Electron Paramagnetic Resonance and Spectroscopic Characteristics of Electrogenerated Mixed-Valent Systems $[(\eta^5 \text{-} C_5 M \text{e}_5)M(\mu \text{-} L)M(\eta^5 \text{-} C_5 M \text{e}_5)]^+$ (M = Rh, Ir; L = **2,5-Diiminopyrazines) in Relation to the Radicals** $[(\eta^5 \text{-} C_5\text{Me}_5) \text{C} \text{M}(\mu \text{-} \text{L}) \text{M} \text{C} \text{M}(\eta^5 \text{-} C_5\text{Me}_5)]^+$ and $[(\eta^5 \text{-} C_5 \text{Me}_5) \text{M}(\mu \text{-} \text{L}) \text{M} \text{Cl}(\eta^5 \text{-} C_5 \text{Me}_5)]^{2+}$

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Electrochemical reduction of the dinuclear $[(\eta^5 - C_5M_e)(C_M/L)MCl(\eta^5 - C_5M_e)]^{2+}$ ions (M = Rh, Ir; L = 2,5bis(1-phenyliminoethyl)pyrazine (bpip) and 2,5-bis[1-(2,6-dimethylphenyl)iminoethyl]pyrazine (bxip)) proceeds via the paramagnetic intermediates $[(\eta^5 \text{-} C_5 \text{Me}_5) \text{Cl}(u \text{-} L) \text{MCl}(\eta^5 \text{-} C_5 \text{Me}_5)]^+$ (L = bpip) or $[(\eta^5 \text{-} C_5 \text{Me}_5) \text{M}(u \text{-} L) \text{-} C_5 \text{Me}_5]$ $MCl(\eta^5-C_5Me_5)|^{2+}$ (L = bxip) and $[(\eta^5-C_5Me_5)M(\mu-L)M(\eta^5-C_5Me_5)]^+$. Whereas the first is clearly a radical species with a small *g* anisotropy, the chloride-free cations are distinguished by structured intervalence charge transfer (IVCT) bands in the near-infrared region and by rhombic electron paramagnetic resonance features between $g =$ 1.9 and $g = 2.3$, which suggests considerable metal participation at the singly occupied MO. Alternatives for the d configuration assignment and for the role of the bisbidentate-conjugated bridging ligands will be discussed. The main difference between bpip and bxip systems is the destabilization of the chloride-containing forms through the bxip ligand for reasons of steric interference.

Introduction

In contrast to the numerous mixed-valent complexes L_nM^k - $(\mu$ -L)M^{k+1}L_n with d^{5}/d^{6} configurated metal centers M and bridging ligands $L¹⁻⁶$, there have been relatively few⁷⁻¹³ such compounds with other d^{n-1}/d^n configurations. In the course of research on the coupling of organometallic redox reaction centers through mediating bridging ligands μ ; η ⁴-L,¹⁴⁻¹⁸ we have

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now been able to study the systems $[(\eta^5{\text{-}}C_5Me_5)M(\mu{\text{-}}L)M(\eta^5{\text{-}}C_5Me_5)]$ (C_5Me_5) ⁺ (M = Rh, Ir; L = 2,5-bis(1-phenyliminoethyl)pyrazine (bpip) or 2,5-bis[1-(2,6-dimethylphenyl)iminoethyl]pyrazine (bxip) by electron paramagnetic resonance (EPR) and by

bxip

absorption spectroscopy in the near-infrared, the typical region for intervalence charge transfer (IVCT) transitions. These results, as obtained through in situ EPR and OTTLE (optically transpar-

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ent thin-layer electrolysis)¹⁹ spectroelectrochemistry, will be compared with the EPR and UV/vis/NIR data of related paramagnetic precursor complexes $[(\eta^5-C_5Me_5)CIM(\mu-L)MC]$ $(\eta^5$ -C₅Me₅)]⁺ and $[(\eta^5$ -C₅Me₅)M(μ -L)MCl(η^5 -C₅Me₅)]²⁺.

Bpip has been used in a number of cases $16-18,20,21$ as a symmetrically bisbidentate-conjugated bridging ligand containing a central pyrazine ring that, in unsubstituted form, is the bridge in the prototypical mixed-valent Creutz-Taube ion.^{1-4,22,23} The bxip ligand is presented here for the first time. Its conception was based on structural results obtained for a pentamethylcyclopentadienyliridium complex of 1,4-bis(2,6 dimethylphenyl)-1,4-diazabutadiene,

which revealed electronic as well as steric stabilization of the coordinatively unsaturated metal center. $24-26$

Experimental Section

Materials. The ligand bpip and the complexes $[(\eta^5 \text{-} C_5 \text{Me}_5) \text{C} \text{M}(\mu - \text{Me}_5)]$ bpip)MCl($η$ ⁵-C₅Me₅)](PF₆)₂ have been described previously.^{16-18,20,21}

