5d and 6p orbitals from each Pt center decreasing from $1^{\bullet-}$ to $2^{\bullet-}$.

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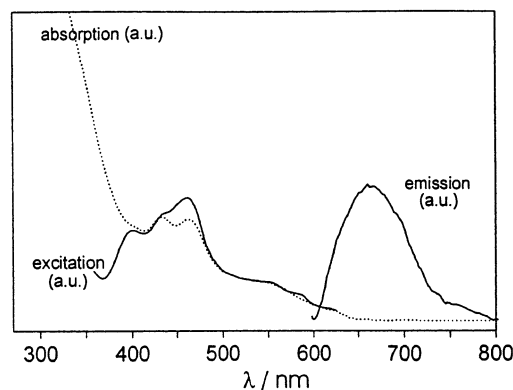


Figure 7. Absorption, emission, and excitation spectra ($\lambda = 700 \text{ nm}$) of 2 in dmf at room temperature.

Excited State: Absorption, Emission, and Excitation Spectra, Spectroelectrochemistry, and Resonance Raman Spectroscopy. Owing to the stabilized $[\text{PtCl}_2]$ -centered HOMOs (Tables 6 and 7), both 1 and 2 exhibit metal-to-ligand charge transfer (MLCT) absorptions at comparatively high energies, despite the low-lying π^* MOs as evident from strongly shifted reduction potentials. Both compounds are luminescent.

The absorption spectrum of mononuclear 1 shows an intense long-wavelength band around 400 nm which exhibits slightly negative solvatochromism ($\lambda_{\text{max}} = 396 \text{ nm}$ in MeCN, 402 nm in dmf, 413 nm in CH_2Cl_2). A closer look at the low-energy side reveals a very weak shoulder around 470 nm. The dinuclear complex 2 exhibits a band system in dmf solution with maxima at 435 and 465 nm. There is an intense band at 315 nm and a broad feature at the low-energy side around 550 nm. Due to the poor solubility of 2 in most solvents its solvatochromism could not be investigated.

The mononuclear complex shows emission at ambient temperatures. In the solid, the excitation at 480 nm into the weak long-wavelength absorption band leads to a weak and broad emission at 670 nm. Resonance Raman experiments gave essentially the same results. The excitation spectrum at $\lambda = 650 \text{ nm}$ gave main features at 401 and 472 nm. These maxima coincide with absorption bands observed in dmf solution at 405 and 470 sh nm. On the other hand, no emission was observed in a dmf solution of 1 . Only when samples were excited between 320 and 360 nm, a weak emission could be detected at 440 nm. The corresponding excitation spectrum yielded an intense, relatively narrow band with a maximum at 340 nm, responsible for the emission activity.

Emission spectroscopy of dinuclear 2 (Figure 7) gave essentially the same results for the solid and the dmf solution at room temperature. Excitation at 490 nm resulted in broad unstructured emission bands with maxima at 700 nm for the solid or at 660 nm in dmf, respectively. Resonance Raman experiments yielded essentially similar emission results. The excitation spectrum of 2 in the solid ($\lambda = 720 \text{ nm}$) showed maxima at 398, 439 sh, 470, and 502 nm that in dmf ($\lambda = 700 \text{ nm}$) gave maxima at 405, 435 sh, 460, and 550 nm. These values are in good agreement with the absorption maxima found at 400 sh, 435, 465, and ca. 550 nm for the dmf solution of 2 with two d^8 metal centers. While dinuclear

Table 9. Selected Calculated Lowest TD-DFT Singlet Transition Energies and Oscillator Strengths for **1**

