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Multistep Oxidation of Diethynyl Oligophenylamine-Bridged Diruthenium and Diiron Complexes

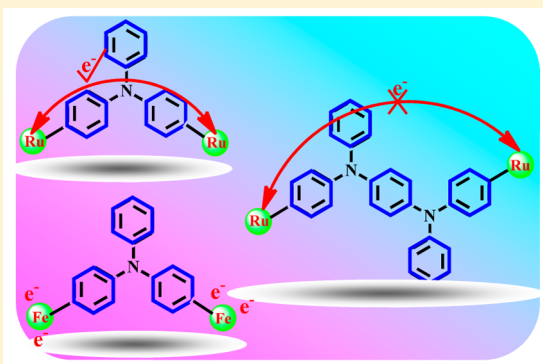
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[§] Supporting Information

ABSTRACT: Homo-dinuclear nonlinear complexes $[\{M(dppe)-Cp^*\}_2\{\mu-(C\equiv C)_2X\}]$ ($dppe = 1,2$ -bis(diphenylphosphino)ethane; $Cp^* = \eta^5-C_5Me_5$; $X =$ triphenylamine (TPA), $M = Ru$ (**1a**) and Fe (**1b**); $X = N,N,N',N'$ -tetraphenylphenylene-1,4-diamine (TPPD), $M = Ru$ (**2a**)) were prepared and characterized by 1H , ^{13}C , and ^{31}P NMR spectroscopy and single-crystal X-ray diffraction (**1a**, **2a**). Attempts to prepare the di-iron analogue of **2a** were not successful. Experimental data obtained from cyclic voltammetry, square wave voltammetry, UV–vis–NIR (NIR = near-infrared) spectro-electrochemistry, and very informative IR spectro-electrochemistry in the $C\equiv C$ stretching region, combined with density functional theory calculations, afford to make an emphasizing assessment of the close association between the metal–ethynyl termini and the oligophenylamine bridge core as well as their respective involvement in sequential one-electron oxidations of these complexes. The anodic behavior of the homo-bimetallic complexes depends strongly both on the metal center and the length of the oligophenylamine bridge core. The poorly separated first two oxidations of di-iron complex **1b** are localized on the electronically nearly independent Fe termini. In contrast, diruthenium complex **1a** exhibits a significantly delocalized character and a marked electronic communication between the ruthenium centers through the diethynyl–TPA bridge. The ruthenium–ethynyl halves in **2a**, separated by the doubly extended and more flexible TPPD bridge core, show a lower degree of electronic coupling, resulting in close-lying first two anodic waves and the NIR electronic absorption of $[2a]^+$ with an indistinctive intervalence charge transfer character. Finally, the third anodic waves in the voltammetric responses of the homo-bimetallic complexes are associated with the concurrent exclusive oxidation of the TPA or TPPD bridge cores.



INTRODUCTION

In recent years mixed-valence (MV) states of binuclear complexes have attracted considerable attention,^{1,2} providing important model systems for intramolecular charge transfer processes and offering broad prospects for building highly functionalized molecules with interesting electronic and optoelectronic properties essential for molecular scale electroactive materials and devices.^{3–6} In this respect, numerous studies focused in the past decades on rigid and π -conjugated bridging ligands connecting two redox-active metallic termini, as the simplest models for the electron-transfer phenomena in the MV systems.^{7–10} Comprehensive studies have explored a wide range of systems designed to provide some insight into electron transfer over a long distance.¹¹ However, increasing the chain length to a modest degree decreased significantly solubility of the prototypical π -conjugated systems in common organic solvents, and the complexity of their syntheses essentially arose. These factors present obstacles for the relevant fundamental research and restrict the longitudinal extension and processability in electronic devices.¹¹ In this

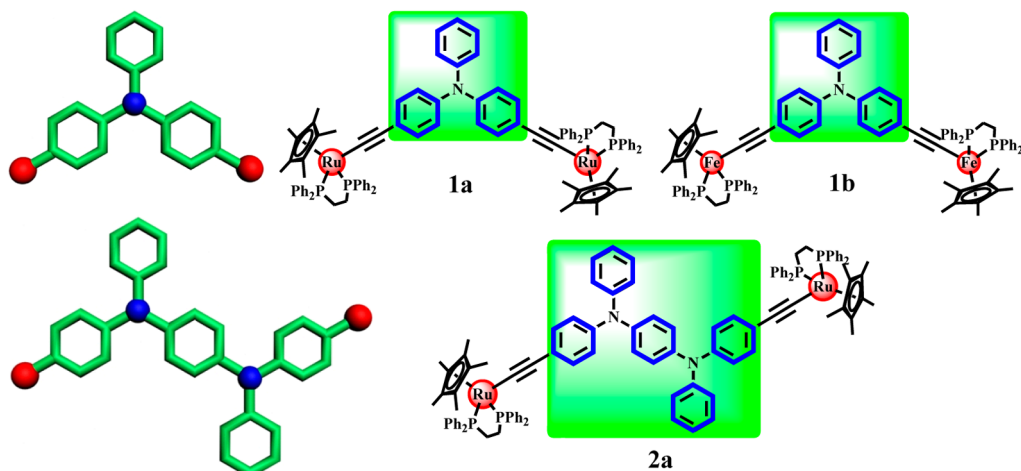
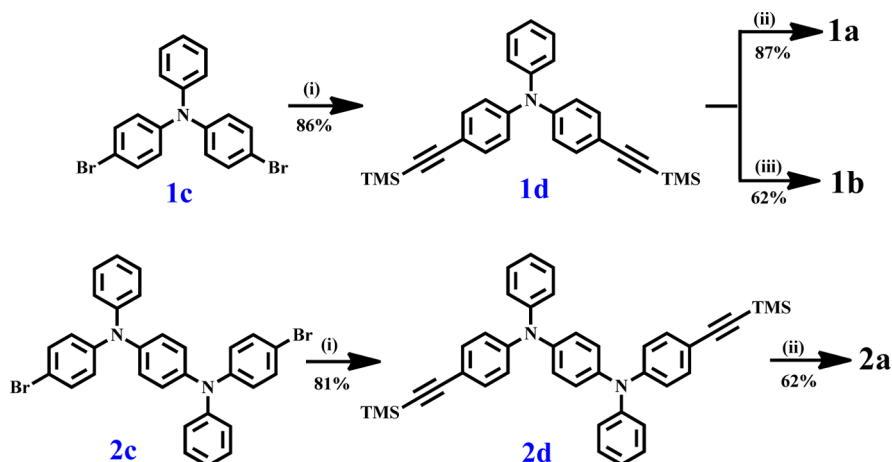
respect, continued search for novel or modified perspective systems to eliminate the above-mentioned drawbacks is very meaningful and challenging.

Triphenylamine (TPA) and other mono- and oligomeric triarylamine derivatives have widely been used as selective one-electron reductants, electrocatalysts, and hole-transporting materials in organic optoelectronic devices, such as photoconductors, photorefractive materials, and organic light-emitting devices. This is ascribed to their favorable redox properties, with one or more readily accessible oxidations and high stabilities of the corresponding radical cations with appropriate substitution patterns.^{12–14} Moreover, TPA has also been considered as an ideal redox center to study the intermolecular electron transfer processes in MV systems.^{15–17} Despite the efforts, reports on mixed-valence systems featuring the TPA unit as (a component of) a bridging ligand are relatively limited in number compared with the π -conjugated

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Chart 1. Studied Diethynyl Oligophenylamine-Bridged Homo-Bimetallic Ruthenium and Iron Complexes

Scheme 1. General Synthetic Routes to Complexes 1a, 1b, and 2a^a

^aReagents and conditions: (i) $[\text{Pd}(\text{PPh}_3)_4]$, CuI, Et_3N , TMSA, in THF; (ii) $[\text{RuCl}(\text{dppe})\text{Cp}^*]$, KF, MeOH/THF; (iii) K_2CO_3 in CH_3OH , $[\text{FeCl}(\text{dppe})\text{Cp}^*]$, NaBPh_4 , $t\text{-BuOK}$.

systems,¹⁸ let alone a detailed discussion of their electronic properties.^{18d} In another aspect, a nonrigid TPA-based structure with appropriate substituents^{12–14} offers a viable solution to the problem of poorly soluble elongated systems and the difficulty in synthesizing large dendrimers. Notably, Onitsuka and co-workers synthesized successfully a series of ruthenium–acetylide dendrimers with the tris(4-ethynylphenyl)amine bridging ligand up to the second generation by using a convergent method.¹⁸ⁱ In this work we focus on TPA and scarcely reported tetraphenylphenylenediamine (TPPD) in the core of the diethynyl-terminated bridge. The external “ $\text{Ru}(\text{dppe})\text{Cp}^*$ ” and “ $\text{Fe}(\text{dppe})\text{Cp}^*$ ” (dppe = 1,2-bis(diphenylphosphino)ethane; Cp^* = $\eta^5\text{-C}_5\text{Me}_5$) units completing the studied homo-bimetallic chains (Chart 1) are particularly suited as reference redox-active termini in molecular wire models to study their MV properties, as documented by the previously reported work.^{2d–h,19} The assembly of multiple redox-active components was anticipated to exhibit peculiar electrochemical and photophysical properties. Hereinafter we detail our efforts to explore to which degree the redox, spectroscopic, and bonding properties and electronic communication in the investigated homo-bimetallic series are affected by the varied combination of the oligophenylamine

bridge core and the metal centers. The input data for the discussion have been obtained by using controlled-potential voltammetry and UV–vis–NIR/IR (NIR = near-infrared) spectro-electrochemistry combined with density functional theory (DFT) calculations.

