Sandwich and half-sandwich metal complexes derived from

cross-conjugated 3-methylene-penta-1,4-diynes

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Abstract

The cross-conjugated ethynyl-vinylidene [Ph₂C=C(≡CH){C(H)=CRu(PPh₃)₂Cp}]PF₆ ([4a]PF₆), and [FcC(H)=C(≡CH){C(H)=CRu(PPh₃)₂Cp}]PF₆ ([4b]PF₆), and ethynyl-alkynyl Ph₂C=C(≡CH){C≡CRu(PPh₃)₂Cp} (5a), and FcC(H)=C(≡CH){C≡CRu(PPh₃)₂Cp} (5b) compounds (Cp = η⁵-cyclopentadienyl) have been prepared from reactions of the known 3-methylene-penta-1,4-diynes Ph₂C=C(≡CH)₂ (3a) and [FcCH=C(≡CH)₂] (3b) with [RuCl(PPh₃)₂Cp]. The compounds derived from 3b incorporating the more electron-rich alkene proved to be unstable during work-up, and attempts to prepare bis(ruthenium) complexes from 3a and 3b or from transmetallation reactions of the bis(alkynylgold) complex FcCH=C(≡CAuPPh₃)₂ (7) with RuCl(PPh₃)₂Cp were unsuccessful. The related bis- and tris(ferrocenyl) derivatives Ph₂C=C(C≡CFc)₂ (6a) and FcCH=C(C≡CFc)₂ (6b) were more readily obtained from Pd(II) / Cu(I) catalysed cross-coupling reactions of FcC≡CH with the 1,1-dibromo vinyl complexes PhC=CBr₂ (1a) and FcC(H)=CBr₂ (1b). Cyclic voltammetry of 6a and 6b using n-Bu₄N[PF₆] as the supporting electrolyte shows broad, overlapping waves arising from the sequential oxidation of the ferrocenyl moieties in electronically and chemically similar environments. Electrostatic effects between the ferrocenyl moieties are enhanced in solutions of the weakly ion-pairing electrolyte n-Bu₄N[B{C₆H₃(CF₃)₂-3,5}₄], leading to better resolution of the individual electrochemical processes. The comparative IR spectroelectrochemical response of 6a and 6b suggest the vinyl ferrocene moiety in 6b undergoes oxidation before the ethynyl ferrocene fragments. There is no evidence of electronic coupling between the metalloocene moieties and [6a]⁺, [6b]ⁿ⁺ (n = 1, 2) are best described as Class I mixed-valence compounds.

Introduction

Ferrocene has long been used as an electrophore in the design and study of mixed-valence (MV) compounds.¹ In the 1970s Cowan demonstrated that biferrocenyl was able to undergo one-electron oxidation to give a Class II MV compound with a characteristic inter-valence charge transfer (IVCT) band centred at ca. 1900 nm (5260 cm⁻¹).²⁻⁶ Studies of the redox properties of related
1,2-bis(ferrocenyl)ethenes also revealed thermodynamically stable MV compounds to be formed on one-electron oxidation. Although spectroscopic studies of the Z-isomers were complicated by isomerisation, available data support the description of these compounds as less strongly coupled Class II systems. Studies of MV complexes derived from 1,4-bis(ferrocenyl)buta-1,3-dienes such as FcC(H)=C(H)C(H)=C(H)Fc are also complicated by a range of issues relating to facile interconversion between the cis and trans isomers. Related structures in which the diene fragment is (conceptually) constrained into a heterocyclic structure have proven to be more amenable to study, and a combination of spectroscopic and computational studies have indicated that these heterocyclic bridged MV compounds can be considered as Class II MV complexes, with the diene-like backbone contributing significantly to the optical charge transfer transition.

The thermodynamic stability of MV derivatives of the structurally simple, linear α,ω-bis(ferrocenyl) oligoyne Fc(C≡C)ₙFc decreases as the oligoyne chain length increases, such that the ferrocene redox processes in 1,6-bis(ferrocenyl)hexa-1,3,5-triyne are heavily overlapped. Although the 1,8-bis(ferrocenyl)octa-1,3,5,7-tetrayne has been prepared, we are unaware of any studies of mixed-valence analogues of this compound; in the case of the 1,12-bis(ferrocenyl)dodeca-1,3,5,7,9,11-hexayne the ferrocene moieties give rise to two genuinely independent oxidation processes in a 0.1 M n-Bu₄NPF₆ (1:1 CH₂Cl₂/NCMe) electrolyte. Although spectroscopic data from these longer MV complexes are limited, those that are available, indicate that the interactions between the ferrocenyl moieties decrease with increasing the bridge length.

In contrast, oligoenediyl- and oligoynediyl-bridged complexes of electron-rich ruthenium phosphine moieties generally give rise to much more extensively delocalised radical cations on oxidation, as in [{Ru(PPh₃)₂(CO)Cl}₂(µ-CH=CHCH=CH)], [{Ru(PEt₃)₃(CO)Cl}₂(µ-CH=CHCH=CH)], and [{Ru(PPh₃)₂(CO)Cl(NC₅H₄COOEt-4)}₂(µ-CH=CHCH=CH)] and the various spectroscopically identified rotamers of {Cp’(L₂)Ru}₂(µ-C≡CC≡C) (Ru(L₂)Cp’ = Ru(PPh₃)₂Cp).
Ru(PMe$_3$)(PPh$_3$)Cp,$^{20}$ Ru(dppe)Cp,$^{21}$ Ru(dppe)Cp* $^{21}$). However, the substantial contributions from the unsaturated bridging ligand to the semi-occupied molecular orbitals (SOMOs) suggests that such systems are better described in terms of bridge-localised radical cations than true mixed-valence complexes.$^{22}$

In addition to linear systems, a variety of compounds featuring more than two ferrocene or ruthenium moieties linked through a common but branched bridging ligand have been explored. Interest in such multiple electrophore complexes include the fundamental mechanisms and methods of analysis of charge transfer through ‘2-dimensional’ frameworks,$^{25}$ and assessment of the optoelectronic properties of branched molecules.$^{26-28}$ Exemplary molecular systems include 1,3,5-tris(ferrocenylythynyl)benzene,$^{29}$ and Astruc's 1,2,3,4,5,6-hexakis(ferrocenylythynyl)benzenes,$^{30,31}$ which exhibit electrostatic interactions between the ferrocenyl moieties upon oxidation, if not true electronic delocalisation. The use of electrolytes with weakly ion-pairing anions such as n-Bu$_4$N[B{C$_6$H$_3$(CF$_3$)$_2$-3,5}$]$_4$] enhances through-space (or Coulombic) interactions between nearby electrophores, leading to the resolution of individual electrochemical events.$^{32}$ For further example, although there is no significant electronic coupling between the ferrocene moieties in tetraferrocenyl(nickel dithiolene), in n-Bu$_4$N[B{C$_6$H$_3$(CF$_3$)$_2$-3,5}$]$_4$] four distinct oxidation events (and two dithiolene based reductions) are observed by differential pulse voltammetry.$^{33}$