The ligand bxip was prepared as follows. A mixture containing 2,5 diacetylpyrazine (100 mg, 0.609 mmol), freshly distilled 2,6-dimethylaniline (238 mg, 1.34 mmol), $4 \text{ Å molecular sieves (3 g)}$, and acidic alumina (20 mg) was heated in toluene (50 mL) under reflux for 24 h. The hot solution was filtered and reduced in volume to 30 mL, and the yellow crystals were collected. Washing with *n*-heptane, recrystallizing from diethyl ether, and drying under vacuum yielded the pale-yellow product in 109 mg yield (48%). Found: C, 76.54; H, 8.26; N, 14.06%. C24H30N4 (374.53) requires C, 76.97; H, 8.07; N, 14.96%. NMR *δ*^H (acetone- d_6): 2.12 (12H, s, CH₃), 2.68 (6H, s, CH₃), 6.45 (2H, t, CH_{Ph}), 6.82 (4H, d, CH_{Ph}), 9.17 (2H, s, CH_{pz}) ppm; ³*J* (CH_{Ph}) = 7.45 Hz.

The complexes [($η$ ⁵-C₅Me₅)ClM($μ$ -bxip)MCl($η$ ⁵-C₅Me₅)](PF₆)₂ were obtained in a similar manner as the bpip compounds^{20,21} by reacting [M(η ⁵-C₅Me₅)Cl₂]₂ with 2 equiv of AgPF₆ in acetone, filtration, and addition of the free bridging ligand (1 equiv). After 1 h the deeply colored solutions were filtered and reduced in volume, and the filtrate was then treated with 1 equiv of Bu_4NPF_6 to yield the purplish-black (Ir) or brown-red (Rh) product in about 70% yield.

 $[(\eta^5-C_5Me_5)ClIr(\mu\text{-}bxi)IrCl(\eta^5-C_5Me_5)](PF_6)_2$ (cis (A) and trans (B) isomers in 1:3 ratio). Found: C, 37.62; H, 4.21; N, 3.60%. C₄₄H₅₆-Cl2F12Ir2N4P2 (1390.27) requires C, 38.12; H, 4.07; N, 4.04%. NMR *δ*_H (CD₃NO₂): 1.65 (A) and 1.66 (B) (s, 30H, ($η$ ⁵-C₅Me₅)), 2.29 (B) and 2.35 (A) (s, 6H, CH₃(phenyl)), 2.44 (B) and 2.45 (A) (s, 6H, CH₃-(phenyl)), 2.82 (B) and 2.87 (A) (s, 6H, CH_3 (imine)), 7.48 (A,B) (m, 6H, CH_{Ph}), 9.57 (B) and 9.58 (A) (s, 2H, CH_{pz}) ppm.

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 $[(\eta^5-C_5Me_5)CIRh(\mu-bxip)RhCl(\eta^5-C_5Me_5)](PF_6)$ ₂ (cis and trans isomers in 1:1 ratio). Found: C, 43.84; H, 5.28; N, 4.88%. $C_{44}H_{56}Cl_2$ -F12N4P2Rh2 (1207.62) requires C, 43.67; H, 4.67; N, 4.64%. NMR *δ*^H (CD₃NO₂): 1.64 and 1.65 (s, 30H, ($η$ ⁵-C₅Me₅)), 2.22 and 2.25 (s, 6H, CH3(phenyl)), 2.46 and 2.48 (s, 6H, CH3(phenyl)), 2.83 and 2.86 (s, 6H, CH₃(imine)), 7.50 (m, 6H, CH_{Ph}), 9.57 and 9.58 (s, 2H, CH_{pz}) ppm.

Instrumentation. EPR spectra were recorded in the X band on a Bruker System ESP 300 equipped with a Bruker ER035M gauss meter and a HP 5350B microwave counter. A continuous flow cryostat ESR 900 from Oxford Instruments was used for liquid He cooling. ¹ H NMR spectra were taken on a Bruker AC 250 spectrometer, and UV/vis/ NIR absorption spectra were recorded on a Bruins Instruments Omega 10 spectrophotometer. Cyclic voltammetry was carried out in solutions containing 0.1 mol dm⁻³ in 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 an internal reference. Digital simulation was performed using the program ESP (C. Nervi)²⁶ suitable for staircase voltammetry. Spectroelectrochemical measurements were performed using an optically transparent thin-layer electrode (OTTLE, Pt-grid) cell19 for UV/vis spectra and a two-electrode capillary for EPR studies.²⁷