RESULTS AND DISCUSSION

Syntheses and Characterization. The general synthetic route to homo-dinuclear metal complexes **1a**, **1b**, and **2a** is outlined in Scheme 1. Bridge precursors **1d** and **2d** were obtained in yields higher than 80%, having used Pd/Cu-catalyzed Sonogashira coupling reactions between 4-bromo- N -(4-bromophenyl)- N -phenylaniline (**1c**), N^1,N^4 -bis(4-bromophenyl)- N^1,N^4 -diphenylbenzene-1,4-diamine (**2c**), and trimethylsilylacetylene (TMSA), respectively. Subsequently, the trimethylsilyl (TMS) termini in compounds **1d** and **2d** were deprotected with KF or K_2CO_3 to give the terminal bis(alkyne) and then reacted with $[\text{RuCl}(\text{dppe})\text{Cp}^*]$ and $[\text{FeCl}(\text{dppe})\text{Cp}^*]$ to obtain the corresponding target complexes **1a**, **2a**, and **1b**, respectively. Unfortunately, deprotected **2d** did not coordinate to the $\text{Fe}(\text{dppe})\text{Cp}^*$ termini by this procedure,²⁰ and the intended diethynyl-TPPD diiron complex (**2b**) was not obtained. Notably, no obvious differences have been observed

in comparative NMR responses of related diruthenium complexes **1a** and **2a**, specifically, the dppe signals in the ^1H and ^{31}P NMR spectra and the $\text{Ru}-\text{C}\equiv\text{C}$ signals in the ^{13}C NMR spectra.

X-ray Crystallography. The molecular structures of solid **1a** and **2a** were resolved by single-crystal X-ray diffraction. Pertinent diffraction parameters are given in Table S1 (see the Supporting Information). Important bond lengths (Å), bond angles (deg), and $\text{Ru}\cdots\text{Ru}$ distances (Å) in the crystal structures are collected in Table 1. The structures of **1a** and **2a** feature

Table 1. Selected Bond Lengths (Å), Angles (deg), and Inter-Ruthenium Distances (Å) in the Crystal Structures of Complexes **1a** and **2a**

complex 1a			
Ru(1)–C(37)	1.999 (3)	C(43)–C(44)	1.378 (4)
Ru(1)–P(1,2)	2.250 (9), 2.258 (8)	C(39)–C(44)	1.395 (5)
C(37)–C(38)	1.209 (4)	C(45)–C(46)	1.384 (4)
C(38)–C(39)	1.436 (4)	C(46)–C(47)	1.378 (5)
C(39)–C(40)	1.401 (5)	C(47)–C(48)	1.371 (5)
C(40)–C(41)	1.382 (4)	N(1)–C(42)	1.426 (3)
C(41)–C(42)	1.387 (4)	N(1)–C(45)	1.403 (5)
C(42)–C(43)	1.391 (4)	Ru(1)···Ru(2)	14.19
P(1)–Ru(1)–P(2)		83.3 (3)	
Ru(1)–C(37)–C(38)		174.9 (3)	
C(37)–C(38)–C(39)		175.6 (3)	
complex 2a			
Ru(1)–C(27)	1.991 (5)	C(36)–C(37)	1.343 (2)
Ru(1)–P(1,2)	2.246 (3), 2.248 (3)	C(37)–C(38)	1.390 (2)
C(27)–C(28)	1.214 (9)	C(38)–C(39)	1.394 (2)
C(28)–C(29)	1.448 (7)	C(39)–C(40)	1.348 (2)
C(29)–C(30)	1.375 (9)	C(35)–C(40)	1.402 (8)
C(30)–C(31)	1.360 (8)	C(41)–C(42)	1.363 (9)
C(31)–C(32)	1.390 (9)	C(42)–C(43)	1.393 (9)
C(32)–C(33)	1.378 (9)	C(41)–C(43)	1.390 (8)
C(33)–C(34)	1.389 (7)	N(1)–C(32)	1.428 (5)
C(29)–C(34)	1.392 (2)	N(1)–C(41)	1.424 (6)
C(35)–C(36)	1.392 (2)	Ru(1)···Ru(2)	20.68
P(1)–Ru(1)–P(2)		84.4 (3)	
Ru(1)–C(37)–C(38)		178.5 (3)	
C(37)–C(38)–C(39)		176.1 (3)	

approximately linear $\text{Ru}-\text{C}\equiv\text{C}-\text{C}$ moieties, the pairs of $\text{Ru}-\text{C}\equiv\text{C}$ and $\text{C}\equiv\text{C}-\text{C}$ angles being 174.9° , 175.6° and 178.5° , 176.1° , respectively. As seen in Figure 1 and Figure S1(A) (Supporting Information), the crystallographic analysis manifests typical pseudo-octahedral geometry around the ruthenium centers. The $\text{C}\equiv\text{C}$ bond lengths [1.209 Å (**1a**) and 1.214 Å (**2a**)] comply with a carbon–carbon triple bond, and the bonding parameters of the $\text{Ru}(\text{dppe})\text{Cp}^*$ group, including the $\text{Ru}-\text{C}$ and $\text{Ru}-\text{P}$ distances and $\text{P}-\text{Ru}-\text{P}$ angles, compare well with the data reported for a range of similar systems.^{21,22} Elongation of the bridging ligand core from monoamine (TPA) in **1a** to diamine (TPPD) in **2a** caused the $\text{Ru}(1)\cdots\text{Ru}(2)$ distance to increase considerably from 14.19 to 20.68 Å. These data are consistent with the corresponding calculated results presented hereinafter.

Some additional interesting features can be found in the crystal packing (Figure S1(B)). Complex **2a** shows typical lamellar packing characteristics with the parallel arrangements of the bridge and metallic termini in each layer from the side view of the lamellar structure, the benzene rings 1, 3 and 2, 4 of the diamine bridge core (Figure S1(B)) being oriented in opposite directions and arranged in parallel with each other.

Electrochemical Properties. The anodic behavior of complexes **1a**, **1b**, and **2a** was investigated by cyclic voltammetry (CV) and square-wave voltammetry (SWV) in deaerated dichloromethane containing $1 \times 10^{-1}\text{ M}$ $n\text{-Bu}_4\text{NPF}_6$ as the supporting electrolyte (Figure 2). The TMS-terminated precursors **1d** and **2d** (Scheme 1) were also studied for comparison (see Supporting Information, Figure S2). The relevant electrochemical data are summarized in Table 2.

Complexes **1a** and **1b** exhibit three successive oxidation steps. The anodic behavior of **2a** is more complex, and only the first four anodic processes were investigated. Comparing the electrochemical behavior of diruthenium complexes **1a** and **2a**, the potentials of their first three anodic steps ($E_{1/2}(1,2,3)$) are similar. The main difference arises in the wave splitting $\Delta E_{1/2}(1-2)$ and the thermodynamic stability (comproportionation) constant K_c of $[\mathbf{2a}]^+$, which is considerably smaller than that determined for $[\mathbf{1a}]^+$ (Table 2). This result has likely its origin in the extended bridge core in **2a** and limited electronic communication between two redox active $\text{Ru}-\text{C}\equiv\text{C}$ units, resembling in this respect an earlier reported series of

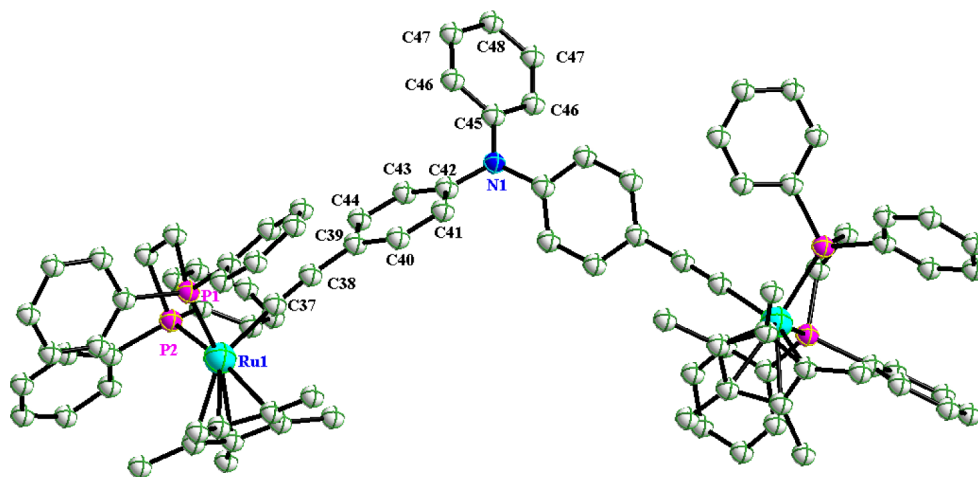


Figure 1. X-ray crystal structure of **1a** shown with thermal ellipsoids at the 50% probability level. Hydrogen atoms were omitted for clarity. Full crystallographic details are given in Supporting Information. CCDC No. 1435472.