Cross-conjugated $\pi$-systems are potential scaffolds upon which to further develop such interests,$^{34-36}$ and attention is now being directed towards the synthesis and properties of complexes featuring cross-conjugated bridging ligands such as ene-yynes.$^{37-39}$ We have recently described a range of mono(ferrocenyl) derivatives of 3-methylene-penta-1,4-diyne, which were obtained from cross-coupling reactions of alkynes with 1,1-dibromo-2-ferrocenyl ethane,$^{40}$ and also the step-wise assembly of 1,1,2,2-tetrakis(ferrocenylythynyl)ethane.$^{41}$ The latter revealed well-resolved sequence of four ferrocene-based oxidations in n-Bu$_4$N[B{C$_6$H$_3$(CF$_3$)$_2$-3,5}$]$_4$], although there
was no clear spectroscopic evidence for any significant coupling between the ferrocene / ferrocenium moieties. Now attention is turned to related compounds featuring both ruthenium half-sandwich and ferrocene electrophores. In this contribution further synthetic explorations of metal complexes of 3-methylene-penta-1,4-diyne are described, together with the electrochemical characteristics of the tris(ferrocenyl) complex FcC(H)=C(C≡Cfc)₂.

### Results & Discussion

The half-sandwich bis(phosphine) ruthenium complexes, exemplified by [RuCl(PPh₃)₂Cp], undergo facile reaction with terminal alkynes, HC≡CR, to give vinylidene complexes (e.g. [Ru(C≡C(H)R)(PPh₃)₂Cp]⁺) which in turn are readily deprotonated to the acetylide complexes (e.g. [Ru(C≡CR)(PPh₃)₂Cp]).

Complexes of this type continue to serve as workhorses for the exploration of vinylidene and acetylide ligand chemistry since the inception of the field in the 1980s. However, attempts to access the chemistry of ruthenium complexes featuring cross-conjugated ligand systems by reaction of [RuCl(dppe)Cp*] with penta-3-one-1,4-diyne (HC≡CC(=O)C≡CH) were unsuccessful; instead, reaction of [Ru(C≡CH)(dppe)Cp*] with oxalyl chloride gave the bimetallic complex [{Ru(dppe)Cp*}₂{µ-C≡C(CO)C≡C}] in good yield, providing a platform for further synthetic transformations of the cross-conjugated bridging ligand. However, a more direct route to such complexes is still to be developed.

Benzophenone, PhC(=O)Ph, is conveniently transformed to 1,1-dibromo-2,2-diphenyl ethene 1a (79%) on reaction with CBr₄ and PPh₃. The dibromoethene 1a is in turn smoothly cross-coupled ([Pd(PPh₃)₄] / Cul catalyst) with trimethylsilylacetylene, Me₃SiC≡CH, to give the bis(alkyne) Ph₂C=C(C≡CSiMe₃)₂ (2a, 84%), and desilylated (K₂CO₃ / MeOH) to give Ph₂C=C(C≡CH)₂ (3a, 95%) (Scheme 1). Reactions of 2a and 3a with two equivalents of [RuCl(PPh₃)₂Cp] were explored in a range of solvents (THF, MeOH, CH₂Cl₂) and conditions (2a: KF, n-Bu₄NF; 3a: NH₄PF₆, n-Bu₄NPF₆, KPF₆; room temperature – reflux; 1 – 24 h). In all cases, in situ NMR monitoring revealed the formation of the monovinylidene [4a]PF₆ and unreacted [RuCl(PPh₃)₂Cp]. The synthesis of [4a]PF₆ was subsequently optimised from the 0.9:1 reaction of [RuCl(PPh₃)₂Cp]
with 3a (CH₂Cl₂ / KPF₆ / reflux / 20 hrs) and isolated in 79% yield by precipitation with diethyl ether (Scheme 1). The ³¹P NMR spectrum of [4a]PF₆ is characterised by a singlet at δ 41.62 ppm, whilst in the ¹H NMR spectrum the vinylidene ligand is evinced by a triplet at δ 4.92 ppm (Jₕₜ = 2.5 Hz) and the free ethynyl moiety by a singlet at δ 3.09 ppm. The Cp ligand gave a low field resonance typical of the {Ru(PPh₃)₂Cp}⁺ fragment at δ 5.23 ppm. The IR spectrum contained the anticipated vinylidene ν(C=C) (1629 cm⁻¹), alkyne ν(C≡C) (2108 cm⁻¹) and ethene (1483 cm⁻¹) bands. Other spectroscopic and analytical data were in accord with the proposed structure, which was confirmed by single crystal X-ray analysis (vide infra).

Scheme 1 The synthesis of 1a – 5a.

Treatment of a suspension of [4a]PF₆ in methanol with KO'Bu gave the ethynyl-alkynyl complex 5a as a bright yellow precipitate (62%) (Scheme 1). In the ³¹P NMR spectrum, the half-sandwich fragment gave rise to a singlet at δ 50.35 ppm, whilst the Cp moiety was detected as a singlet in the ¹H NMR spectrum at δ 4.22 ppm, together with a singlet from the free ethynyl proton at δ 2.77 ppm. The IR spectrum contained ν(C≡C) bands from both free (2105 cm⁻¹) and coordinated (2044 cm⁻¹) alkyne moieties in addition to the ethene ν(C=C) band at 1482 cm⁻¹.
Again, other spectroscopic and analytical data, and a single crystal X-ray structure determination, confirmed the structure. Further attempts to metallate the free ethynyl fragment in 5a were also unsuccessful.

Table 1. Selected bond lengths (Å) and angles (°) from the crystallographically determined structures of [4a]PF₆, and 5a.

