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Binuclear methylphosphinidine complexes of cyclopentadienylruthenium carbonyls: effects of the higher ligand field strength of ruthenium derivatives relative to iron derivatives†

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The structures and energetics of the binuclear methylphosphinidene complexes of cyclopentadienyl-ruthenium carbonyls of the type $[\text{MePRu}_2(\text{CO})_n\text{Cp}_2]$ ($n = 4, 3, 2, 1$) have been investigated for comparison with their previously studied iron analogues. For the tetracarbonyls and tricarbonyls $[\text{MePM}_2(\text{CO})_n\text{Cp}_2]$ ($n = 4, 3$) substituting ruthenium for iron has relatively little effect on the energetically preferred structures. Thus such structures have two-electron donor bridging MeP groups with no metal–metal bond for the tetracarbonyls and a metal–metal single bond for the tricarbonyls. This leads to favored 18-electron configurations for both ruthenium atoms in all cases. However, the higher ligand field strengths of ruthenium complexes relative to analogous iron complexes have major effects on the energetically preferred structures for the dicarbonyls and monocarbonyls $[\text{MePM}_2(\text{CO})_n\text{Cp}_2]$ ($M = \text{Fe, Ru}; n = 2, 1$). Thus the 11 lowest energy structures for the dicarbonyl $[\text{MePFe}_2(\text{CO})_2\text{Cp}_2]$ are triplet or quintet spin state structures whereas the 6 lowest energy structures for the ruthenium analogue $[\text{MePRu}_2(\text{CO})_2\text{Cp}_2]$ are all singlet structures. These low-energy singlet $[\text{MePRu}_2(\text{CO})_2\text{Cp}_2]$ structures include species in which both ruthenium atoms attain the favored 18-electron configurations in different ways: either by a Ru–Ru single bond and an agostic C–H–Ru interaction from the methyl group, a Ru–Ru single bond and a four-electron donor bridging MeP ligand with P= Ru double bonds, or a formal Ru= Ru double bond with a two-electron donor bridging MeP ligand. The 8 lowest energy structures for the diiron monocarbonyl $[\text{MePFe}_2(\text{CO})\text{Cp}_2]$ are all triplet or quintet spin structures whereas the lowest energy structure for the diruthenium monocarbonyl $[\text{MePRu}_2(\text{CO})\text{Cp}_2]$ by more than 20 kcal mol^{−1} is a singlet structure with a formal Ru= Ru double bond and bridging CO and four-electron donor MeP groups. Thermochemical information predicts such monocarbonyl derivatives to be the dominant binuclear decarbonylation products of the tricarbonyls $[\text{RPRu}_2(\text{CO})_3\text{Cp}_2]$ by thermal or photochemical methods.

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1. Introduction

Alkyl- and arylphosphinidenes provide examples of species that are not isolable in the free state but that can occur as stable ligands in transition metal complexes, particularly as bridges between two metal atoms.¹ In such environments they can function either as two-electron donors or four electron donors. A two-electron donor phosphinidene ligand bridging a pair of metal atoms is similar to a bridging carbonyl group by forming P–M single bonds with each of the metal atoms (Fig. 1). This leaves a stereochemically active phosphorus lone pair so that the phosphorus atom lies significantly outside of the $[\text{M}_2\text{C}(\text{alkyl})]$ plane. Experimental examples of two-electron donor phosphinidene ligands in binuclear cyclopentadienyl-metal carbonyl chemistry include iron and ruthenium derivatives of the type $[(\mu\text{-RP})\text{M}_2(\mu\text{-CO})(\text{CO})_2\text{Cp}_2]$ ($\text{Cp} = \eta^5\text{-C}_5\text{H}_5$; $\text{M} =$