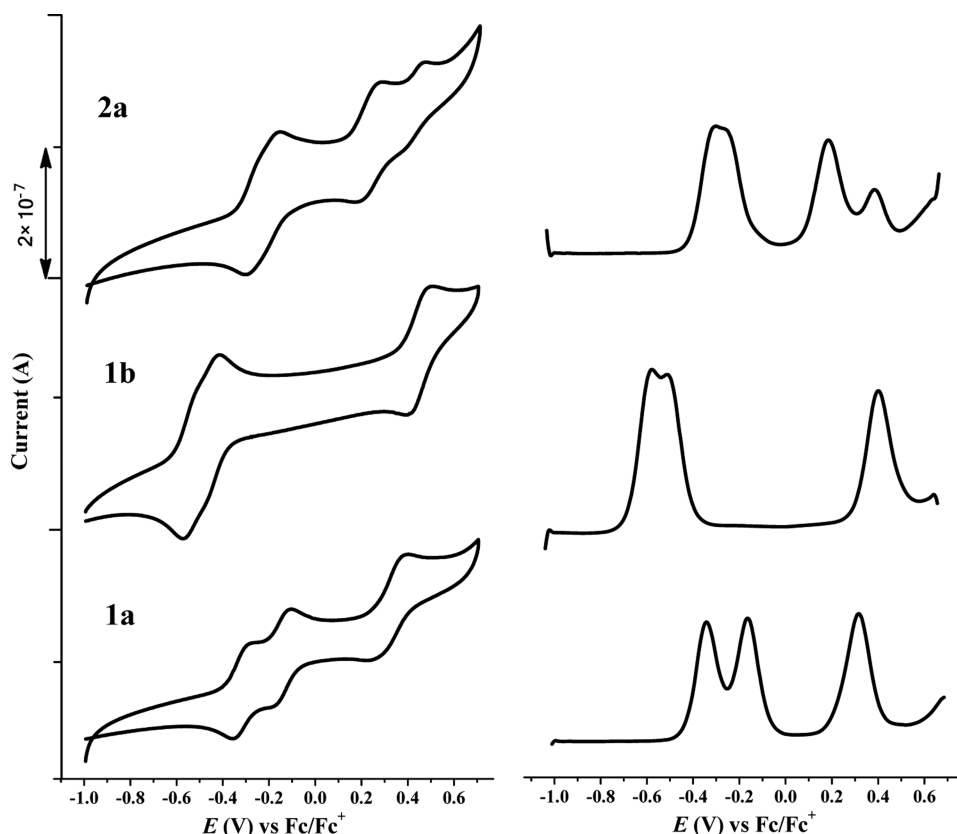


Figure 2. (left) CV of complexes **1a**, **1b**, and **2a** in $\text{CH}_2\text{Cl}_2/n\text{-Bu}_4\text{NPF}_6$ at $\nu = 50 \text{ mV s}^{-1}$. (right) Corresponding SWV of complexes **1a**, **1b**, and **2a** at $f = 10 \text{ Hz}$ and $t_p = 25 \text{ mV}$.

Table 2. Electrochemical Data for Complexes 1a, 1b, and 2a, TMS-Terminated Reference Compounds 1d and 2d, a Dinuclear Vinyl–Ruthenium Complex Related to 1a, and Reference Phenylamines^a

compound	$E_{1/2}(1) \text{ (V)}$	$E_{1/2}(2) \text{ (V)}$	$E_{1/2}(3) \text{ (V)}$	$E_{1/2}(4) \text{ (V)}$	$\Delta E_{1/2}(1-2) \text{ (mV)}$	K_c^b
1a	−0.36	−0.18	0.30		180	1.1×10^3
Ru–vinyl^c	−0.17	0.15	0.59		320	3.8×10^5
1b	−0.56	−0.50	0.41		60	10
1d	0.66					
TPA^d	0.70 ^{e,f}					
2a	−0.31	−0.26	0.19	0.39	50	7
2d	0.21	0.68				
TPPD^d	0.14 ^f	0.63				

^aThe anodic potentials are referenced against the standard ferrocene/ferrocenium (Fc/Fc^+) redox couple. ^bThe comproportionation constants K_c were estimated using the expression $K_c = \exp(\Delta E/25.69 \text{ mV})$ at 298 K, with input data recorded at $\nu = 50 \text{ mV s}^{-1}$. ^c $[\{\text{Cl}(\text{CO})(\text{P}(\text{t}\text{Bu})\text{R}_3)_2\text{Ru}-\text{CH}=\text{CH}-\text{C}_6\text{H}_4-\}_2\text{N}-\text{C}_6\text{H}_4\text{OCH}_3]$. ^dReference 18f. ^eIrreversible anodic peak potential. ^fPotential determined against an Ag/AgCl reference electrode; $E_{1/2}(\text{Fc}/\text{Fc}^+) = +0.43 \text{ V}$ versus Ag/AgCl .

oligothiophene-bridged diruthenium diethynyl complexes.²¹ The additional fourth anodic wave of **2a** at a higher potential can safely be ascribed to the second oxidation of the diamine core, based on comparison with the anodic potentials of the TMS-terminated diethynyl TPPD reference compound **2d**. The obvious difference in the composition of **1a** and **1b** is the metallic redox center. Interestingly, the replacement of ruthenium in **1a** with iron in **1b** resulted in a slightly negative shift of the first anodic wave (at $E_{1/2}(1)$) and significantly decreased comproportionation constant K_c (Table 2). It will be shown in the spectro-electrochemical section that the formally $\text{Fe}(\text{II})$ centers in the terminal positions of **1b** oxidize in a more localized fashion compared to the $\text{Ru}(\text{II})$ derivative, **1a**, as also encountered in the literature for related compounds.^{3c,8b,18h,23}

Accordingly, the two formally $\text{Fe}(\text{II})$ centers in **1b** communicate poorly through the twisted ethynyl–phenylene–N–phenylene–ethynyl bridge, in line with the small $\Delta E_{1/2}(1-2)$ and K_c values for **[1b]⁺**. Consequently, it was hardly possible to record IR and UV–vis–NIR spectra of “pure” **[1b]⁺** in the course of the corresponding spectro-electrochemical experiments described in the following section. A less severe restriction applies in this regard for **[2a]⁺**.

A significantly greater involvement of the TPA core in the electrochemical oxidation has been reported^{18d} for a bridged dinuclear vinyl–ruthenium complex, causing a stronger electronic interaction between the ruthenium–vinyl linker subunits in the corresponding one-electron oxidized cation and a larger K_c value (Table 2). The different auxiliary ligands at the

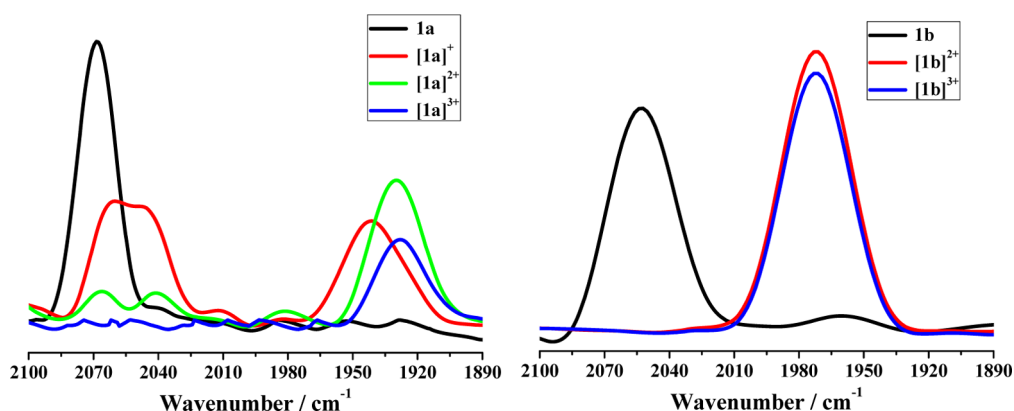


Figure 3. IR spectra recorded in the $\nu(\text{C}\equiv\text{C})$ region for complexes **1a** and **1b** in different oxidation states generated in $\text{CH}_2\text{Cl}_2/1 \times 10^{-1} \text{ M } n\text{-Bu}_4\text{NPF}_6$ at 298 K within an OTTLE cell. For the chemical oxidation of **1b** to $[\mathbf{1b}]^+$ see [Supporting Information](#), Figure S6.

193 ruthenium center, specifically, *trans*- $\text{RuCl}(\text{CO})\{\text{P}(i\text{-Pr})_3\}_2$, also
194 play a role in the control of the electronic conjugation between
195 the TPA bridge core and the ruthenium–linker (vinyl vs
196 ethynyl) subunits. For example, smaller differences in the
197 anodic behavior exist between TPA-based mono- and trinuclear
198 vinyl–ruthenium^{18d} and ethynyl–ruthenium²⁴ complexes with
199 similar ruthenium coordination sites.

200 **IR and UV–vis–NIR Spectro-Electrochemistry.** IR and
201 UV–vis–NIR spectro-electrochemical studies were undertaken
202 to get insight into the nature of the physical and electronic
203 structures in different oxidation states within the investigated
204 three redox series.

205 The IR spectra of neutral complexes **1a**, **1b**, and **2a** are
206 characterized by a strong single $\nu(\text{C}\equiv\text{C})$ band ([Figure 3](#),
207 [Figure S3](#) ([Supporting Information](#)), and [Table 3](#)). By contrast,

individual ethynyl–ruthenium moieties in the oxidation
process.