state	main character (in %)	G98/B3LYP (vacuum)		G98/B3LYP PCM (CH ₂ Cl ₂)		expt (CH ₂ Cl ₂): λ _{max} /ε ^b
		trans. energy ^a (wavelength) ^a	osc str	trans. energy ^a (wavelength) ^a	osc str	
¹ B ₂	98 (6a ₂ → 9b ₁) (HOMO → LUMO)	2.00 (620)	0.002	2.56 (484)	0.002	470 sh
¹ A ₁	77 (8b ₁ → 9b ₁); 19(6a ₂ → 7a ₂) (HOMO - 2 → LUMO)	2.51 (494)	0.044	3.00 (412)	0.049	413/2200
¹ B ₂	96 (6a ₂ → 10b ₁) (HOMO → LUMO + 1)	2.82 (439)	0.013	3.41 (363)	0.012	365 sh
¹ A ₁	54(6a ₂ → 7a ₂); 40 (8b ₁ → 10b ₁) (HOMO → LUMO + 2)	3.06 (405)	0.004	3.06 (405)	0.004	390 sh
¹ A ₁	55 (8b ₁ → 10b ₁); 24 (6a ₂ → 7a ₂); 14 (8b ₁ → 9b ₁) (HOMO - 2 → LUMO + 1)	3.37 (367)	0.094	3.89 (318)	0.066	300/7200

^a Transition energies in eV; wavelengths in nm. ^b Absorption maxima in nm; molar extinction coefficients in M⁻¹ cm⁻¹.

Table 10. Selected Calculated Lowest TD-DFT Singlet Excitation Energies and Oscillator Strengths for **2**

state	main character (in %)	G98/B3LYP (vacuum)		G98/B3LYP PCM (CH ₂ Cl ₂)		expt (CH ₂ Cl ₂): λ _{max} /ε ^b
		trans. energy ^a (wavelength) ^a	osc str	trans. energy ^a (wavelength) ^a	osc str	
¹ B _{2u}	98 (5b _{1g} → 8b _{3u}) (HOMO → LUMO)	1.69 (734)	0.002	2.06 (603)	0.001	550 br ^c
¹ B _{1u}	84 (6b _{2g} → 8b _{3u}), 14 (5b _{1g} → 5a _u) (HOMO - 4 → LUMO)	2.07 (598)	0.094	2.42 (513)	0.093	465/3300
¹ B _{2u}	92 (4a _u → 7b _{2g}) (HOMO - 1 → LUMO + 1)	2.58 (480)	0.013	3.22 (385)	0.014	435/3200 ^d
¹ B _{1u}	74 (5b _{1g} → 5a _u), 84 (7 _{3u} → 7b _{2g}) (HOMO → LUMO + 2)	2.75 (452)	0.021	3.10 (399)	0.009	

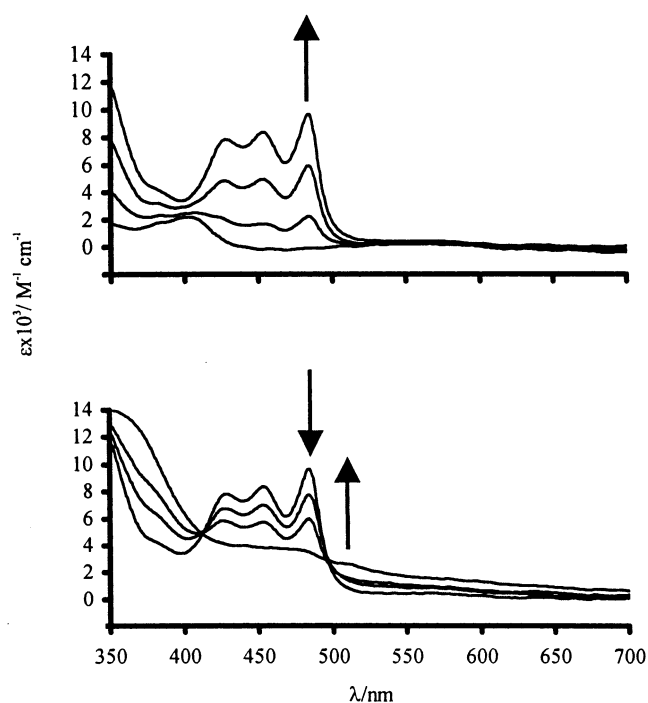
^a Transition energies in eV; wavelengths in nm. ^b Absorption maxima in nm; molar extinction coefficients in M⁻¹ cm⁻¹. ^c Weak absorption tails down to ca. 650 nm. ^d Alternatively, this band may be assigned as a vibronic sideband of the 465 nm band.

complexes of bpym with d⁶ metal centers are typically poor emitters,^{24,55} there are dicopper(I) (d¹⁰) complexes with strong luminescence in the solid and in solution.⁴²