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†Electronic supplementary information (ESI) available: Table S1. Initial $[\text{MePRu}_2(\text{CO})_n\text{Cp}_2]$ structures. Table S2. Distance table for the lowest-lying $[\text{Cp}_2\text{Ru}_2\text{PMe}]$ structures. Table S3. Distance table for the lowest-lying $[\text{MePFe}_2(\text{CO})\text{Cp}_2]$ structures. Table S4. Distance table for the lowest-lying $[\text{MePRu}_2(\text{CO})_2\text{Cp}_2]$ structures. Table S5. Distance table for the lowest-lying $[\text{MePRu}_2(\text{CO})_3\text{Cp}_2]$ structures. Table S6. Distance table for the lowest-lying $[\text{MePRu}_2(\text{CO})_4\text{Cp}_2]$ structures. Table S7. Orbital energies and HOMO/LUMO gaps. Complete Gaussian reference (ref. 20). Separated concatenated xyz-file containing the Cartesian coordinates of the optimized structures which can be visualized with a variety of software such as Mercury CCDC which is a free program. See DOI: <https://doi.org/10.1039/d4dt02279c>

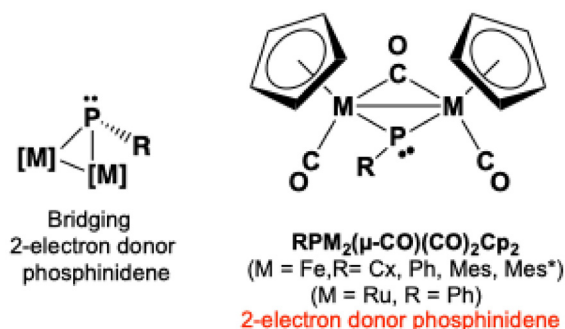


Fig. 1 Two-electron donor phosphinidene ligands and their occurrence in cyclopentadienylmetal carbonyl chemistry of iron and ruthenium.

Fe,² R = cyclohexyl, phenyl, mesityl, and 2,4,6-*t*Bu₃C₆H₂; M = Ru,³ R = phenyl). Such species are related to the well-known [Cp₂M₂(μ-CO)₂(CO)₂] (M = Fe,^{4–6} Ru^{7,8}) with a phosphinidene ligand replacing one of the carbonyl groups.

Phosphinidene ligands can also serve as four-electron donor bridging ligands through involvement of the phosphorus lone pair. Such a bridging four-electron donor phosphinidene ligand has either P=M double bonds or P → M dative bonds to each of the metal atoms (Fig. 2). Four-electron donor bridging phosphinidene ligands ideally have the phosphorus atom in the [M₂C(alkyl)] plane and shorter P=M or P → M distances than the P–M distances of two-electron donor bridging phosphinidene ligands. A four-electron donor bridging phosphinidene ligand is found in the experimentally known molybdenum complex [(μ-2,4,6-*t*Bu₃C₆H₂P)Mo₂(CO)₄Cp₂].⁹

Decarbonylation of the binuclear cyclopentadienyliron carbonyl [Cp₂Fe₂(μ-CO)₂(CO)₂] using different methods yields two stable products of interest (Fig. 3). Thus photolysis of [Cp₂Fe₂(μ-CO)₂(CO)₂] gives a tricarbonyl [Cp₂Fe₂(μ-CO)₃], which is of interest in representing a stable triplet state organometallic molecule with a formal Fe=Fe σ + 2/2π double bond similar to the O=O double bond in dioxygen.^{10–12} However, pyrolysis of [Cp₂Fe₂(μ-CO)₂(CO)₂] gives a very stable tetranuclear derivative [Cp₄Fe₄(μ₃-CO)₄] consisting of a central Fe₄ tetrahedron in which each face is bridged by a μ₃-CO carbonyl group.¹³ A binuclear [Cp₂Fe₂(CO)₂] derivative is a possible intermediate in the formation of the tetranuclear [Cp₄Fe₄(μ₃-CO)₄]. A Fe≡Fe formal triple bond is required for each iron atom to have the favored 18-electron configuration in such a [Cp₂Fe₂(CO)₂] derivative.^{14,15} Density functional theory studies on [Cp₂Fe₂(CO)_n] (n = 4, 3, 2, 1) systems are consistent with these experimental results.¹⁶