<table>
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<tr>
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<th>[4a]PF₆</th>
<th>5a</th>
<th>[4a]PF₆</th>
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<td>Ru1–C1</td>
<td>1.842(3)</td>
<td>2.0017(15)</td>
<td>Ru1–C1–C2</td>
<td>172.2(2)</td>
</tr>
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<td>1.328(4)</td>
<td>1.2202(2)</td>
<td>C1–C2–C3</td>
<td>125.6(2)</td>
</tr>
<tr>
<td>C2–C3</td>
<td>1.475(3)</td>
<td>1.430(2)</td>
<td>C2–C3–C4</td>
<td>115.1(2)</td>
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<tr>
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<td>2.2831(4)</td>
<td>C2–C3–C6</td>
<td>132.2(2)</td>
</tr>
<tr>
<td>Ru1–P2</td>
<td>2.3705(7)</td>
<td>2.2923(4)</td>
<td>P1–Ru1–C1</td>
<td>87.49(8)</td>
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<tr>
<td>C3–C4</td>
<td>1.435(4)</td>
<td>1.444(2)</td>
<td>P2–Ru1–C1</td>
<td>98.39(8)</td>
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<tr>
<td>C4–C5</td>
<td>1.190(4)</td>
<td>1.189(2)</td>
<td>C3–C6</td>
<td>1.371(4)</td>
</tr>
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</table>

The molecular structures of [4a]PF₆ and 5a are plotted in Figures 1 and 2, and illustrate the rather congested environment of the ostensibly reactive alkyne moiety. Selected bond lengths and angles summarised in Table 1. The vinylidene cation [4a]+ (Figure 1) features the expected short Ru=C [1.842(3) Å] and C=CH [1.328(4) Å] distances, and linear Ru=C=C fragment [172.2(2)°], consistent with related examples described elsewhere.⁴⁸⁻⁵² The vinylidene ligand deviates slightly from co-planarity with the ene-yne substituent (C(1)-C(2)-C(3)-C(4) -26.3(3)°); the orientation of vinylidene ligands relative to the half-sandwich fragment has been discussed on previous occasions.⁵³,⁵⁴ Although relatively free rotation of the vinylidene complex around the C(2)-C(3) bond might be anticipated given the significant single-bond character (C(2)-C(3) 1.475(3) Å), the larger steric effects of the C(81) phenyl ring compared with the C(4)=C(5)H ethynyl fragment leads to the half-sandwich group adopting a position s-cis with respect to C(4)=C(5). It is likely that the further formation of both the initial π-complex and subsequent rearrangement to the bis(vinylidene) expected on reaction of [4a]+ with RuCl(PPh₃)₂Cp are disfavoured on steric grounds.
Figure 1 A plot of the cation [4a]$^+$ from the crystallographically determined structure of [4a]PF$_6$. Hydrogen atoms have been removed for clarity. Ellipsoids are plotted at 50%.

Similar arguments can be invoked from the acetylide complex 5a (Figure 2). The formal bond orders within the cross-conjugated ligand are reflected in the C(1)=C(2) (1.2202(2) Å), C(2)-C(3) (1.430(2) Å), C(3)=C(6) (1.375(2) Å), C(3)-C(4) (1.444(2) Å) and C(4)≡C(5) (1.189(2) Å) bond lengths. It seems probable that the combination of the flanking phenyl groups at C(6) and the metal alkynyl fragment again hinder access of a second {Ru(PPh$_3$)$_2$Cp}$^+$ to the C(4)=C(5) triple bond and therefore prevent further reaction.
Given the ready access to 1,1-diethynyl-2,2-diphenyl-ethenes from 1a an alternative approach to cross-conjugated bi-metallic derivatives was explored. Reaction of 1a with ethynyl ferrocene (FcC≡CH) under the usual cross-coupling conditions gave the bis(ferrocenyl) compound 6a in moderate (48%) yield as a bright orange solid (Scheme 2).

Scheme 2. The preparation of 6a.

The conversion of ferrocene carboxaldehyde (FcCHO) to 1,1-dibromo-2-ferrocenylethene (1b) is known; most usually 1b serves as a precursor to both ethynyl ferrocene (FcC≡CH), and 1,1-diethynyl-2-ferrocenyl ethenes. The
Pd(0) / Cu(I) catalysed cross-coupling reactions of 1b with trimethylsilylacetylene gives 2b (76%), which in turn is readily desilylated to give 3b (94%) (Scheme 3). In a manner similar to that described for 6a, the cross-coupling of 1b with ethynyl ferrocene gave the tris(ferrocenyl) complex 6b (68%) (Scheme 3).

Scheme 3. The preparation of 2b, 3b and 6b

In a manner similar to that described above for 3a, reactions of 3b with one equivalent of [RuCl(PPh₃)₂Cp] and NH₄PF₆ gave the mono-vinyldene complex [Ru{C=C(H)C(=CHFc)(C≡CH)}(PPh₃)₂Cp]PF₆, but which proved to be unstable on deprotonation with KOBu'. All attempts at reaction of 3b with excess [RuCl(PPh₃)₂Cp] proved unsuccessful, yielding brightly coloured blue/green compounds that could not be satisfactorily characterised.

Metallation of 2b with [AuCl(PPh₃)] took place smoothly in methanol solutions of NaOH to give [FcC(H)=C(C≡C̷AuPPh₃)] (7, 71%), which was characterised by the usual combination of spectroscopic and elemental analytical methods (Scheme 4). The ³¹P NMR spectrum gave resonances at δ 42.20 and 42.41 ppm corresponding to the inequivalent phosphine ligands, whilst Au-C resonances
were observed in the $^{13}$C NMR spectrum as doublets ($J_{CP} = 27$ Hz in each case) at 105.53 and 102.91 ppm. The $\nu$(AuC≡C) (2102 cm$^{-1}$) and $\nu$(FcC=CR$_2$) (1481 cm$^{-1}$) bands in the IR spectrum were also characteristic of 7. Unfortunately, despite the success of transmetallation reactions between gold acetylide complexes and half-sandwich iron and ruthenium complexes,$^{56}$ reactions of 7 with [RuCl(PPh$_3$)$_2$Cp] or [RuCl(dppe)Cp*] failed to yield the desired acetylide complexes.

Scheme 4. The preparation of 7.