Results and Discussion

Characterization of Precursors. The precursor complexes $[(\eta^5$ -C₅Me₅)ClM(μ -bpip)MCl(η^5 -C₅Me₅)](PF₆)₂, M = Rh and Ir, have been described previously.20,21 The new compounds bxip and $[(\eta^5-C_5Me_5)CIM(\mu-bxip)MC](\eta^5-C_5Me_5)](PF_6)$ ₂ were prepared accordingly. The complexes were obtained as mixtures of cis and trans isomers

with respect to the arrangement of Cl and $(\eta^5$ -C₅Me₅) ligands vs the central molecular plane, which, however, does not significantly affect the EPR and (spectro)electrochemical results.20,21,28

Cyclic Voltammetry. Dinuclear complex ions of the general formula $[(C_nR_n)HaM(\mu-L)MHa(C_nR_n)]^{2+}$, $n = 5$ and $M = Rh$ or Ir, $n = 6$ and M = Ru and Os, are known to undergo typical electron transfer/atom transfer sequences as depicted in the Scheme 1.20,21,28 Essential features of this scheme as elucidated by cyclic voltammetry are the reversible one-electron reduction (at potential E_1) of the precursor dications before a second electron (at *E*2) causes the first chloride dissociation ("electron reservoir" behavior of L)^{20,21,28} and the dissociative third oneelectron addition at *E*3, which leads to a chloride-free mixedvalent intermediate. The fourth electron at *E*⁴ produces a neutral

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^{305.}

Figure 1. Cyclic voltammograms of $[(\eta^5 \text{-} C_5\text{Me}_5) \text{CIRh}(\mu \text{-} b \text{xip}) \text{RhCl}$ ($η$ ⁵-C₅Me₅)](PF₆)₂ in DMF/0.1 M Bu₄NPF₆ at 100 mV/s.

compound that may be reduced one more time at very negative potentials (E_5) .

The main difference between corresponding bpip and bxip compounds as now observed here lies in the diminished reversibility of the first ligand-centered reduction for the bxip complexes (Figure 1). While the expected cathodic shift of potentials due to the introduction of electron-releasing methyl substituents is marginal, the apparent and structurally documented²⁴ steric effect of the space-filling 2,6-dimethylphenyl group is to disfavor the more congested chloridecontaining M(III) precursor and to kinetically stabilize the chloride-free system with its lower coordination number at the metal center. Accordingly, the diiridium complex of the bxip ligand exhibits a cyclic voltammetric response with two return peaks (E_1^{pa}, E_{1x}^{pa}) for the first reduction process, corresponding to chloride-containing and chloride-free species (Table 1).25 For the rhodium system, the reversible counterpeak relating to chloride-containing species is detectable at $E^{pa} = -0.46$ V only at scan rates above 500 mV/s. In contrast to the situation for bpip complexes,²¹ the reoxidation peaks are well separated in the case of the bxip analogues, which allows for an evaluation of the rate constants for chloride dissociation from $[(\eta^5{\text{-}}C_5Me_5)$ -ClM(μ -bxip)MCl(η ⁵-C₅Me₅)]⁺. The values $k = 5.9 \pm 0.9 \text{ s}^{-1}$ and $k = 1.1 \pm 0.2$ s⁻¹ for M = Rh and M = Ir, respectively,

Table 1. Peak Potentials from Cyclic Voltammetry*^a* of Complexes [($η$ ⁵-C₅Me₅)ClM($μ$ -L)MCl($η$ ⁵-C₅Me₅)](PF₆)₂

	μ -L = bpip		μ -L = bxip	
F^b	$M = Ir$	$M = Rh$	$M = Ir$	$M = Rh$
$E_1^{\rm pc}$ $E_2^{\rm pc}$ $E_3{}^{pc}$ $E_4{}^{pc}$ $E_5^{\rm pc}$ E_1x^{pa} E_2 ^{pa} E_1 ^{pa} E_{2v} ^{pa} E_3 ^{pa}	-0.44 -0.83 $-1.54c$ $-1.54c$ ≤ -2.7 $-0.36c$ $-0.36c$ -1.23 -1.44	-0.50 -0.76 -1.25 -1.39 -2.45 $-0.43c$ $-0.43c$ -1.03 -1.33	-0.46 -0.78 -1.52^{c} -1.52^{c} -2.69 -0.19^{c} $-0.19c$ -0.39 -0.52 -1.10 -1.45	-0.53 -0.93 -1.23 -1.37 -2.27 $-0.31c$ $-0.31c$ -0.56 -0.84
$E_4{}^{pa}$ E_5^{pa}	≤ -2.6	-2.37		-1.31 -2.20

^a In CH3CN (bpip compounds) or DMF (bxip complexes) with 0.1 M Bu₄NPF₆ at 100 mV/s scan rate. Potentials vs $\text{FeCp}_2^{+/0}$. Superscript pc: cathodic peak. Superscript pa: anodic peak. *^b* For identification of processes see scheme in the main text. *^c* Two overlapping oneelectron peaks according to spectroelectrochemistry.

were obtained by fitting the experimental cyclic voltammograms using the ESP²⁶ simulation program. (The fitting procedure was applied to experimental cyclic voltammograms involving the first two reduction and corresponding return peaks. The error limit refers to the standard deviation obtained from a set of cyclic voltammograms at different scan rates.)