Experimental studies show the decarbonylation of the binuclear phosphinidene cyclopentadienyliron carbonyl complexes [(μ-RP)Fe₂(μ-CO)(CO)₂Cp₂] to be much more complicated than that of [Cp₂Fe₂(μ-CO)₂(CO)₂] discussed above.¹⁷ In particular, trinuclear and tetranuclear decarbonylation products were found in which the phosphinidene phosphorus lone pair becomes involved in the metal–ligand bonding. Furthermore, the structure of the decarbonylation product was found to be dependent on the alkyl or aryl group of the phosphinidene ligand.

In order to gain some insight into possible intermediates for the decarbonylation of [(μ-RP)Fe₂(μ-CO)(CO)₂Cp₂] derivatives we undertook a density functional theory study of the simplest such derivatives, namely the methylphosphinidene

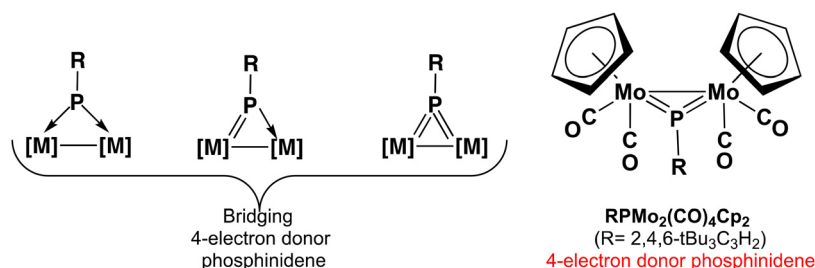


Fig. 2 Four-electron donor phosphinidene ligands and their occurrence in cyclopentadienylmetal carbonyl chemistry of molybdenum.

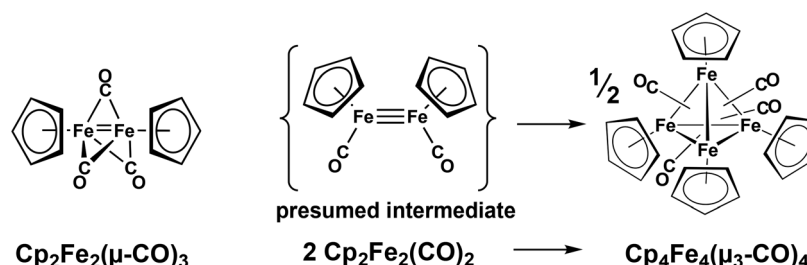


Fig. 3 Decarbonylation products of [Cp₂Fe₂(μ-CO)₂(CO)₂].



derivatives $[(\mu\text{-MeP})\text{Fe}_2(\text{CO})_n\text{Cp}_2]$ ($n = 4, 3, 2, 1$).¹⁸ The low-energy structures of the unsaturated dicarbonyl and monocarbonyl $[(\mu\text{-MeP})\text{Fe}_2(\text{CO})_n\text{Cp}_2]$ ($n = 2, 1$) systems were all found to be high-spin triplet and quintet structures rather than the singlet structures with iron-iron double and triple bonds found in their $[\text{Cp}_2\text{Fe}_2(\text{CO})_{n+1}]$ ($n = 2, 1$) analogues. The observation of low-energy high-spin $[(\mu\text{-MeP})\text{Fe}_2(\text{CO})_n\text{Cp}_2]$ ($n = 2, 1$) structures with significant spin density on the iron atoms suggests mechanistic possibilities for forming new iron-iron bonds leading to the experimentally observed trinuclear and tetranuclear clusters.