Single crystals of the tris(ferrocenyl) complex 6b suitable for X-ray diffraction were obtained by slow evaporation of a dichloromethane solution (Figure 3). Pertinent bond lengths and angles are given in Table 2. As expected, the 1,1-dialkynyl ethene portion of the molecule is essentially planar, with key C=C (1.366(5) Å), and C≡C (1.193(5) and 1.198(5) Å) bond lengths consistent with the formal valence bond description. There is no evidence from the bond lengths for any substantive difference in conjugation between the ferrocene moieties along the cis- or trans- ene-yne pathways.
Figure 3. A plot of the molecular structure of 6b, selected hydrogen atoms removed, showing the atom labeling scheme (thermal ellipsoids plotted at 50 %).

Table 2. Bond lengths (Å) and angles (deg) from the crystal structure of 6b.

<table>
<thead>
<tr>
<th>Bond Length or Angle</th>
<th>Value</th>
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<td>C11–C1–C2</td>
<td>128.2(3) Å</td>
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<tr>
<td>C1–C11</td>
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</tr>
<tr>
<td>C1–C2–C3</td>
<td>120.0(3) deg</td>
</tr>
<tr>
<td>C1–C15</td>
<td>1.435(5) Å</td>
</tr>
<tr>
<td>C1–C2–C5</td>
<td>124.1(3) deg</td>
</tr>
<tr>
<td>C1–C2</td>
<td>1.366(5) Å</td>
</tr>
<tr>
<td>C2–C3–C4</td>
<td>178.5(4) deg</td>
</tr>
<tr>
<td>C2–C3</td>
<td>1.437(5) Å</td>
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<tr>
<td>C2–C5–C6</td>
<td>176.2(4) deg</td>
</tr>
<tr>
<td>C2–C5</td>
<td>1.193(5) Å</td>
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<tr>
<td>C3–C4–C21</td>
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<tr>
<td>C3–C4</td>
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<td>C6–C31</td>
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</table>

The redox chemistry of 6a and 6b was explored by cyclic voltammetry in a dichloromethane solution containing 10⁻¹ M n-Bu₄NX as the supporting electrolyte (X = [PF₆] or [B(C₆H₅)(CF₃)₂-3,5]₄ ([BARf₄])) (Table 3, Figure 4). When the more strongly ion-pairing or associating electrolyte is employed (X = [PF₆]), the voltammogram of 6a exhibits a single, chemically reversible electrochemical process. The anodic-to-cathodic peak separation (ΔEₚ(1)) for this wave approaches twice that of the internal ferrocene/ferrocenium reference couple.
consistent with the overlapped one-electron processes of the two ethynyl ferrocenyl moieties. In the same electrolyte, the tris(ferrocenyl) complex 6b shows two close lying waves, with apparent peak currents approximately in the ratio 1:2 suggesting that the vinyl-ferrocene fragment undergoes oxidation at a measurably different potential than the two (electrochemically indistinguishable) ethynyl-ferrocene moieties (Table 3, Figure 4). The reversible, apparently one electron, oxidation can be further assigned on the basis of comparison with the potentials observed for oxidation of FcCH=C(C≡CR)₂ complexes structurally related to the vinyl-ferrocene moiety. The second redox wave is assigned to the overlapping oxidation processes of the two ethynylferrocene groups, with the ΔEₚ of this wave approaching twice that of the internal reference and a halfwave potential consistent with that observed for the ethynyl ferrocene moieties in complex 6a (Table 3).

Cyclic voltammograms obtained from solutions of an electrolyte containing the weakly coordinating anion [BARF₄]⁻ exhibit greater separation of the individual redox processes of 6a and 6b. This observation is consistent with results from other poly(ferrocene) compounds in electrolytes containing weakly coordinating anions, with an increase in separation of the individual redox events to give two (6a) or three (6b) reversible, one electron process that can be attributed sequential oxidation of the two ethynylferrocene fragments (6a) or the vinyl and two ethynylferrocenyl fragments (6b). The greater separation of the redox processes separation in the voltammetric experiments conducted with the weakly coordinating [BARF₄]⁻ anion, and hence larger comproportionation constant Kc (Table 3), suggested that there would be a better prospect of observing the individual charge states of each complex in spectroelectrochemical experiments when the appropriate electrolyte was employed.
Figure 4. Cyclic voltammetry of 6a (upper) 6b (lower) in CH$_2$Cl$_2$ solution containing $10^{-1}$ M $n$-Bu$_4$NX as the supporting electrolyte ($X = $ [PF$_6$]) (dotted lines) or [B$_3$H$_3$(CF$_3$)$_2$-3,5]$_4$ (solid lines),
Table 3. Electrochemical data from 6a and 6b in a dichloromethane solution containing 10^{-1} M n-BuNX as the supporting electrolyte (X = [PF_6] or [B(C_6H_3(CF_3)_{2-3,5})_4])

<table>
<thead>
<tr>
<th></th>
<th>6a</th>
<th>6b</th>
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<tbody>
<tr>
<td>X</td>
<td>[PF_6]</td>
<td>[BARf_4]^-</td>
</tr>
<tr>
<td>E_{1/2}(1)/V</td>
<td>0.202</td>
<td>0.089</td>
</tr>
<tr>
<td>E_{1/2}(2)/V</td>
<td>0.201</td>
<td>0.202^a</td>
</tr>
<tr>
<td>E_{1/2}(3)/V</td>
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<tr>
<td>ΔE(1-2)/V</td>
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^a from differential pulse voltammetry by the method of Taube and Richardson.57

Spectroelectrochemical investigations were undertaken in order to better assess the interactions between the ferrocenyl moieties in 6a and 6b. Spectroelectrochemical measurements were carried out in CH_2Cl_2 solutions containing 10^{-1} M n-BuN[B(C_6H_3(CF_3)_{2-3,5})_4] as supporting electrolyte to maximize the thermodynamic stability of the individual redox states with respect to redox disproportionation.

The IR spectra of the bis(ferrocenyl) compound [6a]^{n+} (n = 0–2) give rise to a ν(C≡C) band envelope which steadily increases in intensity over the course of the two oxidation steps (Figure 5). The asymmetrically-shaped band envelopes of [6a]^{n+} display unresolved shoulders indicative of multiple, overlapping ν(C≡C) bands. The ν(C=C) band at 1481 cm^{-1} is unaffected by the change in the oxidation state, while a new electronic absorption band grows at ca. 4150 cm^{-1} consistent with the localized d-d transition of ferrocenium.41,58
Figure 5. Reversible IR spectral changes in the ν(C≡C) region accompanying oxidation of the ferrocenyl moieties of [6a]n+ in CH2Cl2 / 10⁻¹ M n-Bu4N [B{C₆H₃(CF₃)₂-3,5}]₄ within an OTTLE cell.