The second reduction of $[(\eta^5-C_5Me_5)CIM(\mu-bxip)MC](\eta^5-C_5-\eta^2)$ $Me₅$)]²⁺ (peak B in Figure 1) is followed by fast chloride dissociation; cyclic voltammetry within peak B does not exhibit any counterpeak at scan rates up to 10 V/s. Furthermore, no new anodic peak appears when the potential sweep is extended from the first reduction peak (A) behind the second (B); however, the cyclic voltammogram is then showing increased height of the reoxidation peak (H). Hence, the potentials of the reoxidation processes $[(\eta^5-C_5M_{\text{e}_5})M(\mu-bxi)p)MC](\eta^5 (C_5Me_5)$]^{+ \rightarrow 2+ \rightarrow 3+ are close, which results in the disproportion-} ation of the species $[(\eta^5$ -C₅Me₅)M(μ -bxip)MCl(η^5 -C₅Me₅)]²⁺. The electrode mechanism of the two-electron reduction of $[(\eta^5 C_5Me_5$)ClM(μ -bxip)MCl(η ⁵-C₅Me₅)]²⁺ can thus be interpreted as consecutive one-electron transfers, coupled to a series of

Scheme 1

$$
+CI^{+}C_{1}^{+}C_{2}^{+}M_{2}^{+}C_{3}^{+}M_{3}^{+}C_{4}^{+}M_{4}^{+}C_{5}^{+}M_{5}^{+}C_{6}^{+}M_{6}^{+}C_{7}^{+}M_{7}^{+}C_{8}^{+}M_{8}^{+}C_{9}^{+}M_{9}^{+}C_{1}^{
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\uparrow\uparrow\cdot C_{5}Me_{5})M(\mu-L)MCl(\eta^{5}\cdot C_{5}Me_{5})] \longrightarrow [\eta^{5}\cdot C_{5}Me_{5})M(\mu-L)M(\eta^{5}\cdot C_{5}Me_{5})] \uparrow\downarrow E_{4}
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\uparrow\downarrow E_{4}
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chemical reactions that include chloride dissociations, and a disproportionation equilibrium of the species $([(\eta^5-C_5Me_5)M (\mu$ -bxip)MCl(η ⁵-C₅Me₅)]²⁺ \rightarrow $[(\eta$ ⁵-C₅Me₅)M(μ -bxip)MCl(η ⁵- (C_5Me_5)]⁺ + [(η ⁵-C₅Me₅)M(μ -bxip)MCl(η ⁵-C₅Me₅)]³⁺). Further evidence for the disproportionation reaction comes from the decreased height of the second reduction peak (B, when compared with A), clearly seen at low scan-rates, and from spectroelectrochemically observed direct formation of the twoelectron reduction product during the first step of the electrolysis.

The mixed-valent species $[(\eta^5$ -C₅Me₅)M(μ -bxip)MCl(η^5 - C_5Me_5]⁺ undergoes the expected ECE reduction mechanism (overlapping peaks C, D), leading to the chloride-free species [(*η*5-C5Me5)M(*µ*-bxip)M(*η*5-C5Me5)] via [(*η*5-C5Me5)M(*µ*-bxip)- M(*η*5-C5Me5)]+. Reoxidation at peaks E, F, G, H is a chemically reversible process during which chloride ions return to the coordination sphere (Scheme 1).

The singly reduced paramagnetic species $[(\eta^5-C_5M_{\rm g})M(\mu$ bxip)MCl(η^5 -C₅Me₅)]²⁺ could be electrogenerated in sufficient amounts for EPR spectroscopy. Similarly, the intermediate species $[Cp^*M(\mu-L)MCp^*]^+$ could be characterized by EPR and OTTLE spectroelectrochemistry, although the difference between E_3 ^{pc} and E_4 ^{pc} can be quite small, ranging from 140 mV ($M = Rh$) to less than 50 mV ($M = Ir$). The pertinent values for the systems described in this work are summarized in Table 1.

The effect of metal replacement (Rh by Ir) results especially in cathodic shifts of E_3^{pc} and E_4^{pc} , confirming the interpretation of these processes as more metal-centered, in contrast to the presumably more ligand-centered processes *E*1.