Substituting ruthenium for iron in the $[(\mu\text{-MeP})\text{M}_2(\text{CO})_n\text{Cp}_2]$ derivatives is expected to lead to stronger ligand fields thereby disfavoring the high spin structures observed as low-energy structures for the iron derivatives. Accordingly, we have now used similar density functional methods to investigate the ruthenium derivatives $[(\mu\text{-MeP})\text{Ru}_2(\text{CO})_n\text{Cp}_2]$ ($n = 4, 3, 2, 1$) related to the experimentally known³ $[\text{PhPRu}_2(\mu\text{-CO})(\text{CO})_2\text{Cp}_2]$. In contrast to the iron systems, we find that the low-energy structures for the unsaturated ruthenium derivatives $[(\mu\text{-MeP})\text{Ru}_2(\text{CO})_n\text{Cp}_2]$ ($n = 2, 1$) are singlet spin state structures with examples of structures with Ru= Ru formal double bonds and/or four-electron donor methylphosphinidine ligands. These theoretical studies predict that decarbonylation by photolysis or pyrolysis of ruthenium derivatives $[(\mu\text{-RP})\text{Ru}_2(\mu\text{-CO})(\text{CO})_2]$ is likely to lead to very different products than that of the corresponding iron derivatives.

2. Theoretical methods

The initial $[\text{MePRu}_2(\text{CO})_n\text{Cp}_2]$ chemical structures studied in this work have been designed by considering a $[\text{MePRu}_2\text{Cp}_2]$ unit followed by systematic placement of carbonyl groups as terminal and/or bridging ligands (coordinating to the metal ions through the carbon as well as both the carbon and oxygen atoms). Various Cp-Ru-CO orientations were also considered. This led to 95 different $[\text{MePRu}_2(\text{CO})_n\text{Cp}_2]$ ($n = 1$ to 4) starting structures computed as singlets, triplets, and quintets.

Full geometry optimizations were performed by using the PBE1PBE DFT functional¹⁹ and the def2-TZVP basis set as implemented in the Gaussian 09 software package²⁰ applying an ultrafine integration grid and tight convergence criteria. The energies include the zero-point and thermal corrections at 273 K. The nature of the stationary points was characterized by their harmonic vibrational frequencies. Saddle point structures with imaginary vibrational frequencies were reoptimized by following the normal modes to ensure that genuine minima were obtained. The $\nu(\text{CO})$ frequencies reported in the paper are scaled with a factor of 0.96. All of the lowest-energy structures have substantial HOMO-LUMO gaps ranging from 3.44 to 4.23 eV (Table S8 in the ESI†).

Only the lowest energy and thus potentially chemically significant structures are presented in detail in this paper. A larger number of structures of higher energy is presented in

the ESI.† The optimized structures are designated as **Ru2PCOn-mX**, where n designates the number of carbonyl groups, m designates the energy ordering in terms of relative energy as compared to the global minimum of each family, and X designates the spin state as S (singlet), T (triplet), or Q (quintet).