The IR spectra of complex [6b]n⁺ (n = 0–3, Figure 6) exhibit very weak ν(C≡C) bands in the neutral (n = 0) and monocationic (n = 1) species. As oxidation proceeds, the band envelope becomes more intense, although the appearance of this broad, structured band envelope differs little between the di and tri (n = 2, 3) cationic states (Figure 6, Table 4). Again, the vinyl ν(C=C) (1552 cm⁻¹) is unchanged as a function of the redox state of the molecule and ferrocenium based electronic transitions are observed at ca. 4150 cm⁻¹. A comparison of the trends in intensity of the ν(C≡C) bands as a function of oxidation state change in 6a and 6b is consistent with the initial oxidation of the vinyl ferrocene moiety drawn from the electrochemical data above. Further examination of the NIR region failed to identify any electronic transition that could be uniquely associated with the mixed-valence states [6a]+, [6b]+ and [6b]2+, and therefore these systems are described as Class I systems in the framework of the Robin-Day classification scheme. This behaviour parallels that of other multi-ferrocene complexes such as the tetraferrocenyl(nickel dithiolene) complex studied in the Geiger group.33
Figure 6. Reversible IR spectral changes in the ν(C≡C) region accompanying oxidation of the ferrocenyl moieties of \([6b]^{n^+}\) in CH\(_2\)Cl\(_2\) / 10\(^{-1}\) M \(n\)-Bu\(_4\)N \([B\{C_6H_3(CF_3)_2-3,5\}_4]\) within an OTTLE cell.

Table 4. IR ν(C≡C) / cm\(^{-1}\) data for \([6a]^{n^+}\) and \([6b]^{n^+}\) obtained spectroelectrochemically in CH\(_2\)Cl\(_2\) / 10\(^{-1}\) M \(n\)-Bu\(_4\)N\([B\{C_6H_3(CF_3)_2-3,5\}_4]\) within an OTTLE cell.

<table>
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Conclusion

The monometallic ethynyl-vinylidene complex \([\text{Ph}_2\text{C}=\text{C}=(\text{CH})=\text{CRu}(\text{PPh}_3)_2\text{Cp}]\)\([\text{PF}_6\) ([4a]PF₆) and the related ethynyl-alkynyl complex \(\text{Ph}_2\text{C}=\text{C}=(\text{CH})=\text{CRu}(\text{PPh}_3)_2\text{Cp}\) \([5a]\) were successfully prepared from \(\text{Ph}_2\text{C}=\text{C}=(\text{CH})\) \([3a]\) and characterised. However, analogous compounds derived from the more electron-rich ligand precursor \([\text{FcCH}=\text{C}=(\text{CH})_2]\) \([3b]\) proved to be unstable during work-up. Furthermore, attempts to prepare bis(ruthenium) complexes from \(3a\) and \(3b\) or from transmetallation reactions of the bis(alkynylgold) complex \(\text{FcCH}=\text{C}=(\text{CH})_2\text{AuPPh}_3\) \([7]\) with \(\text{RuCl}(\text{PPh}_3)_2\text{Cp}\) were unsuccessful. Instead, bis- and tris(ferrocenyl) compounds \(\text{Ph}_2\text{C}=\text{C}(\text{C}=\text{CFc})_2\) \([6a]\) and \(\text{FcCH}=\text{C}(\text{C}=\text{CFc})_2\) \([6b]\) were more readily obtained from Sonogashira-like Pd(II)/Cu(I) catalysed cross-coupling reactions of \(\text{FcCH}=\text{C}=(\text{CH})\) with the 1,1-dibromo vinyl complexes \(\text{PhC}=\text{C}=(\text{Br})_2\) \([1a]\) and \(\text{FcC}(\text{H})=\text{C}=(\text{Br})_2\) \([1b]\). Analysis of the multi-ferrocene compounds \(6a\) and \(6b\) by cyclic voltammetry in \(\text{CH}_2\text{Cl}_2\) solutions containing \(n\)-\(\text{Bu}_4\text{NPF}_6\) or \(n\)-\(\text{Bu}_4\text{NBAr}_4\) as supporting electrolyte indicates a significant contribution from electrostatic effects to the separation of the individual ferrocene-based redox processes, and hence better resolution of the individual electrochemical processes in the electrolyte featuring the more weakly coordinating anion, \([\text{BAr}_4\text{F}_4]^-\). The trends in the electrochemical potentials and the IR spectroelectrochemical response of \(6a\) and \(6b\) indicate the vinyl ferrocene moiety in \(6b\) undergoes oxidation before the ethynyl ferrocene fragments. There is no evidence of electronic coupling between the metallocene moieties and \([6a]^{\pm}, [6b]^{\pm}\) \((n = 1, 2)\) are best described as Class I mixed-valence compounds. Efforts to reduce the steric bulk of the half-sandwich metal fragment (e.g. \(\text{Ru(dppe)}\text{Cp}, \text{Ru(PMe}_3)_2\text{Cp}\)) or alter the orientation of the intermediate vinylidenes (e.g. \(\text{Mo(dppe)}(\text{η}_1\text{C}_2\text{H}_7)\)) to promote further metallation of the cross-conjugated ligand will form the basis of future work.