EPR Spectroscopy of Paramagnetic Intermediates. According to the redox scheme, at least two paramagnetic intermediates should be accessible in each case by prolonged electrolysis for in situ EPR characterization. Whereas the first, the one-electron reduction products of the precursor compounds, presented no problems and yielded relatively narrow signals at ambient temperatures, the next odd-electron species, the threeelectron reduction products $[(\eta^5 \text{-} C_5\text{Me}_5)M(\mu\text{-}L)M(\eta^5 \text{-} C_5\text{Me}_5)]^+,$ were only observable by EPR at lower temperatures in glassy frozen solutions, indicating rapid relaxation and thus sizable 4d and 5d metal contributions. This contrast between the oneand three-electron reduction products is also confirmed immediately by the larger *g* anisotropy, ∆*g*, of the latter, the spectra being distinguished by three well-separated *g* components (Figure 2, Table 2). The EPR spectra shown here also illustrate that—depending on the electrolysis conditions—both paramagnetic products can be detected simultaneously, owing to the EPR electrolysis cell design,²⁷ which allows for only slow vertical

diffusion in a narrow tube.
Unfortunately, hyperfine coupling from metals $(^{103}Rh$ ($I =$ U_2 , 100% natural abundance) or ^{191,193}Ir ($I = 3/2$, 37.3 and 62.7%) natural abundance))²⁹ or other nuclei such as $14N$ could not be detected except for the previously described^{20,21} $[(\eta^5 - C_5M_e)^2]$ $CIRh(\mu\text{-bpip})RhCl(\eta^5\text{-}C_5Me_5)]^+$ system. The other paramagnetic species exhibit broader lines due to unresolved hyperfine structure and higher *g* anisotropy (Table 2).

Although the chloride-containing radicals invariably exhibit ∆*g* values smaller than those of the chloride-free species $[(\eta^5$ -C₅Me₅)M(μ -L)M(η^5 -C₅Me₅)]⁺, the ratio between pertinent ∆*g* values is much larger for the dirhodium systems (about 20) than for corresponding diiridium compounds (ca. 3.5). Alternatively, the difference between corresponding Rh and Ir species is very pronounced for the radicals $[(\eta^5 - C_5M_{\text{e}_5})\text{Cl}M(\mu - L)M\text{Cl}$ -

Figure 2. X band EPR spectra of $[(\eta^5 - C_5M_e)Rh(\mu - b\dot{p})Rh(\eta^5 - C_5 - \dot{p})]$ Me5)]+ (upper spectrum) and [(*η*⁵ -C5Me5)Ir(*µ*-bpip)Ir(*η*5-C5Me5)]+ (lower spectrum), generated from cathodic three-electron reduction of precursors $[(η⁵-C₅Me₅)CIM(μ-bpip)MCl(η⁵-C₅Me₅)](PF₆)₂ in DMF/$ 0.1 M Bu₄NPF₆. The signals marked "x" are due to the respective [(*η*5-C5Me5)ClM(*µ*-bpip)MCl(*η*5-C5Me5)]⁺ intermediates.

Table 2. *g* Factors of Paramagnetic Dirhodium and Diiridium Species*^a*

М	L	g_1	82	83	Δg	$\langle g \rangle$		
$[(\eta^5$ -C ₅ Me ₅)ClM(μ -L)MCl(η^5 -C ₅ Me ₅)] ⁺								
Rh	bpip	1.9975	1.9975	1.990	0.0076	1.995^b		
Ir	bpip	2.009	2.000	1.916	0.093	1.975		
$[(\eta^5-C_5Me_5)M(\mu-L)MC1(\eta^5-C_5Me_5)]^{2+}$								
Rh	bxip	2.015	2.001	1.996	0.019	2.004		
Ir	bxip	2.0411	2.0068	1.9362	0.1049	1.995		
$[(\eta^5 - C_5Me_5)M(\mu - L)M(\eta^5 - C_5Me_5)]^+$								
Rh	bpip	2.1656	2.0474	1.9516	0.2140	2.055		
Rh	bxip	2.167	2.043	1.950	0.217	2.053		
Ir	bpip	2.225	2.055	1.905	0.320	2.061		
Ir	bxip	2.227	2.048	1.910	0.317	2.061		

^a Electrogenerated through cathodic reduction of precursors (Scheme 1) in CH3CN/0.1 M Bu4NPF6 (bpip complexes) or DMF/0.1 M Bu4NPF6 (bxip systems). *g* values from simulations of spectra were taken at 3.5 K. *^b* Hyperfine resolution (see ref 21).

 $(\eta^5$ -C₅Me₅)]⁺ and $[(\eta^5$ -C₅Me₅)M(μ -L)MCl(η^5 -C₅Me₅)]²⁺ but less marked for complexes $[(\eta^5 \text{-} C_5 \text{Me}_5) \text{M}(\mu \text{-} L) \text{M}(\eta^5 \text{-} C_5 \text{Me}_5)]^+$. This result is somewhat unexpected, given the distinctly higher spin-orbit coupling factors of iridium (5d) relative to corresponding rhodium $(4d)$ states.²⁹ For instance, the organometallic d^{5}/d^{6} mixed-valent ions $[(R_{3}P)_{2}(OC)_{3}M(\mu-L)M(CO)_{3}(PR_{3})_{2}]^{+}$, $L =$ pyrazine, $R =$ isopropyl, and $M =$ Mo, W, showed a much more pronounced difference between the 4d system $(M = Mo:$ g_{\perp} = 2.0655, g_{\parallel} = 1.9755) and the 5d analogue (M = W: g_1 = 2.197, $g_2 = 2.158$, $g_3 = 1.931$.^{5,6} However, a comparable EPR behavior as described here has been reported for M^IM^{II} species with direct metal-metal interaction.³⁰