The second anodic step toward $[\mathbf{1a}]^{2+}$ led to the
disappearance of both $\nu(\text{C}\equiv\text{C})$ bands close to the wave-
number of the neutral parent state, while a new single band at
1930 cm^{-1} gained intensity. This behavior suggests the
complete oxidation of both $\text{Ru}-\text{C}\equiv\text{C}$ centers in $[\mathbf{1a}]^{2+}$.
Importantly, no prominent low-energy shift of the $\nu(\text{C}\equiv\text{C})$
mode was observed for the subsequent anodic generation of the
tricationic product $[\mathbf{1a}]^{3+}$, characterized by a single absorption
band marking a symmetric electronic structure. The very little
 $\nu(\text{C}\equiv\text{C})$ shift compares well with the invariable $\nu(\text{C}\equiv\text{C})$
wavenumber encountered for the TMS-terminated diethynyl
TPA reference **1d** and its monocation ([Supporting](#)
[Information](#), [Figure S4](#)), suggesting the dominantly TPA-
localized one-electron oxidation of $[\mathbf{1a}]^{2+}$.

The anodic conversion of **2a** to $[\mathbf{2a}]^{2+}$ was marked by
gradual disappearance of the parent $\nu(\text{C}\equiv\text{C})$ absorption at
2068 cm^{-1} and the growth of a new absorption band at 1934
 cm^{-1} belonging to the dication ([Supporting Information](#),
[Figure S3](#)). The red shift of 134 cm^{-1} is close to that of 138
 cm^{-1} observed for the oxidation of **1a** to $[\mathbf{1a}]^{2+}$ ([Table 3](#)),
indicating comparable participation of the ethynyl linkers in the
oxidation of **2a** to the symmetric dication, in line with the
theoretical description (see below). The small separation of the
first two anodic waves of **2a** implies that singly oxidized $[\mathbf{2a}]^+$
can hardly be observed in the pure form. The IR spectrum in
[Figure S3](#) corresponding to the redox equilibrium with the
highest concentration of $[\mathbf{2a}]^+$ was selected with the aid of the
characteristic NIR electronic absorption of the monocation (see
below) simultaneously monitored in the course of the careful
anodic electrolysis within the OTTLE cell. It can safely be
concluded that $[\mathbf{2a}]^+$ exists only in a single conformation
detectable on the time scale inherent to IR spectroscopy,
differently from $[\mathbf{1a}]^+$ (see above). The IR $\nu(\text{C}\equiv\text{C})$
absorption of $[\mathbf{2a}]^+$ overlaps with those of parent **2a** and
ultimately $[\mathbf{2a}]^{2+}$ ([Table 3](#)), rendering the monocation with
weak electronic coupling between the $\text{Ru}-\text{C}\equiv\text{C}$ termini not
exceeding that in $[\mathbf{1a}]^+$, in line with the corresponding CV
responses ([Table 2](#)).

The third anodic step producing stable $[\mathbf{2a}]^{3+}$ was not
accompanied by any $\nu(\text{C}\equiv\text{C})$ shift, causing merely a strongly
diminished intensity of the $\nu(\text{C}\equiv\text{C})$ band at 1934 cm^{-1}
([Figure S3](#)). This behavior reflects the dominant oxidation of
the TPPD bridge core in $[\mathbf{2a}]^{2+}$ to the corresponding radical
cation. This assignment is supported by the invariable $\nu(\text{C}\equiv\text{C})$

Table 3. Spectro-Electrochemically Determined $\nu(\text{C}\equiv\text{C})$
Wavenumbers (cm^{-1}) for $[\mathbf{1a}]^{n+}$, $[\mathbf{1b}]^{n+}$, $[\mathbf{1d}]^{n+}$ and $[\mathbf{2a}]^{n+}$,
 $[\mathbf{2d}]^{n+}$

complex	$n = 0$	$n = 1$	$n = 2$	$n = 3$	$n = 4$
$[\mathbf{1a}]^{n+}$	2068 (s)	2059 (m), 2046 (m), 1941 (m)	1930 (m)	1928 (w- m)	
$[\mathbf{1b}]^{n+}$	2052 (s)	^a	1972 (s)	1972 (s)	
$[\mathbf{1d}]^{n+}$	2065	2065			
$[\mathbf{2a}]^{n+}$	2068 (s)	2066, 1935 ^b	1934 (s)	1933 (w)	^c
$[\mathbf{2d}]^{n+}$	2066 (m)	2066 (m)	2066 (s)		

^aThe $\nu(\text{C}\equiv\text{C})$ absorption of $[\mathbf{1b}]^+$ is not reported due to the small K_c value and a low conversion evidenced by UV–vis–NIR spectroscopy ([Supporting Information](#), [Figure S6](#)). ^bThis absorption does not correspond to the pure form of $[\mathbf{2a}]^+$ due to small K_c but to the maximum conversion reached by electrochemical oxidation of **2a** determined by parallel UV–vis–NIR monitoring ([Supporting Information](#), [Figure S12](#)). ^cNo $\nu(\text{C}\equiv\text{C})$ band perceptible.

208 the $\nu(\text{C}\equiv\text{C})$ pattern of one-electron-oxidized $[\mathbf{1a}]^+$ is more
209 complex, consisting of a strongly shifted, fairly broad, and
210 slightly asymmetric absorption band at 1941 cm^{-1} and two
211 overlapping $\nu(\text{C}\equiv\text{C})$ bands at 2059 and 2046 cm^{-1} , that is,
212 near the absorption of parent **1a** at 2068 cm^{-1} . These spectral
213 changes comply with the presence of a symmetry-broken
214 radical cation $[\mathbf{1a}]^+$ on the time scale of IR spectroscopy ($1 \times$
215 10^{-11} to 1×10^{-12} s), existing in different conformations
216 (rotamers)²⁵ with slightly different participation of the

wavenumber in the TMS-terminated reference redox series $[2d]^{n+}$ ($n = 0, 1, 2$); see Table 3 and Supporting Information, Figure S4.

The poorly resolved first two anodic steps of **1b** (Table 2 and Figure 2) resemble the oxidation of **2a**. We note that the $\nu(\text{C}\equiv\text{C})$ shift induced by the oxidation of **1b** to $[1b]^{2+}$ is merely 80 cm^{-1} compared to 134 cm^{-1} for $2a \rightarrow [2a]^{2+}$, and 138 cm^{-1} for $1a \rightarrow [1a]^{2+}$ featuring the same molecular bridge (Table 3). This difference points to significantly more Fe-centered oxidation of the Fe–C $\equiv$ C termini, as also confirmed by DFT calculations (see below). The electronic interaction between the Fe centers in $[1b]^+$ through the TPA core becomes strongly limited, resulting in dominant disproportionation of the monocation to **1b** and $[1b]^{2+}$. Therefore, $[1b]^+$ is hardly detectable by IR spectroscopy (Table 3 and Supporting Information, Figures S5 and S6), and only the characteristic NIR absorption (Supporting Information, Figure S6) reveals its presence. The negligible change in the $\nu(\text{C}\equiv\text{C})$ wavenumber accompanying the third anodic step producing $[1b]^{3+}$ resembles the formation of $[1a]^{3+}$ (Figure 3), both bearing the TPA-core oxidation characteristics.

The UV–vis–NIR spectra of complexes **1a**, **1b**, and **2a** and TMS-terminated bridges **1d** and **2d** in the different oxidation states were recorded by using the spectro-electrochemical monitoring or stepwise chemical oxidation, as shown in Figures 4 and 5 and Supporting Information, Figures S7–S12. The relevant data are collected in Table 4.

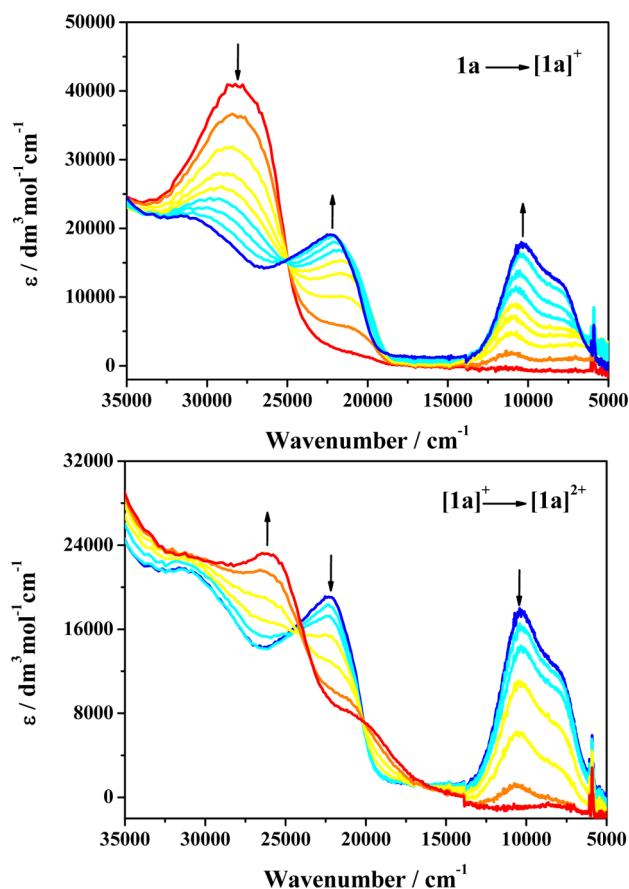


Figure 4. UV–vis–NIR spectral changes recorded during the oxidation of complex **1a** to $[1a]^+$ (top) and $[1a]^{2+}$ (bottom) in $\text{CH}_2\text{Cl}_2/1 \times 10^{-1}\text{ M } n\text{-Bu}_4\text{NPF}_6$ at 298 K within an OTTE cell.