3. Results and discussion

3.1. Structures of the tetracarbonyl $[\text{MePRu}_2(\text{CO})_4\text{Cp}_2]$

The metal atoms in the tetracarbonyls $[\text{MePM}_2(\text{CO})_4\text{Cp}_2]$ ($M = \text{Fe}, \text{Ru}$) with a two-electron donor bridging MeP ligand have the favored 18-electron configuration without the need for a metal-metal bond. Thus the low-energy structures for the ruthenium tetracarbonyl derivative $[\text{MePRu}_2(\text{CO})_4\text{Cp}_2]$ are singlet structures that are very similar to those of its iron analogue¹⁸ (Fig. 4 and Table 1). The lowest energy such structure **Ru2PCO4-1S** has a long non-bonding Ru...Ru distance of ~ 4.20 Å and all terminal CO groups. The next higher energy $[\text{MePRu}_2(\text{CO})_4\text{Cp}_2]$ structure, namely **Ru2PCO4-2S** at 11.4 kcal mol⁻¹ in energy above **Ru2PCO4-1S**, also has a long non-bonding Ru...Ru distance of ~ 4.28 Å. However, one of the CO groups in **Ru2PCO4-2S** bridges an Ru-P bond whereas the other three CO groups remain terminal groups. The CO group in **Ru2PCO4-2S** bridging a Ru-P bond exhibits a low $\nu(\text{CO})$ frequency, namely 1734 cm⁻¹. This $\nu(\text{CO})$ frequency is not only more than 200 cm⁻¹ below the terminal $\nu(\text{CO})$ frequencies at 1958, 2008, and 2056 cm⁻¹ in **Ru2PCO4-2S**, similar to its iron analogue, but also ~ 100 cm⁻¹ below the $\nu(\text{CO})$ frequencies of $\mu\text{-CO}$ groups bridging Ru-Ru bonds in the $[\text{MePRu}_2(\text{CO})_3\text{Cp}_2]$ structures discussed below. The WBI values of 0.74 to 0.78 for the Ru-P bonds in **Ru2PCO4-1S** and **Ru2PCO4-2S** bonds are consistent with formal single bonds.

The lowest four unoccupied molecular orbitals of **Ru2PCO4-1S** (Fig. 5) have dominant components on the metal centers, suggesting a formal description of di-Ru(η) bound to a di-anionic phosphinidene.

3.2. Structures of the tricarbonyl $[\text{MePRu}_2(\text{CO})_3\text{Cp}_2]$

The metal atoms in the tricarbonyls $[\text{MePM}_2(\text{CO})_3\text{Cp}_2]$ ($M = \text{Fe}, \text{Ru}$) with a two-electron donor bridging MeP ligand have the favored 18-electron configuration if there is a metal-metal

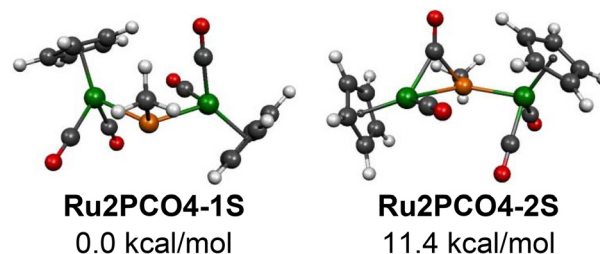
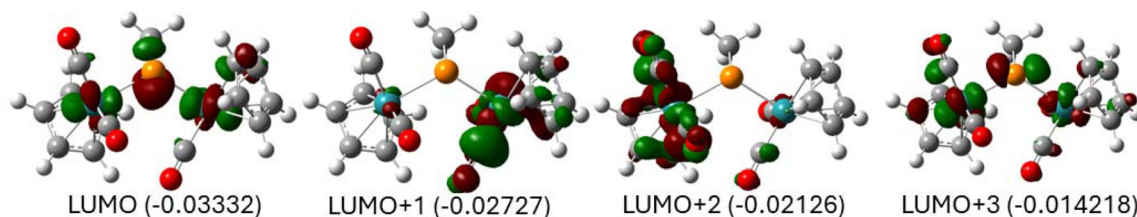


Fig. 4 The two lowest energy $[\text{MePRu}_2(\text{CO})_4\text{Cp}_2]$ structures lying within 20 kcal mol⁻¹ of the lowest energy structure.