Experimental Section

**General Conditions** All reactions were carried out under an atmosphere of dry nitrogen using standard Schlenk techniques. Reaction solvents were dried and
distilled or purified by passage through an Innovative Technologies SPS-400 solvent purification system, and degassed before use. Other solvents were standard reagent grade and used as received. No special precautions were taken to exclude air or moisture during workup except where otherwise indicated. The compounds [RuCl(PPh₃)₂Cp]₅⁹ 1,1-dibromo-2,2-diphenyl ethene (1a)⁴⁷ 1,1-dibromo-2-ferrocenyl ethene (1b)⁴⁰ 1,1-bis(trimethylsilylethynyl)-2,2-diphenyl-ethene (2a)⁶⁰ 1,1-bis(trimethylsilylethynyl)-2-ferrocenyl-ethene (2b)⁴⁰ 1,1-bis(ethynyl)-2,2-diphenyl-ethene (3a)⁶⁰ and [1,1-bis(ethynyl)-2-ferrocenyl-ethene (3b)]⁴⁰ were prepared by literature routes. Ethynyl ferrocene (FcC≡CH) was synthesised from 1b,⁵⁵ [AuCl(PPh₃)] from HAuCl₄,⁶¹ and [Pd(PPh₃)₄] from PdCl₂.⁶² Carbon tetrabromide (CBr₄) was sublimed before use. All other reagents were commercially available and used as received. NMR spectra were recorded at 23 °C on a Varian NMR Systems 700 spectrometer using CD₂Cl₂ as the solvent. Chemical shifts were determined relative to internal solvent signals,⁶³,⁶⁴ or external 85% H₃PO₄ (δ = 0.00 ppm). Assignment of the ¹H and ¹³C NMR data was supported by gradient selected ¹³C, ¹H HMQC and HMBC experiments. ASAP-MS spectra were recorded from solid aliquots on an LCT Premier XE mass spectrometer (Waters Ltd., U.K.) or Xevo QToF mass spectrometer (Waters Ltd., U.K.) in which the aliquot is vapourised using hot N₂, ionized by a corona discharge, and carried to the TOF detector (working range m/z 100−1000). A melting point tube was dipped in to a sample solution (~1 mg/mL) and introduced into the spectrometer, where a temperature ramp from 50 °C to 450 °C vaporises the sample enabling atmospheric pressure chemical ionisation (APCI) to occur. MS data is processed using MassLynx 4.1.

Cyclic voltammetry was carried out using an EcoChemie Autolab PG-STAT 30 or a Palm Instruments EmStat2 potentiostat, with a platinum disc working electrode, a platinum wire counter electrode, and a platinum wire pseudo-reference electrode, from solutions in dichloromethane containing either 0.1 M n-Bu₄NPF₆ or 0.1 M n-Bu₄N[B{C₆H₃(CF₃)₂-3,5}]₄ (n-Bu₄NBAR₄) as the electrolyte. Measurements with ν = 100, 200, 400 and 800 mV.s⁻¹ showed that the ratio of the anodic to cathodic peak currents varied linearly as a function of the square root of scan rate in all cases. The decamethylferrocene/decamethylferrocinium
(FeCp*₂/[FeCp*₂⁺) couple was used as an internal reference for potential measurements such that the couple falls at -0.55 V (CH₂Cl₂ / n-Bu₄NPF₆) or -0.62 V (CH₂Cl₂ / n-Bu₄NBF₄) relative to external FeCp₂/[FeCp₂⁺ at 0.00 V. 65,66 IR spectra IR spectroelectrochemical experiments at room temperature were recorded on a Bruker Vertex 70v FT-IR with an OTTLE cell equipped with a Pt-minigrid working electrode and CaF₂ windows. 67 The optical path of the cell was ca. 0.2 mm. The concentrations of the ferrocenyl compounds and the supporting electrolyte (n-Bu₄NBF₄) used in these measurements were 1.3 × 10⁻² and 3 × 10⁻¹ mol dm⁻³, respectively. Elemental analyses were performed at the London Metropolitan University.

**Preparation of [Ru{C=C(H)C(=CH)=CPh₂}(PPh₃)₂Cp]PF₆ ([4a]PF₆)**

A solution of 3a (100 mg, 0.438 mmol), [RuCl(PPh₃)₂Cp] (286 mg, 0.394 mmol) and KPF₆ (81 mg, 0.438 mmol) in CH₂Cl₂ (15 ml) was heated at reflux point for 20 hours. The solution was allowed to cool, concentrated in vacuo to ca. 2 ml and then added drop-wise to vigorously stirred, ice-cold diethyl ether causing precipitation of a pale red solid. The precipitate was collected by filtration, washed with hexanes (3 x 5 ml) and diethyl ether (3 x 5 ml) and air dried. Yield 329 mg, 79 %. Crystals suitable for X-ray diffraction were grown by slow diffusion of diethyl ether into a CH₂Cl₂ solution of the product. ¹H NMR (CD₂Cl₂): δ 3.09 (1H, s, H₅), 4.92 (1H, t, J = 2.5 Hz, H₃), 5.23 (5H, s, H₆), 7.02 – 7.05 (12H, m, H₇), 7.23 – 7.26 (15H, m, H₈ and H₉), 7.29 – 7.34 (4H, m, H₁₀ and H₁₁), 7.40 – 7.44 (6H, m, H₁₂, H₁₃ and H₁₄). ³¹P NMR (CD₂Cl₂): δ 41.62 (s). ¹³C NMR (CD₂Cl₂): δ 81.7 (C₈), 84.7 (C₉), 90.7 (C₁₀), 94.9 (C₁₁ t, J = 2 Hz), 104.2, 118.4, 127.8, 128.1 (C₁₂, C₁₃, C₁₄, C₁₅), 128.5 (C₁₆), 128.6 (C₁₇ or C₁₈ dd, Jₙ = J'ₙ = 5 Hz), 128.8 (C₁₉ or C₂₀ dd, Jₚ = J'ₚ = 5 Hz), 129.4, 130.4, 131.1 (C₂₁, C₂₂, C₂₃), 133.0 (C₂₄ dd, ²Jₙ = ⁵Jₚ = 5 Hz, 139.9,
140.5, 145.9 (C₆, C₇, C₈), C₉ not detected. MALDI MS(+) m/z 919.1 [M⁺]. IR (CH₂Cl₂) ν(C≡C) 2108 cm⁻¹, ν(CH=CRu) 1629 cm⁻¹, ν(C=C) 1483 cm⁻¹. Found C 66.53, H 4.40 % required C 66.58, H 4.45 %.

Crystal data for [4a]PF₆: C₅₉H₄₇P₂Ru × PF₆ × 2CH₂Cl₂, M = 1233.80, monoclinic, space group P 2₁/c, a = 12.1012(2), b = 31.5936(4), c = 14.8655(2) Å, β = 103.096(2)°, U = 5535.6(1) Å³, F(000) = 2512.0, Z = 4, Dc = 1.480 mg m⁻³, µ = 0.623 mm⁻¹. 92790 reflections were collected yielding 14013 unique data (Rₘₑᵣᵍ = 0.0549). Final wR₂(F²) = 0.1018 for all data (678 refined parameters), conventional R₁(F) = 0.0443 for 11904 reflections with l ≥ 2σ, GOF = 1.093.