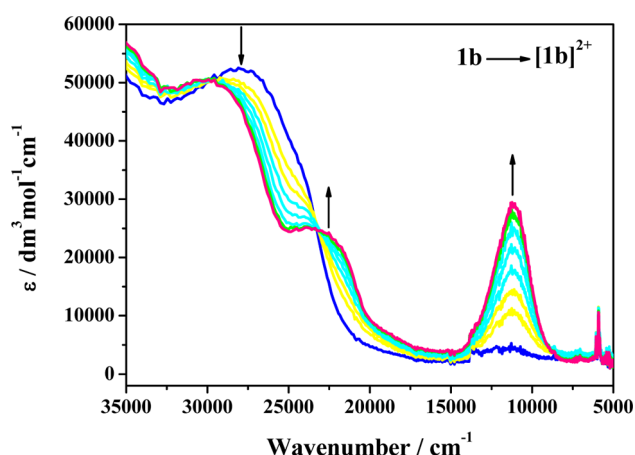


Figure 5. UV–vis–NIR spectral changes recorded during the oxidation of complex **1b** to $[1b]^{2+}$ in $\text{CH}_2\text{Cl}_2/1 \times 10^{-1}\text{ M } n\text{-Bu}_4\text{NPF}_6$ at 298 K within an OTTE cell.

Table 4. UV–Vis–NIR Electronic Absorption of Complexes **1a**, **1b**, **1d**, **2a**, **2d**, and Their Oxidized Forms in Dichloromethane/ $n\text{-Bu}_4\text{NPF}_6$

complex	$\tilde{\nu}_{\text{max}}$ (cm^{-1}) (ϵ_{max} ($\text{dm}^3\text{ mol}^{-1}\text{ cm}^{-1}$))
1a	28 100 (41 300)
$[1a]^+$	22 200 (19 500), 10 200 (18 100), 7800 (12 400)
$[1a]^{2+}$	26 000 (23 300), 20 500 (7600)
$[1a]^{3+}$	23 800 (21 900), 8300 (10 400)
1b	27 500 (52 700)
$[1b]^+$	^a
$[1b]^{2+}$	29 600 (50 800), 22 500 (24 700), 11 200 (29 400)
$[1b]^{3+}$	16 800 (32 000), 9700 (19 000)
1d	41 400 (28 600), 28 500 (40 700)
$[1d]^+$	13 500 (12 200)
2a	29 200 (56 600)
$[2a]^+$	^b 30 400, 21 600, 10 300, 8400sh
$[2a]^{2+}$	29 800 (44 700), 22 000 (8500)
$[2a]^{3+}$	23 900 (22 700), 8700 (22 200)
2d	29 600 (43 200)
$[2d]^+$	23 400 (23 600), 10 400 (19 100), 8100 (12 400)
$[2d]^{2+}$	38 600 (28 300), 13 900 (44 800)

^a Detectable by an unresolved weak NIR absorption between 10 000 and 4000 cm^{-1} (see Supporting Information, Figure S6). ^b Molar absorptivity not reported.

The diethynyl–TPA complex **1a** exhibits a pronounced broad absorption band at $\sim 2.8 \times 10^4\text{ cm}^{-1}$, most likely stemming from the $\pi \rightarrow \pi^*$ intraligand transition with some metal-to-ligand charge transfer contribution, in line with the reported similar systems.^{26–28} The smooth oxidation of neutral **1a** to $[1a]^+$ conformers leads to the appearance of new intense visible and NIR absorptions and strongly diminished parent UV absorption (Figure 4, top). The asymmetric NIR absorption of $[1a]^+$ corresponds to an overlap of two sub-bands obtained by deconvolution of the Gaussian function (Supporting Information, Figure S9 and Table S2). On further oxidation to dicationic species $[1a]^{2+}$, the characteristic NIR absorption of $[1a]^+$ gradually disappeared (Figure 4, bottom). The final, well-separated anodic step producing $[1a]^{3+}$ led to appearance of a new NIR absorption band below $1 \times 10^4\text{ cm}^{-1}$ (Figure S7) resembling the electronic absorption of oxidized reference $[1d]^+$ at $1.35 \times 10^4\text{ cm}^{-1}$ (Figure S10) and, therefore, belonging to the TPA radical cation.²⁹ It has to be considered

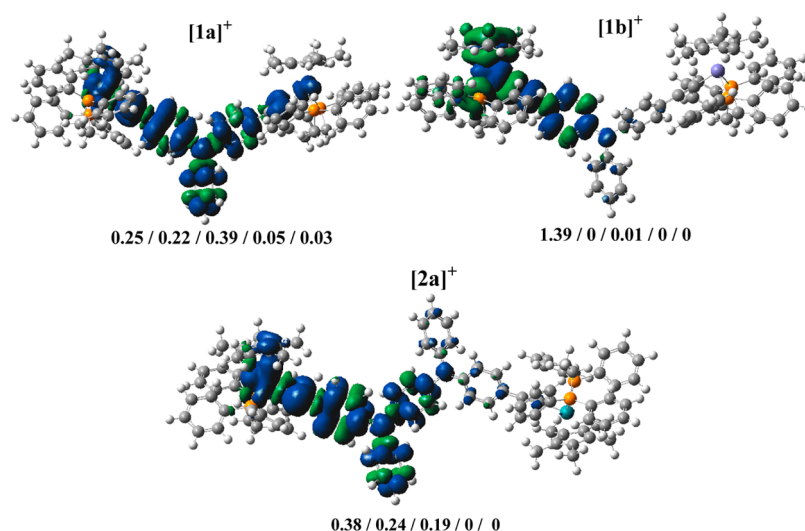


Figure 6. Calculated spin density distribution in $[1a]^+$ and $[2a]^+$ (Ru/C $\equiv$ C/arylamine/C $\equiv$ C/Ru), and $[1b]^+$ (Fe/C $\equiv$ C/arylamine/C $\equiv$ C/Fe). Contour values: ± 0.04 (e/bohr 3) $^{1/2}$. BLYP35/6-31G* (Ru/Fe: Lanl2DZ)/CPCM/CH $_2$ Cl $_2$.

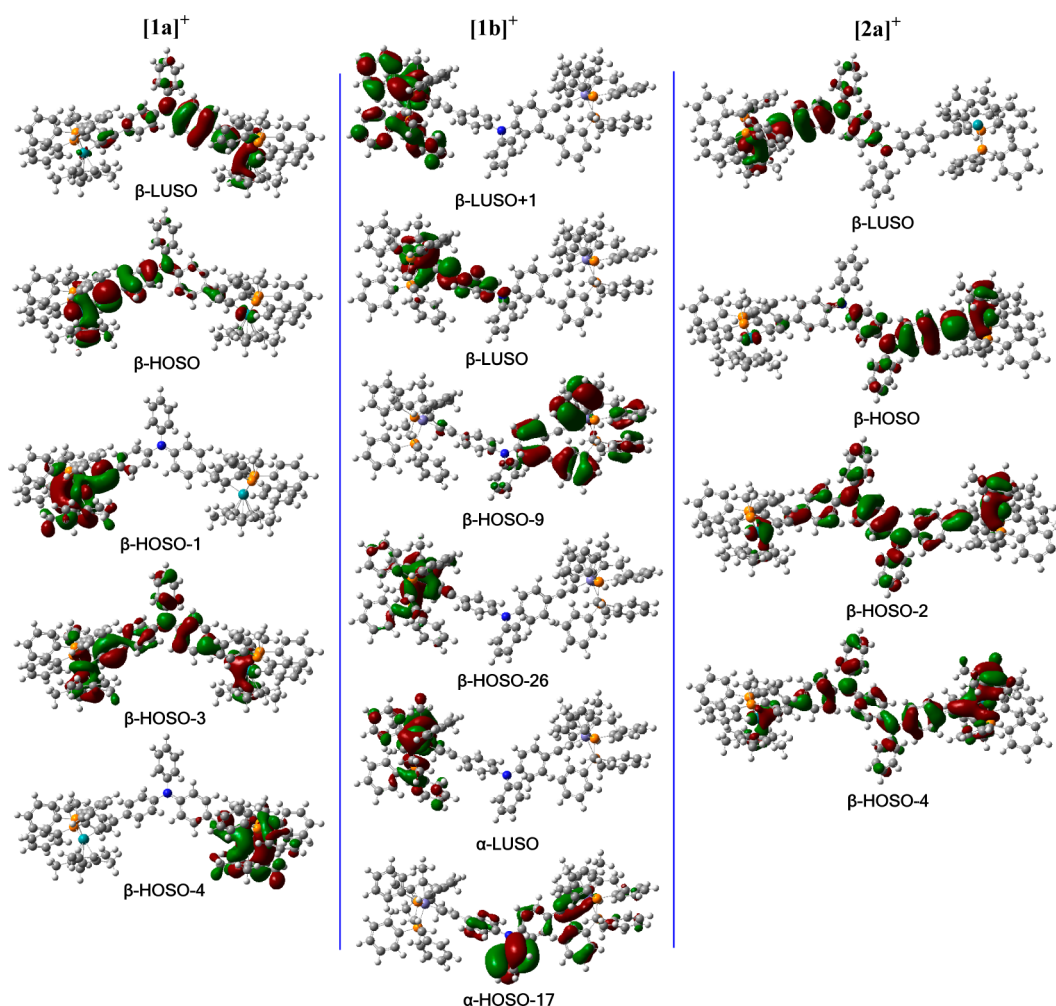


Figure 7. Spin orbitals involved in the major electronic excitations in $[1a]^+$ (left), $[1b]^+$ (middle), and $[2a]^+$ (right) presented in Table 5 (D = doublet). BLYP35/6-31G* (Ru/Fe: Lanl2DZ) /CPCM/CH $_2$ Cl $_2$.

that the oxidized Ru–C $\equiv$ C units in precursor $[1a]^{2+}$ somewhat affects the absorption of the cationic bridge-core in $[1a]^{3+}$, causing its different position and band shape compared to $[1d]^+$. Thus, the third anodic step of **1a** is mainly localized

on the TPA core, in agreement with the foregoing voltammetric and IR spectro-electrochemical results.