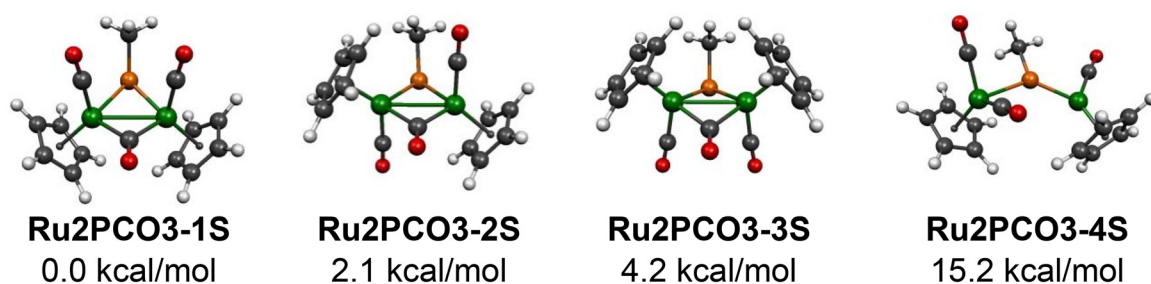
Table 1 The two [MePRu₂(CO)₄Cp₂] structures lying within 20 kcal mol⁻¹ of the lowest energy structure

Structure (symmetry)	ΔE	Ru–Ru interaction		Ru–P interactions		P distance (Å)	ΔE Fe
		Distance	WBI	Distances	WBI	Above Ru ₂ C plane	Analog
Ru2PCO4-1S (C_1)	0.0	4.202	0.02	2.45, 2.45	0.78, 0.76	0.79	1.1
Ru2PCO4-2S (C_1)	11.4	4.277	0.04	2.33, 2.37	0.74, 0.75	0.14	8.8

**Fig. 5** Lowest unoccupied orbitals of Ru2PCO4-1S.

formal single bond. The three lowest-energy [MePRu₂(CO)₃Cp₂] structures, namely the three stereoisomers **Ru2PCO3-1S**, **Ru2PCO3-2S**, and **Ru2PCO3-3S**, all lying within 5 kcal mol⁻¹ of energy, are of this type with Ru–Ru single bond distances of ~ 2.67 Å having WBI values of 0.33 and corresponding very closely to the three lowest-energy structures of their iron analogues [MePFe₂(CO)₃Cp₂] (Fig. 6 and Tables 2 and 3). The two-electron donor bridging MeP ligands in these three structures have Ru–P single bond distances of ~ 2.36 Å with the phosphorus atom ~ 1 Å outside of the Ru₂C(methyl) plane indicating a stereochemically active lone pair. One of the three CO groups in each of the three structures bridges the Ru–Ru bond. The remaining two CO groups are terminal CO groups with one on each ruthenium atom. The bridging CO groups in

each of the three structures **Ru2PCO3-1S**, **Ru2PCO3-2S**, and **Ru2PCO3-3S** exhibit a $\nu(\text{CO})$ frequency around 1820 cm⁻¹ which is significantly lower than the terminal $\nu(\text{CO})$ frequencies at 1980 cm⁻¹ and higher (Table 3). Both **Ru2PCO3-1S** and **Ru2PCO3-3S** have the pair of Cp rings in *cis* orientations relative to the Ru–Ru bond. However, in **Ru2PCO3-1S** the terminal CO groups are *cis* relative to the MeP bridge whereas in **Ru2PCO3-3S** the terminal CO groups are *trans* to the MeP bridge and the bridging MeP group. Structure **Ru2PCO3-2S** has the two Cp rings and the two terminal CO groups in *trans* orientations relative to the Ru–Ru bond. The WBI values for the Ru–P bonds in **Ru2PCO3-1S**, **Ru2PCO3-2S**, and **Ru2PCO3-3S** all fall in the range 0.77 to 0.80 consistent with formal single bonds.