Apart from the metal-dependent *g* anisotropy, ∆*g*, there are few EPR differences between bpip and bxip containing compounds $[(\eta^5$ -C₅Me₅)M(μ -L)M(η^5 -C₅Me₅)]⁺, and remarkably, the calculated average $\langle g \rangle$ is essentially identical at about $g = 2.055$ for all four systems (Table 2). In contrast, the chloridecontaining radicals exhibit a much greater variety in terms of *g* splitting symmetry (axial or rhombic), ∆*g* (strongly increased for Ir systems, larger for bxip than for bpip species), and $\langle g \rangle$

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(lower for Ir and bpip species). The main reason for this variability lies in the different composition, that is, $[(\eta^5{\rm -}C_5Me_5)-$ ClM(μ -bpip)MCl(η ⁵-C₅Me₅)]⁺ vs $[(\eta$ ⁵-C₅Me₅)M(μ -bxip)MCl- $(\eta^5$ -C₅Me₅)²⁺. Apparently, the dissociation of one chloride already causes increased metal participation at the singly occupied MO (SOMO), indicated by larger ∆*g* and 〈*g*〉 values. Part of the variability may also lie in structural differences between bpip and bxip complexes, the (partially) chloridecontaining species with approximately tetrahedral configuration at the metal probably show a more sensitive response in terms of distortion toward the steric requirements resulting from the 2,6-dimethylphenyl substituent in bxip. As for [∆]*g*, the spinorbit coupling constants of M(III) ions are higher than those of the corresponding metals in lower oxidation states, which could explain the strong increase of ∆*g* on going from Rh(III) to Ir(III) radicals.

Isotropic or axial *g* component splitting is common for π radicals and their complexes, whereas rather symmetrical rhombic splitting has been observed for metal/ligand mixed situations at lower symmetries.¹⁴ The higher $\langle g \rangle$ values for $[(\eta^5$ -C₅Me₅)M(μ -L)M(η^5 -C₅Me₅)⁺ relative to those of $[(\eta^5$ -C₅- $Me₅$)ClM(μ -L)MCl(η ⁵-C₅Me₅)⁺ reflect a situation known for most d^{5}/d^{6} mixed-valent species, that is, occupied MOs lying rather close to the SOMO, which also causes low-energy IVCT transitions (HOMO \rightarrow SOMO; see below).^{5,6} On the other hand, the configuration in $[(\eta^5 \text{-} C_5\text{Me}_5) \text{C} \text{M}(\mu \text{-} L) \text{M} \text{C} \text{M}(\eta^5 \text{-} C_5\text{Me}_5)]^+$ produces a singly occupied L-centered MO with closer-lying unoccupied MOs (d or other π^* MOs of L), which results in lowered *g* values relative to g (electron) = 2.0023.³¹

The paramagnetic product $[(\eta^5 \text{-} C_5\text{Me}_5) \text{M}(\mu \text{-} \text{L}) \text{M}(\eta^5 \text{-} C_5 \text{Me}_5)]$ ⁻ resulting from the last cyclovoltammetrically observed reduction process could not be studied by EPR because of slow dissociation at the very negative potentials required.

UV/Vis/NIR Spectroelectrochemistry of Paramagnetic Intermediates. The appearance of absorptions in the longwavelength region of the visible spectrum or even in the nearinfrared (NIR) is a typical phenomenon associated with mixed valency. $1-13$ These transitions can be approximated as intervalence charge-transfer (IVCT) transitions for a localized formulation $L_n M^k(\mu\text{-}L)M^{k+1}L_n$ or as $\pi \to \pi^*$ transitions of a dimetallic π system in a delozalized description. Numerical approximations as the one by $Hush^{1-4,32}$ have been used to relate these transitions (energy, oscillator strength) with electronic coupling energies.

Using spectroelectrochemical techniques with an optically transparent thin-layer electrochemical cell, 19 we have studied the complexes by absorption spectroscopy in the visible and NIR region. The starting materials are characterized by ¹LMCT (Rh, Ir) and ${}^{3}LMCT$ (Ir) transitions in the visible (LMCT: ligand-to-metal charge transfer).