The stepwise oxidation of **1b** to $[1b]^{2+}$ was accompanied by the instantaneous growth of an intense symmetric absorption

Table 5. Major Electronic Excitations in [1a]⁺, [1b]⁺, and [2a]⁺ Determined by the TD-DFT Method^a

complex	excited state	λ (nm) [$\bar{\nu}$ (cm ⁻¹)]	osc strength (f)	major contributions	assignment	$\bar{\nu}$ (cm ⁻¹) (experiment)
[1a] ⁺	D ₂	1306 [7655]	0.43	β -HOSO→ β -LUSO (77%)	Ru(1)-C≡C(Ph)→Ru(2)-C≡C(Ph) ⁺ IVCT/ ILET	7900
				β -HOSO-4→ β -LUSO (23%)	Ru(2) (dppe)Cp*→Ru(2)-C≡C(Ph) ⁺ CT	
	D ₄	807 [12 390]	0.10	β -HOSO-1→ β -LUSO (60%)	Cp*Ru(1)-C≡C→Ru(2)-C≡C(Ph) ⁺ CT	10 400
				β -HOSO-3→ β -LUSO (33%)	Ru(1)-C≡C-TPA→Ru(2)-C≡C(Ph) ⁺ CT	
[1b] ⁺	D ₄	1051 [9515]	0.13	β -HOSO-26→ β -LUSO (49%)	Cp*(at Fe(2)) ⁺ →Fe(2)-C≡C(Ph) ⁺ ILET	<10 000 (unresolved $\bar{\nu}_{\max}$)
				β -HOSO-9→ β -LUSO (38%)	Cp*(dppe)C≡C→Fe(2)C≡C(Ph) ⁺ CT	
	D ₁₀	568 [17 605]	0.029	β -HOSO-26→ β -LUSO+1 (32%)	Cp*(at Fe(2)) ⁺ →dppe(at Fe(2)) ⁺	not observed
				α -HOSO-17→ α -LUSO (32%)	TPA(C≡C)→FeCp*(dppe) ⁺	
[2a] ⁺	D ₃	1091 [9165]	0.14	β -HOSO-2→ β -LUSO (75%)	Ru(1)-C≡C-TPPD→Ru(2)-C≡C(Ph) ⁺ ILET/IVCT	8400
				β -HOSO→ β -LUSO (25%)	Ru(1)-C≡C(Ph)→Ru(2)-C≡C(Ph) ⁺ IVCT/ ILET	
	D ₆	661 [15 130]	0.061	β -HOSO-4→ β -LUSO (65%)	Ru(1)-C≡C-TPPD→Ru(2)-C≡C(Ph) ⁺ ILET/IVCT	10 300
				β -HOSO-2→ β -LUSO (15%)	Ru(1)-C≡C-TPPD→Ru(2)-C≡C(Ph) ⁺ ILET/IVCT	

^aThe DFT method was BLYP35/6-31G* (Ru/Fe: Lanl2DZ) /CPCM/CH₂Cl₂. D = doublet.

band at 1.12×10^4 cm⁻¹ (Figure 5), reflecting the redox disproportionation of [1b]⁺ to the stable dication and parent 1b, in agreement with the IR monitoring of the initial anodic process (see above). The low-energy absorption of [1b]²⁺ is reminiscent of ligand-to-metal charge transfer transitions in formally Fe(III) complexes [$\{\text{Fe}(\text{dppe})\text{Cp}^*(\text{C}\equiv\text{C}-)\}_n(\text{Ph})\}^{n+}$ ($n = 1, 2$).³⁰ The striking difference in the NIR electronic absorption between [1b]²⁺ and [1a]²⁺, combined with the poorly separated anodic waves in the former case, does not support participation of the TPA bridge core in the initial oxidation of 1b. The latter process occurs probably during the subsequent well-separated oxidation of [1b]²⁺ to [1b]³⁺ (Figure S8), as indicated by the comparison with the similar electronic absorption of reference [1d]⁺ (Figure S10) as well as the invariant IR $\nu(\text{C}\equiv\text{C})$ band during this anodic step (Figure 3, right). The product of the initial one-electron oxidation step, [1b]⁺, is hardly observable during the UV-vis-NIR spectroelectrochemical monitoring of the anodic conversion of 1b to [1b]²⁺ due to the aforementioned redox disproportionation process. However, the weak NIR absorption below 1.1×10^4 cm⁻¹, attributed to unstable [1b]⁺, is clearly seen in the course of the chemical oxidation of 1b with 1 equiv of FcPF₆ (Supporting Information, Figure S6); its assignment, different from an intervalence charge transfer (IVCT), is presented in the following theoretical (time-dependent (TD) DFT) section.

The observed characteristic UV-vis-NIR spectral changes resulting from the stepwise oxidation of TMS-terminated reference compound 2d to the corresponding mono- and dication (Supporting Information, Figure S11) are fully consistent with the spectral evolution reported in the literature³¹ for bare TPPD lacking the ethynyl linkers. By contrast, the partly resolved first and second anodic steps of diruthenium diethynyl-TPPD complex 2a gave rise to UV-vis-NIR absorption changes (Supporting Information, Figure S12) strongly resembling the generation of monocationic diethynyl-TPA-bridged species [1a]⁺, which also matched

their similar IR $\nu(\text{C}\equiv\text{C})$ shifts described in the preceding text. Stable [2a]³⁺ is then characterized by a new UV-vis-NIR absorption (Figure S12, bottom) assigned to the oxidized radical-cationic TPPD core in [2a]³⁺, which is evident from the comparison with the low-lying electronic absorption of reference [2d]⁺ (Figure S11) as well as the negligible change in the $\nu(\text{C}\equiv\text{C})$ wavenumber on oxidation of [2a]²⁺ (Figure S3).

Theoretical Calculations. DFT calculations, using the BLYP35 functional, were performed to gain insight into the electronic structures of the one-electron-oxidized open-shell species [1a]⁺, [1b]⁺, and [2a]⁺, although the latter two cations were not generated in the pure forms during the spectroelectrochemical experiments due to valence disproportionation equilibria resulting in their mixture with the corresponding dications. This applies especially for [1b]⁺ detected only in a very small amount (Supporting Information, Figure S6). The basis set employed here is 6-31G* (Lanl2DZ for Ru and Fe atoms). The BLYP35 method reported by Kaupp and co-workers is appropriate for triarylamine and organometallic complexes similar to our systems.^{25,32} All the calculations were performed on nontruncated complexes [1a]⁺, [1b]⁺, and [2a]⁺ to warrant their accuracy, regardless of the relatively heavy computing burden. To account also for solvent effects, the conductor polarizable continuum model (CPCM) in CH₂Cl₂ was employed for the ground-state structural optimization and analyses as well as in TD-DFT calculations of the electronic excitation energies. The pertinent data are presented in Figures 6 and 7, Supporting Information, Figures S13 and S14, and Table 5.

The DFT results indicate similar geometric changes taking place upon the one-electron oxidation of diruthenium complexes 1a and 2a (see Supporting Information, Figure S14). The anodic electron transfer results in dominant elongation of one of the C≡C bonds and shorter adjacent (ethynyl)C-Ru, (ethynyl)C-C(phenylene), and N-C-

(phenylene) bonds. The differences between the neutral and monocationic states are greater for **1a**. The diethynyl–TPA bridge is more involved in the initial oxidation of **1a** than the diethynyl–TPPD bridge in **2a**, in agreement with the larger separation of the first two anodic waves for **1a** (Figure 2 and Table 2) reflecting a stronger electronic interaction between the Ru centers mediated by the diethynyl–TPA bridge. In contrast, no significant changes in the bonding characteristics of the diethynyl–TPA bridge accompanied the model oxidation of **1b** that is localized exclusively on the metallic termini. The iron–Cp* centers oxidize independently, and the mixed-valence species, $[\mathbf{1b}]^+$, is unstable with respect to redox disproportionation and conversion to $[\mathbf{1b}]^{2+}$, as was revealed by the UV–vis–NIR spectroscopic monitoring of the anodic path. However, the experimental IR spectro-electrochemical results (Table 3) illustrate that the ethynyl linkers are still involved in the initial oxidation of **1b**, although much less than encountered in both diruthenium complexes, **1a** and **2a**.

The important complementary results obtained with DFT for the spin density distribution in $[\mathbf{1a}]^+$, $[\mathbf{1b}]^+$, and $[\mathbf{2a}]^+$ are shown in Figure 6. Apparently, the spin localization in $[\mathbf{1a}]^+$ and, increasingly, in $[\mathbf{2a}]^+$ and $[\mathbf{1b}]^+$ is asymmetric, residing largely on the ruthenium–ethynyl(–phenylene) and iron–Cp* redox centers, respectively, in one-half of the molecule, which is fully consistent with the symmetry-broken triple bond stretching induced by the initial oxidation of **1a**, **2a**, and **1b** (vide supra). Not surprisingly, the asymmetric spin of $[\mathbf{1a}]^+$ exhibits a delocalized distribution in contrast to the strongly localized oxidation of **1b**, again affirming the effect of the different metal centers on the anodic behavior.