**Fig. 6** The four lowest energy [MePRu₂(CO)₃Cp₂] structures lying within 31 kcal mol⁻¹ of the lowest energy structure.**Table 2** The four [MePRu₂(CO)₃Cp₂] structures lying within 31 kcal mol⁻¹ of the lowest energy structure

Structure (symmetry)	ΔE	Ru–Ru interaction		Ru–P interactions		P distance (Å)	ΔE Fe
		Distance	WBI	Distances	WBI	Above Ru ₂ C plane	Analog
Ru2PCO3-1S (C_s)	0.0	2.761	0.33	2.35, 2.35	0.80, 0.80	0.95	0.0
Ru2PCO3-2S (C_1)	2.1	2.774	0.33	2.35, 2.37	0.77, 0.79	1.01	3.3
Ru2PCO3-3S (C_s)	4.2	2.767	0.33	2.37, 2.37	0.77, 0.77	1.07	4.8
Ru2PCO3-4S (C_1)	15.2	4.098	0.06	2.15, 2.38	0.73, 1.46	0.06	—



Table 3 Harmonic $\nu(\text{CO})$ frequencies of the four $[\text{MePRu}_2(\text{CO})_3\text{Cp}_2]$ structures (scaled values in cm^{-1} and IR intensities in km mol^{-1} in parentheses) with the bridging $\nu(\text{CO})$ frequency in *italics*

Structure	$\nu(\text{CO})$ frequencies, cm^{-1}
Ru2PCO3-1S	1822(557), 1981(262), 2014(1286)
Ru2PCO3-2S	1818(556), 1978(1257), 1990(191)
Ru2PCO3-3S	1818(561), 1993(254), 2026(1411)
Ru2PCO3-4S	1956(1138), 2000(743), 2045(676)

The lowest unoccupied molecular orbitals of **Ru2PCO3-1S** (Fig. 7) include an empty σ^* Ru–Ru antibonding component (LUMO) alongside an occupied σ orbital (HOMO–4), confirming a net order of 1 with respect to the Ru–Ru interaction. Also, the fact that four LUMO's are located on the metals confirms a formal description of di-Ru(II) and implicitly a dianionic phosphinidene.

Our previous study of the $[\text{MePFe}_2(\text{CO})_3\text{Cp}_2]$ system¹⁸ revealed three triplet structures within 9 kcal mol^{-1} of the lowest-energy isomer analogous to **Ru2PCO3-1S**. For the analogous ruthenium $[\text{MePRu}_2(\text{CO})_3\text{Cp}_2]$ system no triplet structures were found within 31 kcal mol^{-1} of the lowest energy isomer **Ru2PCO3-1S** relating to the higher ligand field strength in ruthenium complexes relative to their iron analogues. However, in the ruthenium system $[\text{MePRu}_2(\text{CO})_3\text{Cp}_2]$ a fourth singlet structure **Ru2PCO3-4S** was found with an energy of 15.2 kcal mol^{-1} relative to that of **Ru2PCO3-1S** (Fig. 6 and Tables 2 and 3). An analogous iron structure was not found in the $[\text{MePFe}_2(\text{CO})_3\text{Cp}_2]$ system. Structure **Ru2PCO3-4S** differs from its three lower energy isomers in having a long non-bonding Ru...Ru distance of ~ 4.10 Å. Furthermore, the bridging MeP ligand in **Ru2PCO3-4S** is a four-electron donor with a formal $\text{P}=\text{Ru}$ double bond of length 2.15 Å and a WBI value of 1.46 to the ruthenium atom bearing a single terminal CO group and a dative $\text{P} \rightarrow \text{Ru}$ single bond of length 2.38 Å and WBI value of 0.73 to the ruthenium atom bearing two terminal CO groups. Furthermore, the phosphorus atom in the bridging MeP ligand of **Ru2PCO3-4S** lies only 0.06 Å out of the Ru_2C (methyl) plane confirming the participation of its lone pair in the ligand–metal bonding.