The one-electron-reduced forms of the bxip complexes are not sufficiently persistent on the time scale needed for conventional spectroelectrochemical measurements (>1 min). The electrolysis at the first reduction peak leads directly to the twoelectron reduction product because of a disproportionation reaction (see cyclic voltammetry chapter above), and consequently, the second reduction peak (B in Figure 1) is suppressed. The three-electron reduction product $[(\eta^5-C_5Me_5)M(\mu-bxip)$ - $MCl(\eta^5-C_5Me_5)$ ⁺ can be generated when the potential is scanned slowly at the beginning of the overlapping peaks C and D. The one-electron-reduced cations $[(\eta^5 - C_5M_e) \text{C} \text{M}(\mu$ bpip)MCl($η$ ⁵-C₅Me₅)]⁺ exhibit only intraligand (IL) and ligand-

Figure 3. Spectroelectrochemical response for the transitions $[(\eta^5{\text{-}}C_5{\text{-}}\eta^5{\text{-}}C_6{\text{-}}\eta^5{\text{-}}\eta^5{\text{-}}\eta^5{\text{-}}\eta^5{\text{-}}\eta^5{\text{-}}\eta^5{\text{-}}\eta^5{\text{-}}\eta^5{\text{-}}\eta^5{\text{-}}\eta^5{\text{-}}\eta^5{\text{-}}\eta^5{\text{-}}\eta^5{\text{-}}\eta^5{\text{-}}\eta^5{\text{-}}\eta^5{\$ $Me_5)Rh(\mu$ -bxip)RhCl(η ⁵-C₅Me₅)]⁺ \rightarrow [(η ⁵-C₅Me₅)Rh(μ -bxip)Rh(η ⁵-C₅-Me₅)]⁺ (upper spectrum) and [($η$ ⁵-C₅Me₅)Rh(*μ*-bxip)Rh($η$ ⁵-C₅Me₅)]^{(+)→(0)} (lower spectrum) in the UV/vis/NIR region (DMF/0.1 M Bu₄NPF₆).

to-metal charge transfer (LMCT) transitions in the visible according to their formulation as radical complexes.³¹ On the other hand, the chloride-free cations $[(\eta^5{\text{-}}C_5Me_5)M(\mu{\text{-}}L)M(\eta^5{\text{-}}C_5Me_5)]$ (C_5Me_5) ⁺ exhibit a series of at least two bands of medium intensity ($\epsilon \approx 2000 \text{ M}^{-1} \text{ cm}^{-1}$) in the NIR, viz., in the region between 900 and 1800 nm (Figure 3, Table 3). For comparison, the Creutz-Taube ion absorbs at 1585 nm in DMF solution.23

The separation between individual components of about $1100-1600$ cm⁻¹ supports their assignment as vibrational components of one allowed electronic transition.³³⁻³⁶ Chelate complexes of the $(\eta^5$ -C₅Me₅)M fragments are known to exhibit such phenomena because of their rigidity and relatively small number of rotational and vibrational degrees of freedom. $33-36$

As has been observed previously,²¹ the IVCT energy is higher for the iridium species because of the stronger coupling of the 5d metal centers. The difference between bpip and bxip compounds is marginal. The neutral $[(\eta^5 \text{-} C_5\text{Me}_5)M(\mu\text{-}L)M(\eta^5$ - C_5Me_5] complexes also exhibit a series of long-wavelength bands in the near-IR albeit at higher energies than the corresponding cations. According to previous findings for mononuclear compounds ($η^5$ -C₅Me₅)M(α-diimine)^{24-26,33-35} there is extensive charge transfer to the chelate ligand, which allows for possible mixed-valent resonance structures such as $[(\eta^5-C_5-$ Me5)MI (*µ*-L•-)MII(*η*5-C5Me5)].

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Table 3. Absorption Maxima*^a* of Complexes from UV/Vis/NIR Spectroelectrochemistry*^b*