Preparation of cis/trans-[Ru{=C=C(H)C(C≡CH)=CHFc}(PPh₃)₂Cp]PF₆ ([4b]PF₆)

A solution of 3b (100 mg, 0.384 mmol), [RuCl(PPh₃)₂Cp] (252 mg, 0.346 mmol) and KPF₆ (71 mg, 0.384 mmol) in CH₂Cl₂ (15 ml) was heated at reflux for 16 hours. After this time, the solution was allowed to cool, concentrated in vacuo to ca. 3 ml and filtered through Celite into a rapidly stirred diethyl ether to give a red precipitate. The precipitate was collected by filtration, washed with hexanes (3 × 5 ml) and diethyl ether (3 × 5 ml) and air dried. Yield 334 mg, 88% as a mixture of isomers, which was briefly characterised (MALDI MS(+) m/z 950.1 [M-H]⁺. IR (CH₂Cl₂): ν(C≡CH) 1980 cm⁻¹, ν(Ru=C=C) 1632 cm⁻¹; ν(C=C) 1586 cm⁻¹) before being used directly in the attempted preparation of 5b.

Preparation of [Ru{C≡CC(C≡CH)C≡CPh₂}(PPh₃)₂Cp] (5a)
A solution of [4a]PF₆ (85 mg, 0.08 mmol) in methanol (6 ml) was treated with KOBu¹ (42 mg, 0.399 mmol) and the mixture stirred for 10 minutes. The resulting yellow precipitate was collected by filtration, washed with methanol (3 × 5 ml) and dried in air. Yield 45 mg, 62 %. Crystals suitable for X-ray diffraction were grown by slow diffusion of methanol into a CH₂Cl₂ solution of the product.

¹H NMR (CDCl₃): δ 2.77 (s, 1H), 4.22 (s, 5H), 7.05 (t, J = 7.6 Hz, 12H), 7.13 (m, 2H), 7.16 (t, J = 7.4 Hz, 6H), 7.32 - 7.28 (m, 4H), 7.36 (m, 2H), 7.44 - 7.39 (m, 14H).

¹³C NMR (CDCl₃): δ 85.5 (C₅), 106.2, 113.4, 126.5, (C₆, C₇, C₈), 126.6 (C₉), 127.2 (C₆, dd, 3JCP = 6JCP = 5 Hz), 128.2 (C₅), 128.30 (C₇), 130.0, 130.4, 130.7, 131.0 (C₆, C₇, C₈, C₉), 133.8 (C₉, dd, 2JCP = 5JCP = 5 Hz), 138.6, 138.7, 138.8 (C₇, C₈, C₉), 142.3, 142.4 (C₅, C₆), 145.7 (C₇).³¹P NMR (CDCl₃): δ 50.35. MALDI MS(+): m/z 918.2 [M⁺]. IR (CH₂Cl₂) ν(C=CH) 2105 cm⁻¹, ν(C≡CRu) 2044 cm⁻¹, ν(C≡C) 1482 cm⁻¹.

Crystal data for 5a: C₅₉H₄₆P₂Ru × CH₂Cl₂, M = 1002.89, monoclinic, space group P 2₁/n, a = 10.4352(6), b = 24.6836(13), c = 18.6025(10) Å, β = 98.390(1)°, U = 4740.3(4) Å³, F(000) = 2064.0, Z = 4, Dc = 1.405 mg m⁻³, μ = 0.552 mm⁻¹. 83912 reflections were collected yielding 13820 unique data (Rmerge = 0.0307). Final wR²(F²) = 0.0792 for all data (586 refined parameters), conventional R₁(F) = 0.0294 for 11966 reflections with l ≥ 2σl GOF = 1.067.

Preparation of Ph₂C≡C(C≡CFe)₂ (6a)

Ethynyl ferrocene (652 mg, 3.10 mmol), 1a (500 mg, 1.48 mmol), [Pd(PPh₃)₄] (85.0 mg, 73.6 μmol) and Cul (6 mg, 3.15 μmol) were dissolved in triethylamine (25 ml). The reaction mixture was heated to reflux for 17 hours, cooled to ambient temperature and the solvent was removed in vacuo. The residue was purified by column chromatography (eluent: hexanes/dichloromethane (50:50 (v/v))) concentration of the relevant fractions afforded 6a in 47% yield (418 mg, 48%) as an orange solid. ¹H NMR (CDCl₃): δ 4.14 (5H, s, C₅H₅), 4.22 (2H, t, 3JH-H = 2 Hz, C₅H₅C≡C), 4.36 (2H, t, 3JH-H = 2 Hz, C₅H₅C≡C), 7.37–7.43 (6H, m, o-C₅H₅/p-C₅H₅), 7.53 (4H, m, m-C₅H₅). ¹³C NMR (CDCl₃): δ 65.2 (C-1, C₅H₅), 69.4 (C-3/C-5, C₅H₅), 70.3 (C₅H₅), 71.6 (C-2/C-5, C₅H₅), 85.1 (C≡C or (C₅H₅)₂C≡C), 91.6 (C≡C or (C₅H₅)₂C≡C), 103.4 (C≡C or (C₅H₅)₂C≡C), 128.1 (o-C₅H₅), 128.6 (p-C₅H₅), 130.5 (m-C₅H₅), 141.3 (((C₅H₅)₂C≡C)

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153.5 \((i-C_6H_5)\). ASAP MS\((+): m/z\) 597.1 [M+H]\(^+\). Analysis found: C 76.38, H 4.61%; required for \(C_{38}H_{28}Fe_2\): C 76.54, H 4.73%.