Tables 9 and 10 summarize the TD-DFT G98/B3LYP-calculated transition energies of the lowest-lying electronic transitions together with their character for **1** and **2**. Calculations using ADF/BP gave the same transition characters but lower excitation energies and are not listed there. The calculated transition energies correspond to the gas-phase situation and are lower than those observed in solution, a result not unexpected for planar systems open to solvent influence. To estimate the shifts of transition energies due to solvation, the TD-DFT procedure is applied to Kohn–Sham orbitals calculated by the polarized continuum model (PCM).⁴¹ As a consequence of the larger separation of frontier orbitals, the calculated transitions approximated within PCM(CH₂Cl₂) are shifted to higher energies, thus generally corresponding better to the experimental results. The calculated blue shift is in agreement with experimental trends within these complexes.

The lowest-energy transitions calculated for **1** correspond to the excitations from MOs formed mainly by Pt and Cl orbitals into the π* of the bpym ligand. They can be characterized as mixed metal-to-ligand charge transfer (MLCT) transitions with some halide-to-ligand (XLCT) contributions. The lower-lying transitions within **2** have similar character. The lowest broad absorption band is assigned to the HOMO(5b_{1g}) → LUMO(8b_{3u}) transition. The more intense and slightly solvatochromic bands were attributed to the symmetry-allowed MLCT/XLCT transitions (mainly d_{xz} → π*).

On reduction to the radical anions, there are weak (forbidden) intraligand transitions expected in the near-infrared region.^{7,15,23} While these could be detected for **1**^{•-} (Figure 8, Table 4) by OTTLE spectroelectrochemistry,²⁶ the poorly

**Figure 8.** Spectroelectrochemical reduction of **1** to **1**^{•-} (top) and of **1**^{•-} to **1**^{2•-} (bottom) in dmf/0.1 M Bu₄NPF₆.

soluble system **2**/**2**^{•-} did not allow a detection of such features. However, the typical intense band at about 450 nm for dinuclear complexes of bpym^{•-}^{15,23} could be observed (Table 5).

The relatively small symmetrical complexes **1** and **2** are well suited for vibrational spectroscopy, including resonance Raman studies. We have previously reported corresponding results for other bpym-bridged compounds, viz., (μ-bpym)-[M(CO)₄]₂, M = Mo and W.²⁴

Resonance Raman spectra could be obtained for **1** only on excitation at 488 or 458 nm. Both wavelengths lie within the low-intensity absorption bands of **1**. The rR spectra show rather weak resonances at 1565, 1497, 1225, 1018, 923, 878,

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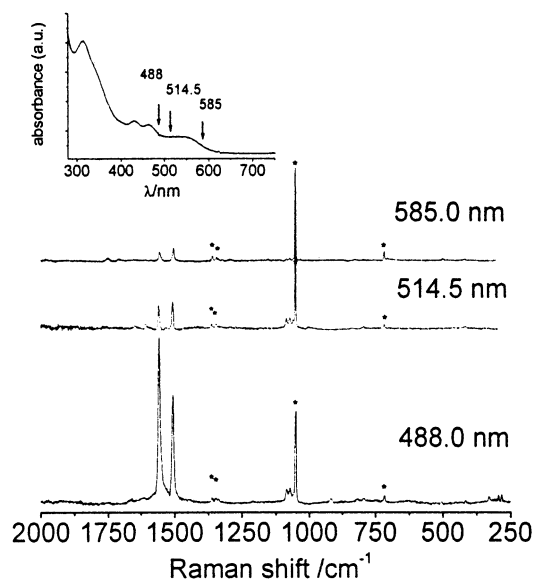


Figure 9. Resonance Raman spectra of **2** in KNO_3 at various wavelengths (absorption spectrum and excitation wavelengths in the insert; asterisks indicate NO_3^- vibrations).

673, 570, and 454 cm^{-1} . The first six of them are quite similar to resonances observed for the free ligand and the related tungsten complex $(\text{bpym})\text{W}(\text{CO})_4$;²⁴ DFT calculations yield values of 1596 and 1510 cm^{-1} for the highest Raman-active vibrations associated with the bpym ligand.