Also worth mentioning is the difference in the spin distribution calculated for radical cations $[\mathbf{1a}]^+$ and $[\mathbf{2a}]^+$ featuring the different bridge cores (Figure 6). In contrast to the delocalized asymmetric spin distribution in $[\mathbf{1a}]^+$, the spin density in $[\mathbf{2a}]^+$ is localized more on one of the Ru centers and less on the bridge core, with no apparent involvement of the remote ruthenium–ethynyl unit, thereby matching well the weak electronic coupling between the ruthenium centers across the elongated TPPD bridge core in $[\mathbf{2a}]^+$ illustrated by the experimental results.

TD-DFT calculations were performed to reproduce the low-energy absorption features in the experimental UV–vis–NIR spectra of the monocationic diethynyl monoamine and diamine complexes and to facilitate their assignment in support of the spin-localized bonding situation (Figure 6). The relevant electronic transitions are presented in Table 5 and depicted in Figure 7. According to the TD-DFT results, the NIR band of $[\mathbf{1a}]^+$ at 7800 cm^{-1} (ν_2 in Figure S9 and Table S2, Supporting Information) has been well-duplicated and can mainly be attributed to the β -HOSO $\rightarrow$ β -LUSO transition (HOSO = highest occupied system orbital; LUSO = lowest unoccupied system orbital). The β -HOSO is primarily localized on the nonoxidized Ru–C $\equiv$ C unit (70%), imparting the corresponding transition an appreciable IVCT character. The electronic coupling parameter, $H_{ab} = 685\text{ cm}^{-1}$, was determined from the Hush formula, $H_{ab} = (2.06 \times 10^{-2}/R_{ab})(\epsilon_{\max}\nu_{\max}\Delta\nu)^{1/2}$ (ref 33), in which R_{ab} is the distance between the ruthenium centers in the X-ray crystal structure (Figure 1). Hence, $[\mathbf{1a}]^+$ can be classified as a moderately coupled Robin-Day Class II mixed-valence compound. The higher-lying electronic transition computed at $12\,390\text{ cm}^{-1}$ is most likely responsible for the absorption band of $[\mathbf{1a}]^+$ at $10\,200\text{ cm}^{-1}$ (ν_1 in Figure S9 and Table S2, Supporting Information). It involves both β -HOSO-1

and β -HOSO-3 so that the associated charge transfer to the oxidized Ru(2)–C $\equiv$ C(Ph) $^+$ site of $[\mathbf{1a}]^+$ also involves the Cp*(Ru) and TPA donor sites.

Complex $[\mathbf{1b}]^+$ exhibits an NIR excitation at 9515 cm^{-1} belonging to β -HOSO-26 $\rightarrow$ β -LUSO and β -HOSO-9 $\rightarrow$ β -LUSO transitions. The dominant former component is localized at the oxidized metallic site, having a partial inter-configurational (IC) character typically observed for similar predominantly Fe(III) systems.^{24,34} The acceptor ethynyl(–phenylene) and donor Cp* frameworks are also involved. The second component represents a charge transfer from a ligand-based π -orbital delocalized in the nonoxidized half of $[\mathbf{1b}]^+$. A convincing experimental evidence for this low-energy electronic absorption in valence-localized $[\mathbf{1b}]^+$ with weakly interacting Fe centers is shown in Figure S6 (Supporting Information). The calculated visible electronic absorption of $[\mathbf{1b}]^+$ could not be verified experimentally (Figure 5 and Supporting Information, Figure S6) due to the redox disproportionation of the cation and the tailing absorption of $[\mathbf{1b}]^{2+}$ in that region.

The asymmetric NIR absorptions of $[\mathbf{2a}]^+$ at ca. 8400 and $10\,300\text{ cm}^{-1}$ have been reproduced less accurately than in the case of $[\mathbf{1a}]^+$, probably due to the exaggerated planar conformation of TPPD seen in the computed model. The calculated excitations encompass the dominant β -HOSO-2 $\rightarrow$ β -LUSO and β -HOSO-4 $\rightarrow$ β -LUSO transitions, respectively. Both β -HOSO-2 and β -HOSO-4 are delocalized over the planar TPPD core and the nonoxidized Ru–C $\equiv$ C unit. Similar to $[\mathbf{1a}]^+$, the β -LUSO resides on the oxidized Ru(2)–C $\equiv$ C(Ph) $^+$ site. The IVCT character of the lowest NIR absorption of $[\mathbf{2a}]^+$ becomes enhanced by the 25% admixture of the β -HOSO $\rightarrow$ β -LUSO component.

CONCLUSIONS

The electrochemical results reveal that homo-bimetallic complexes **1a**, **1b**, and **2a** undergo multistep oxidation processes. The monocationic state of **1a** is stable, with a large K_c value compared to $[\mathbf{1b}]^+$ that tends to disproportionate to the corresponding dication. Likewise, instability was also expected for $[\mathbf{2a}]^+$ with the extended diamine bridge core, based on the similar poorly resolved first two anodic responses of **1b** and **2a**. However, $[\mathbf{2a}]^+$ could still readily be detected with in situ UV–vis–NIR spectroscopy, bearing a strong resemblance to $[\mathbf{1a}]^+$. Combined results of IR and UV/vis/NIR spectro-electrochemistry and DFT/TD-DFT calculations of nontruncated models have demonstrated that (i) there exists an appreciable electronic interaction between the two ruthenium centers in $[\mathbf{1a}]^+$; (ii) the first two oxidation steps of **1b** are largely Fe-localized; (iii) the spectroscopic characteristics of $[\mathbf{2a}]^+$ indicate much weaker electronic interaction between two ruthenium centers through the extended TPPD bridge core; (iv) the subsequent oxidation of the dications of **1a**, **1b**, and **2a** is localized on the arylamine bridge core (as revealed by IR spectro-electrochemistry and the comparison with the anodic behavior of the corresponding TMS-terminated derivatives). The elongated TPPD linker, with more twisted conformation and restricted π -conjugation, indeed, exhibits electronic insulation properties to some extent. Notably, a planar conformation of TPPD in $[\mathbf{2a}]^+$ and its stronger involvement in the initial oxidation than inferred from the experimental data have emerged from DFT calculations of the model complex. The alternative linear π -conjugated oligoarylamine structures will be the subject of our following report on long-range electron transfer systems. Furthermore, this work may

expedient for designing and investigating more diversified systems with multiple redox states.

EXPERIMENTAL SECTION

General Materials. All manipulations were performed under a dry argon gas atmosphere by using standard Schlenk techniques, unless stated otherwise. Solvents were predried and distilled under argon prior to use, except those used directly for spectroscopic measurements, which were of spectroscopic grade. The starting materials 4-bromo-*N*-(4-bromophenyl)-*N*-phenylaniline (**1c**),³⁵ *N*¹,*N*⁴-bis(4-bromophenyl)-*N*¹,*N*⁴-diphenylbenzene-1,4-diamine (**2c**),³⁶ [RuCl(dppe)Cp*],³⁷ and [FeCl(dppe)Cp*]³⁸ were prepared by the procedures described in the literature. Target complexes **1a–1b** and **2a** were prepared along the synthetic route presented in Scheme 1. Other reagents were purchased and used as received.

Syntheses. Intermediate 1d. To a stirred solution of precursor **1c** (806 mg, 2 mmol), CuI (38 mg, 0.2 mmol), and [Pd(PPh₃)₄] (231 mg, 0.2 mmol) in triethylamine (20 mL) and tetrahydrofuran (THF; 30 mL) under an argon atmosphere trimethylsilylacetylene (588 mg, 6 mmol) was added, and the mixture was held at 60 °C for 24 h. After it cooled, the solution was filtered through a bed of diatomaceous earth. The filtrate was evaporated under reduced pressure and purified by silica gel column chromatography (petroleum ether) to give a light yellow solid (753 mg, yield 86%). ¹H NMR (400 MHz, CDCl₃): δ 0.25 (s, 18H, SiMe₃), 6.96 (d, *J*_{HH} = 8 Hz, 4H, Ar–H), 7.06–7.11 (m, 4H, Ar–H), 7.32–7.34 (d, *J*_{HH} = 8 Hz, 5H, Ar–H). ¹³C NMR (100 MHz, CDCl₃): δ 0.03 (SiMe₃), 93.5, 105.1 (C≡C), 116.9, 123.1, 124.1, 124.9, 125.3, 129.3, 129.5, 133.0, 146.5, 147.3 (Ar), as reported in ref 39.

Intermediate 2d. Compound **2d** was prepared from precursor **2c** by a method analogous to that employed for **1d** and purified on a silica gel column (petroleum ether/dichloromethane = 10:1, v/v) to obtain a light yellow solid (980 mg, yield 81%). ¹H NMR (400 MHz, CDCl₃): δ 0.24 (s, 18H, SiMe₃), 6.98–7.10 (m, 14H, Ar–H), 7.28–7.31 (m, 8H, Ar–H). ¹³C NMR (100 MHz, CDCl₃): δ 0.28 (SiMe₃), 93.0, 105.3 (C≡C), 115.8, 121.8, 123.4, 124.7, 125.6, 129.3, 132.8, 142.4, 146.8, 147.7 (Ar). EI-MS: *m/z* = 604.55 [M]⁺. Anal. Calcd for C₄₀H₄₀N₂Si₂: C, 79.42; H, 6.66; N, 4.63. Found: C, 79.63; H, 6.58; N, 4.67%.