**Preparation of FcCH=C(C≡CFc)\(_2\) (6b)**

Ethynyl ferrocene (250 mg, 1.19 mmol), 1b (200 mg, 541 \(\mu\)mol), [Pd(PPh\(_3\))\(_4\)] (32 mg, 27.7 \(\mu\)mol) and Cul (5.00 mg, 27.7 \(\mu\)mol) were dissolved in triethylamine (20 ml). The reaction mixture was heated to reflux for 20 h, cooled to ambient temperature and the solvent was removed in vacuo. The residue was purified by column chromatography (eluent: hexanes to hexanes/dichloromethane (50:50 (v/v))) concentration of the relevant fractions afforded 6b (231 mg, 68%) as a bright red solid. Crystals suitable for X-ray diffraction were grown by slow evaporation from a dichloromethane solution. \(^1\)H NMR (CD\(_2Cl_2\)): \(\delta\) 4.25 (5 H, s, \(C_5H_5\)), 4.26 (2 H, m, \(C_5H_4\)), 4.27 (5 H, s, \(C_5H_5\)), 4.30 (2 H, m, \(C_5H_4\)), 4.33 (5 H, s, \(C_5H_5\)), 4.45 (2 H, m, \(C_5H_4\)), 4.49 (2 H, m, \(C_5H_4\)), 4.59 (2 H, m, \(C_5H_4\)), 4.93 (2 H, m, \(C_5H_4\)), 6.87 (1 H, s, \(CH=C\)). \(^{13}\)C NMR (CD\(_2Cl_2\)): \(\delta\) 65.7 (\(C-C≡C, C_5H_4\)), 65.8 (\(C-C≡C, C_5H_4\)), 69.3 (\(C-H, C_5H_4\)), 69.6 (\(C-H, C_5H_4\)), 69.9 (\(C-H, C_5H_4\)), 70.0 (\(C_5H_5\)), 70.3 (\(C_5H_5\)), 70.4 (\(C_5H_5\)), 70.6 (\(C-H, C_5H_4\)), 71.6 (\(C-H, C_5H_4\)), 71.7 (\(C-H, C_5H_4\)), 80.6 (\(C-CH=C, C_5H_4\)), 84.8 (\(C≡C\)), 86.2 (\(CH=C\)), 86.4 (\(C≡C\)), 93.3 (\(C≡C\)), 100.6 (\(C≡C\)), 142.5 (\(CH=C\)). ASAP-MS\((+): m/z\) 629.0 [M+H]\(^+\). Analysis found: C 68.82, H 4.50%; required for \(C_{36}H_{28}Fe_3\): C 68.84, H 4.49%.

**Crystal data for 6b:** \(C_{36}H_{28}Fe_3\), M = 628.13, monoclinic, space group \(Cc\), \(a = 18.7766(5), b = 12.8250(3), c = 11.6171(3)\) Å, \(\beta = 109.073(1)^\circ\), \(U = 2643.9(1)\) Å\(^3\), \(F(000) = 1288.0\), \(Z = 4\), \(D_c = 1.578\) mg m\(^{-3}\), \(\mu = 1.649\) mm\(^{-1}\). 15594 reflections were collected yielding 6689 unique data \((R_{\text{merge}} = 0.0319)\). Final \(wR_F(F^2) = 0.1101\) for all data \((330\) refined parameters), conventional \(R_F(F) = 0.0427\) for 5980 reflections with \(I \geq 2\sigma(I)\). GOF = 1.065, Flack = 0.41(2)
Preparation of \( \text{FcCH} = \text{C}(\equiv \text{CAuPPh}_3)_2 \) (7)

A solution of \( \text{2b} \) (100 mg, 0.247 mmol as a 25 mg/ml solution in THF)\(^{40}\) was treated with methanol (15 ml) and NaOH (99 mg, 2.47 mmol) and the solution stirred for 30 minutes. After this time, \([\text{AuCl(PPh}_3]\) (247 mg, 0.499 mmol) was added, and the reaction allowed to stir for a further 3 hours. The reaction mixture was filtered, and the precipitate washed with methanol (3 \times 10 ml) and hexane (3 \times 10 ml) and dried in air to give the title compound. Yield 206 mg, 71 \%.

\(^1\)H NMR (CD\(_2\)Cl\(_2\)): \( \delta 4.16 \) (5H, s, \( H_a \)), \( 4.26 \) (2H, vt, \( J = 2 \) Hz, \( H_c \)), \( 4.96 \) (2H, vt, \( J = 2 \) Hz, \( H_b \)), \( 6.56 \) (1H, s, \( H_e \)), \( 7.49 \) (12H, m, \( H_j \) and \( H_p \)), \( 7.54 \) (6H, m, \( H_l \) and \( j \)), \( 7.55 - 7.62 \) (12H, m, \( H_k \) and \( H_d \)). \(^{31}\)P NMR (CD\(_2\)Cl\(_2\)): \( \delta 42.20 \) (s), \( 42.41 \) (s). \(^{13}\)C NMR (CD\(_2\)Cl\(_2\)): \( \delta 69.65 \) (C\(_c \), s), \( 69.78 \) (C\(_b \), s), \( 69.84 \) (C\(_a \), s), \( 69.99 \) (C\(_m \), s), \( 70.02 \) (C\(_g \), s), \( 82.37 \) (C\(_d \), s), \( 102.91 \) (C\(_n \), d, \( J = 27 \) Hz), \( 103.39 \) (C\(_p \), s), \( 105.53 \) (C\(_h \), d, \( J = 27 \) Hz), \( 129.64 \) (C\(_o \), d, \( J = 5 \) Hz), \( 129.7 \) (C\(_r \), d, \( J = 5 \) Hz), \( 130.44 \) (C\(_o \), s), \( 139.22 \) (C\(_o \), s), \( 130.75 \) (C\(_o \), s), \( 132.04 \) (C\(_r \), s), \( 132.06 \) (C\(_r \), s), \( 134.65 \) (C\(_q \), C\(_d \), \( J = 5 \) Hz), \( 134.90 \) (C\(_b \), C\(_d \), \( J = 5 \) Hz). MALDI-MS(+): m/z 1176.1 [M]+. IR (CH\(_2\)Cl\(_2\)) \( \nu(C\equiv\text{CAu}) \) 2102 cm\(^{-1}\), \( \nu(C=\text{C}) \) 1481 cm\(^{-1}\).

Analysis found C 52.93, H 3.34% required C 53.06, H 3.34%.

X-ray Crystallography

The X-ray single crystal data have been collected using \( \lambda \text{MoK} \alpha \) radiation (\( \lambda =0.71073 \AA \)) on a Bruker SMART 6000 (fine-focus sealed tube, graphite monochromator, Monocap optics) (compounds \( \text{5a} \) and \( \text{6b} \)) and Agilent XCalibur.
(Sapphire-3 CCD detector, fine-focus sealed tube, graphite monochromator) \((4a)\) diffractometers equipped with a Cryostream (Oxford Cryosystems) open-flow nitrogen cryostats at the temperature 120.0(2) K. All structures were solved by direct method and refined by full-matrix least squares on \(F^2\) for all data using Olex2 [67] and SHELXTL [68] software. All non-disordered non-hydrogen atoms were refined anisotropically the hydrogen atoms were placed in the calculated positions and refined in riding mode. Disordered atoms in structure \(6b\) were refined isotropically with fixed SOF=0.5. Crystallographic data for the structure have been deposited with the Cambridge Crystallographic Data Centre as supplementary publication CCDC-1518244-1518246.

**Acknowledgments**

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**References**

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