On irradiation at 597, 532, or 514.5 nm into the long-wavelength absorption band of dinuclear **2** the resonance Raman spectra also show weak resonances at 1560, 1509, 1085, 1071, 815, 793, 499, and 416 cm^{-1} (Figure 9). However, upon irradiation at 488 or 458 nm into the partly structured *second* absorption band the rR spectrum showed strong intensity enhancement of some of these bands, particularly those at 1560 and 1509 cm^{-1} . Further resonances are now observed at 1336, 1089, 1077, 924, 790, and 302 cm^{-1} whereas the bands at 499 and 417 cm^{-1} have disappeared. DFT calculations of Raman-active frequencies associated with the high-energy bpym stretching vibrations yield values of 1637 and 1520 cm^{-1} ; the corresponding modes are illustrated in Figure 10.

The resonance Raman results can be discussed in connection with the assignments given above. As long as low-intensity MLCT bands are involved, no strong resonance enhancement is observed for both the mono- and dinuclear complex. Upon irradiation into the second band of the dinuclear complex a very strong enhancement of typical vibration modes of the bridging chelate ligand²⁴ is observed. In view of the findings for related dinuclear complexes $(\mu\text{-bpym})[\text{M}(\text{CO})_4]_2$ ($\text{M} = \text{W}$ or Mo)²⁴ the occurrence of at least two different MLCT transitions has to be considered. The absorption spectra of **1** and **2** are quite similar to those of the molybdenum(0) or tungsten(0) complexes as is the rR behavior upon excitation into the two distinguishable absorption bands. On the basis of the rR results for $(\mu\text{-bpym})[\text{M}(\text{CO})_4]_2$,²⁴ these two bands have been assigned to the HOMO–SLUMO as the lower and the SHOMO–LUMO as the higher, more intense transition. The resonances at 1557

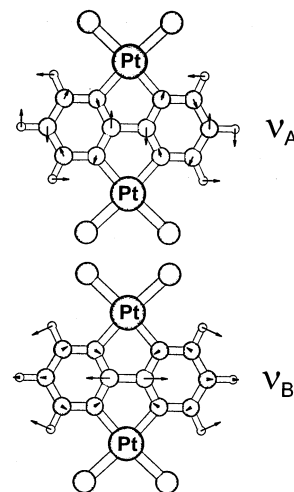


Figure 10. Pictorial representation of Raman active vibrations for **2** calculated at 1637 cm^{-1} (ν_A) and 1520 cm^{-1} (ν_B).

and 1478 cm^{-1} (for $\text{M} = \text{W}$) were found to change in relative intensity on going from band I to band II (the former increases), and in addition, a band at 1332 cm^{-1} emerges which was attributed to the inter-ring vibration $\nu(\text{CC})$ and is consequently enhanced only when the LUMO is the target orbital. For the present dinuclear complex **2** we found exactly the same behavior. The 1560 cm^{-1} band of **2** gradually gains in intensity as compared to the 1506 cm^{-1} band when going through the excitation wavelengths from 597 to 514.5 nm. However the inter-ring vibration $\nu(\text{CC})$ is much weaker than observed for the $\text{M}(\text{CO})_4$ derivatives.²⁴ Similar behavior was observed for the two related complexes $(\text{CO})_5\text{MnRe}(\text{CO})_3(\mu\text{-bpym})\text{W}(\text{CO})_4$ and $(\text{CO})_5\text{MnRe}(\text{CO})_3(\mu\text{-bpym})\text{Re}(\text{CO})_3\text{Br}$ where also a strong enhancement of $\nu(\text{CC})$ was found for the former species whereas the latter compound showed a much weaker such resonance.⁵⁶ Only if there is already an appreciable amount of electron density at the ring-connecting C atoms in the π^* orbitals of bpym in the ground state, the effect on that C–C bond by the CT transition appears to be strong enough to give rise to a significant rR effect for this vibration. In agreement with the other results of this study the $[\text{PtCl}_2]$ complex fragment must then be considered a relatively weak π -donor toward such ligands.

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Supporting Information Available: Details of the X-ray structure determination in CIF format. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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