Homo-Bimetallic Complexes 1a, 1b, and 2a. Target compounds **1a**, **1b**, and **2a** were prepared along the synthetic route presented in Scheme 1.

[[Ru(dppe)Cp*(C≡C)]₂(μ-TPA)] (1a). A solution of [RuCl(dppe)Cp*] (321 mg, 0.50 mmol), **1d** (100 mg, 0.23 mmol), and KF (160 mg, 2.76 mmol) in CH₃OH (20 mL) and THF (5 mL) was heated to reflux under nitrogen atmosphere for 24 h. The crude product was collected by filtration and washed with hexane. The solid was dissolved in dichloromethane and precipitated by slow diffusion of hexane. The solid was filtered off and dried to give **1a** as a yellow powder (312 mg, yield 87%). ¹H NMR (400 MHz, CDCl₃): δ 1.56 (s, 30H, CH₃ of C₅Me₅), 2.03–2.08 (m, 4H, CH₂ of dppe), 2.66–2.71 (m, 4H, CH₂ of dppe), 6.65–6.76 (m, 8H, Ar–H), 6.86–6.87 (m, 1H, Ar–H), 6.99 (d, 2H, *J* = 8 Hz Ar–H), 7.13–7.35 (m, 34H, Ar–H), 7.79 (br, 8H, Ar–H). ¹³C NMR (100 MHz, CDCl₃): δ 10.1 (CH₃ of C₅Me₅), 29.2–31.6 (m, CH₂ of dppe), 92.4 (CH₃ of C₅Me₅), 109.1 (Ru–C≡C), 122.4 (Ru–C≡C), 123.9–148.3 (m, Ar). ³¹P NMR (160 MHz, CDCl₃): δ 78.97 (s, dppe). IR (KBr/cm^{−1}): ν (C≡C) 2062 (w). Anal. Calcd for C₉₄H₉₁NP₄Ru₂: C, 72.34; H, 5.88; N, 0.90. Found: C, 72.56; H, 5.78; N, 0.91%.

[[Fe(dppe)Cp*(C≡C)]₂(μ-TPA)] (1b). A solution of **1d** (87 mg, 0.20 mmol) and K₂CO₃ (61 mg, 0.44 mmol) in CH₃OH (30 mL) and THF (10 mL) was stirred for 10 h under nitrogen atmosphere at room temperature. Then, [FeCl(dppe)Cp*] (275 mg, 0.44 mmol) and Na[BPh₄] (151 mg, 0.44 mmol) were added. After 16 h of stirring, *t*BuOK (52 mg, 0.44 mmol) was introduced, and the mixture was stirred for another 4 h, after which the solvent was evaporated and the residue was extracted with toluene (4 × 10 mL). After the solvent removal, washing with pentane (3 × 10 mL), and vacuum drying, an orange powder was obtained (182 mg, 0.12 mmol, yield 62%). ¹H

NMR (400 MHz, CDCl₃): δ 1.41 (s, 30H, CH₃ of C₅Me₅), 1.96 (br, 4H, CH₂ of dppe), 2.64 (br, 4H, CH₂ of dppe), 6.69 (br, 4H, Ar–H), 6.81–6.83 (m, 5H, Ar–H), 7.02 (d, *J* = 8 Hz, 2H, Ar–H), 7.24–7.34 (m, 34H, Ar–H), 7.91 (br, 8H, Ar–H). ¹³C NMR (100 MHz, CDCl₃): δ 10.1 (CH₃ of C₅Me₅), 30.1–31.6 (m, CH₂ of dppe), 87.5 (CH₃ of C₅Me₅), 121.0 (Fe–C≡C), 122.9–138.7 (m, Ar), 142.9 (Fe–C≡C). ³¹P NMR (160 MHz, CDCl₃): δ 95.63 (s, dppe). IR (KBr/cm^{−1}): ν (C≡C) 2051 (w). Anal. Calcd for C₉₄H₉₁NP₄Fe₂: C, 76.79; H, 6.24; N, 0.95. Found: C, 76.82; H, 6.22; N, 0.96%.

[[Ru(dppe)Cp*(C≡C)]₂(μ-TPPD)] (2a). Complex **2a** was prepared by an analogous method as **1a**, using the following amounts: [RuCl(dppe)Cp*] (464 mg, 0.69 mmol), **2d** (200 mg, 0.33 mmol), and KF (273 mg, 3.96 mmol) dissolved in CH₃OH (20 mL), THF (5 mL). The product was obtained as a light green solid (353 mg, 62% yield). ¹H NMR (400 MHz, CDCl₃): δ 1.56 (s, 30H, CH₃ of C₅Me₅), 1.99–2.10 (m, 4H, CH₂ of dppe), 2.65–2.76 (m, 4H, CH₂ of dppe), 6.70 (d, 4H, *J* = 8 Hz, Ar–H), 6.79 (d, 4H, *J* = 8 Hz, Ar–H), 6.87–6.92 (m, 6H, Ar–H), 7.03 (d, 4H, *J* = 8 Hz Ar–H), 7.17–7.34 (m, 36H, Ar–H), 7.79 (br, 8H, Ar–H). ¹³C NMR (100 MHz, CDCl₃): δ 10.0 (CH₃ of C₅Me₅), 29.4–29.7 (m, CH₂ of dppe), 92.5 (CH₃ of C₅Me₅), 122.6 (Ru–C≡C), 122.9 (Ru–C≡C), 127.1–133.8 (m, Ar). ³¹P NMR (160 MHz, CDCl₃): δ 70.86 (s, dppe). IR (KBr/cm^{−1}): ν (C≡C) 2067 (s). Anal. Calcd for C₁₀₆H₁₀₀N₃P₄Ru₂: C, 73.68; H, 5.83; N, 1.62. Found: C, 73.59; H, 5.80; N, 1.61%.

X-ray Crystallography. Single crystals of complexes **1a** and **2a** suitable for X-ray analysis were grown by layering a solution in dichloromethane with hexane. Crystals with approximate dimensions of 0.20 × 0.20 × 0.10 mm³ for **1a** and 0.20 × 0.20 × 0.20 mm³ for **2a** were mounted on glass fibers for diffraction experiments. Intensity data were collected on a Nonius Kappa CCD diffractometer with Mo Kα radiation (0.710 73 Å) at room temperature. The crystal structures were determined by a combination of direct methods (SHELXS-97)⁴⁰ and Fourier difference techniques, and refined by full matrix least-squares (SHELXL-97).⁴¹ All non-H atoms were refined anisotropically. The hydrogen atoms were placed in ideal positions and refined as riding atoms. The partial solvent molecules were omitted. Further crystal data and details of the data collection are summarized in Table S1. Selected bond distances and angles are given in Table 1.

Physical Measurements. ¹H, ¹³C, and ³¹P NMR spectra were collected on a Varian Mercury Plus 400 spectrometer (400 MHz). ¹H and ¹³C NMR chemical shifts are relative to Si(CH₃)₄, and ³¹P NMR chemical shifts are relative to 85% H₃PO₄. Elemental analyses (C, H, N) were performed with a Vario EIII Chnso instrument. The electrochemical measurements were performed on a CHI 660C potentiostat. A three-electrode single-compartment cell was used for the solution of complexes and supporting electrolyte in dry CH₂Cl₂. The solution was deaerated by argon bubbling on a frit for ~10 min before the measurement. The analyte (complex, ligand) and electrolyte (*n*-Bu₄NPF₆) concentrations were typically 1 × 10^{−3} and 1 × 10^{−1} mol dm^{−3}, respectively. A prepolished 500 μm diameter platinum disk working electrode, a platinum wire counter electrode, and an Ag wire pseudoreference electrode were used. Ferrocene was used as the internal potential reference. Spectro-electrochemical experiments at room temperature were performed with an airtight optically transparent thin-layer electrochemical (OTTLE) cell (optical path length of ca. 200 μm) equipped with a Pt minigrid working electrode and CaF₂ windows.⁴² The cell was positioned in the sample compartment of a Bruker Tensor Fourier transform IR spectrometer (1 cm^{−1} spectral resolution, eight scans) or a Shimadzu UV-3600 UV–vis–NIR spectro-photometer. The controlled-potential electrolyses were performed with a CHI 660C potentiostat. The concentration of analyte samples was ca. 2 × 10^{−3} mol dm^{−3}. Dry 3 × 10^{−1} M *n*-Bu₄NPF₆ was used as the supporting electrolyte.

Computational Details. DFT calculations were performed with the Gaussian 09 program,⁴³ at the BLYP35/6-31G* level of theory. The basis set employed was 6-31G* (Lanl2DZ for Ru and Fe atoms). Geometry optimization was performed without any symmetry constraints. Electronic transitions were calculated by the TD-DFT method. The molecular orbital contributions were generated using the Multiwfn package and plotted using GaussView 5.0. The solution

651 effects in dichloromethane are included for a part of the calculations
652 with the CPCM.⁴⁵

653 ■ ASSOCIATED CONTENT

654 ● Supporting Information

655 The Supporting Information is available free of charge on the
656 ACS Publications website at DOI: 10.1021/acs.inorg-
657 chem.6b02809.

658 Crystallographic information, additional spectro-electro-
659 chemical information, additional calculated DFT data,
660 ¹H, ¹³C and ³¹P NMR spectra of the new compounds.
661 (PDF)

662 Crystallographic data (CIF)

663 Crystallographic data (CIF)

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670 Notes

671 The authors declare no competing financial interest.

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