μ -L = bpip			μ -L = bxip				
$M = Ir$	$M = Rh$	$M = Ir$	$M = Rh$				
$[(\eta^5 - C_5Me_5)CIM(\mu - L)MC1(\eta^5 - C_5Me_5)]^{2+}$							
320 (11.6)	321 (14)	292(7.0)	310(13)				
380 sh (7.7)	370 (11)		400(8.6)				
504 (10.3)	465(10)	514 (9.5)	474 (7.3)				
610 $(2.9)^c$		602^c (5.4)					
$[(\eta^5-C_5Me_5)CIM(\mu-L)MCl(\eta^5-C_5Me_5)]^{*+}$							
360 (10.5)	330 (14)	ℓ	e				
387 (10.7)	402(14)						
480 (11.6)	451 (13)						
533 (8.7)	546 (9)						
636 (7.6)	690(4)						
860 $(0.1)^c$							
	$[(\eta^5 - C_5Me_5)M(\mu - L)MC](\eta^5 - C_5Me_5)]^+$						
300 (17.2)	321 (21)	297 (19)	328 (17)				
382 (16.6)	429 (10)	372(15)	422(8.5)				
530 (18.2)	578 (12)	534 (16)	588 (10)				
625(11.0)	691 (18)	629(10)	702 (16)				
$[(\eta^5 - C_5Me_5)M(\mu - L)M(\eta^5 - C_5Me_5)]^+$							
285^{d}	311 (21)	340^{d}	313(17)				
	362(11)		362(11)				
378	452(7)	376	456 (7.4)				
495 sh		491					
530	578 (7)	527	578 (5.4)				
		626					
680	698 (29)	682	706 (28)				
	805 (12)		820 (15)				
825	877 (9)	825	875 (13)				
1070	1142(0.9)	925					
1225	1396(1.1)	1234	1406(0.8)				
1415	1708 (3.2)	1405	1717(2.3)				
$[(\eta^5\text{-}C_5Me_5)M(\mu\text{-}L)M(\eta^5\text{-}C_5Me_5)]$							
344 (14.7)	298 (19)	344(16)	298 (19)				
412 (13.1)	383 (17)	412 (14)	385 (18)				
527 (11.8)	523(7)	528 (13)	522 (6.9)				
703 (19.5)	698 (33)	702 (29)	698 (34)				
926 (0.9)	886 (6)	928(2.3)	902 (3.4)				
1050(1.9)	1007(5)	1008(3.1)	1016(5.7)				
1205(1.8)	1136(5)	1199 (1.9)	1142 (5.7)				

^{*a*} Absorption maxima λ_{max} in nm (molar extinction coefficients $\epsilon \times$ 10^{-3} in M⁻¹ cm⁻¹). ^{*b*} OTTLE cell: CH₃CN/0.1 M Bu₄NPF₆ for L = hpin and DMF/0.1 M Bu₄NPF₆ for L = hxin ^c³LMCT ^d Observed bpip and DMF/0.1 M Bu₄NPF₆ for L = bxip. ^c ³LMCT. ^{*d*} Observed within comproportionation equilibrium. *^e* Not observed because of low stability (see text).

Concluding Remarks

One-electron uptake by complexes $[(\eta^5-C_5M_{\text{e}_5})\text{Cl}M(\mu-L)$ - $MCl(\eta^5-C_5Me_5)$ ²⁺ with 2,5-diiminopyrazine bridging ligands L produces $L^{\bullet-}$ -containing species after reversible (L = bpip)

or partially irreversible processes $(L = bxi)$. Addition of two more electrons yields chloride-free paramagnetic species $[(\eta^5$ -C₅Me₅)M(μ -L)M(η^5 -C₅Me₅)]⁺ whose EPR data (*g* anisotropy) and spectroelectrochemical characteristics (near-infrared IVCT absorption) suggest a metal-metal mixed-valent situation involving divalent rhodium 37 or iridium.³⁰ The comparatively small *g* anisotropy especially for the iridium systems indicates some $(\eta^5$ -C₅Me₅) coligand participation in the SOMO due to partial covalent bonding; similar effects were noted for organometallic $Mo_{2}^{0,I}$ and $W_{2}^{0,I}$ compounds.^{5,6} In the absence of EPR hyperfine or direct IR vibrational data, the question⁷ of valence localization or delocalization cannot be unambiguously answered; the small potential splitting of less than 150 mV between neigboring redox waves is more compatible with a localized description. The oxidation state and d electron configuration assignment to the metals is also not immediately obvious for compounds $[(\eta^5-C_5Me_5)M(\mu-L)M(\eta^5-C_5Me_5)]^+;$ the ability of $(\eta^5$ -C₅Me₅)M (M = Rh,Ir) fragments to transfer large amounts of negative charge to bound α -diimine acceptor sites^{24-26,33-35} points to a resonance situation

$$
[(\eta^5 - C_5Me_5)M^{II}(\mu - L)M^I(\eta^5 - C_5Me_5)]^+ \rightleftharpoons
$$

\n
$$
[(\eta^5 - C_5Me_5)M^{II}(\mu - L^{\bullet})M^{II}(\eta^5 - C_5Me_5)]^+ \rightleftharpoons
$$

\n
$$
[(\eta^5 - C_5Me_5)M^{III}(\mu - L^{2-})M^{II}(\eta^5 - C_5Me_5)]^+
$$
 (1)

that was similarly observed for diruthenium (II,III) complexes.¹⁴

A π acceptor ligand mediated d^{7}/d^{8} (M^{II}/M^I) mixed-valent configuration would be compatible with the established electronic structure of the $(\eta^5$ -C₅Me₅)M(α -diimine) moiety.²⁴ In contrast, square planar diplatinum(III,II) complexes with their different d orbital ordering are less favored for π ligandmediated metal-metal interaction.¹⁸

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Supporting Information Available: Figures of cyclic voltammograms and EPR spectra. This material is available free of charge via the Internet at http://pubs.acs.org.

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