3.3. Structures of the dicarbonyl $[\text{MePRu}_2(\text{CO})_2\text{Cp}_2]$: major differences between the ruthenium and iron systems

The metal atoms in the dicarbonyls $[\text{MePM}_2(\text{CO})_2\text{Cp}_2]$ ($\text{M} = \text{Fe}, \text{Ru}$) with a two-electron donor bridging MeP ligand require a

formal $\text{M}=\text{M}$ double bond in order to attain the favored 18-electron configuration in a singlet spin state structure. However, for the iron system $[\text{MePFe}_2(\text{CO})_2\text{Cp}_2]$ no fewer than 11 triplet and quintet spin state isomers are found of lower energy than the lowest energy singlet structure lying 9.7 kcal mol^{-1} above the lowest energy higher spin state structure. The corresponding ruthenium system $[\text{MePRu}_2(\text{CO})_2\text{Cp}_2]$ is very different since six singlet structures are found with lower energies than the lowest energy triplet structure **Ru2PCO2-7T** lying 15.3 kcal mol^{-1} above the global minimum **Ru2PCO2-1S** (Fig. 8 and Tables 4 and 5). Also for the iron system $[\text{MePFe}_2(\text{CO})_2\text{Cp}_2]$ six quintet spin state structures were found within 8 kcal mol^{-1} of the lowest energy structure whereas for the analogous ruthenium system no quintet structures were found within 30 kcal mol^{-1} of the lowest energy structures. These differences are a clear reflection of the higher ligand field strength in complexes of the second row metal ruthenium relative to its first row analogue iron. This also suggests that the nature of the products formed by photochemical or thermal decarbonylation of $[\text{RPRu}_2(\text{CO})_3\text{Cp}_2]$ derivatives is likely to be very different than those formed by decarbonylation of their iron analogues.

The three lowest-energy $[\text{MePRu}_2(\text{CO})_2\text{Cp}_2]$ structures, namely **Ru2PCO2-1S**, **Ru2PCO2-2S**, and **Ru2PCO2-3S**, lying within 5 kcal mol^{-1} of energy, are a related series of singlet stereoisomers with single bridging CO and MeP groups (Fig. 8 and Table 4). The bridging MeP group is a two-electron donor as indicated by Ru–P single bond distances of ~ 2.27 and ~ 2.35 Å with WBI values in the range 0.79 to 1.03 and phosphorus distances 0.88 to 1.37 Å above the Ru_2C (methyl) plane. The Ru–Ru bonds in these three structures are clearly formal single bonds with Ru–Ru distances ranging from 2.73 to 2.77 Å and WBIs ranging from 0.33 to 0.37 Å. In both **Ru2PCO2-1S** and **Ru2PCO2-3S** there are C–H–Ru agostic interactions with Ru–H distances of 1.826 and 1.836 Å, respectively, of one of the methyl hydrogens to the ruthenium atom not bearing a terminal CO group. Such an agostic interaction effectively results in the donation of the electron pair forming a methyl C–H bond to the nearby ruthenium atom. This combined with the two-electron donor MeP ligands, the usual two-electron donor CO groups, and a single Ru–Ru bond gives each ruthenium atom in **Ru2PCO2-1S** and **Ru2PCO2-3S** the favored 18-electron configuration. Structure **Ru2PCO2-1S** has a *cis* orientation of the Cp rings and a *trans* orientation of the CO groups whereas **Ru2PCO2-3S** has a *trans* orientation of the Cp

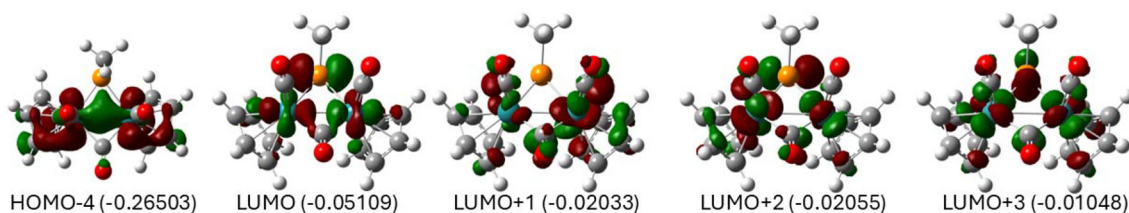


Fig. 7 Frontier molecular orbitals of **Ru2PCO3-1S**.



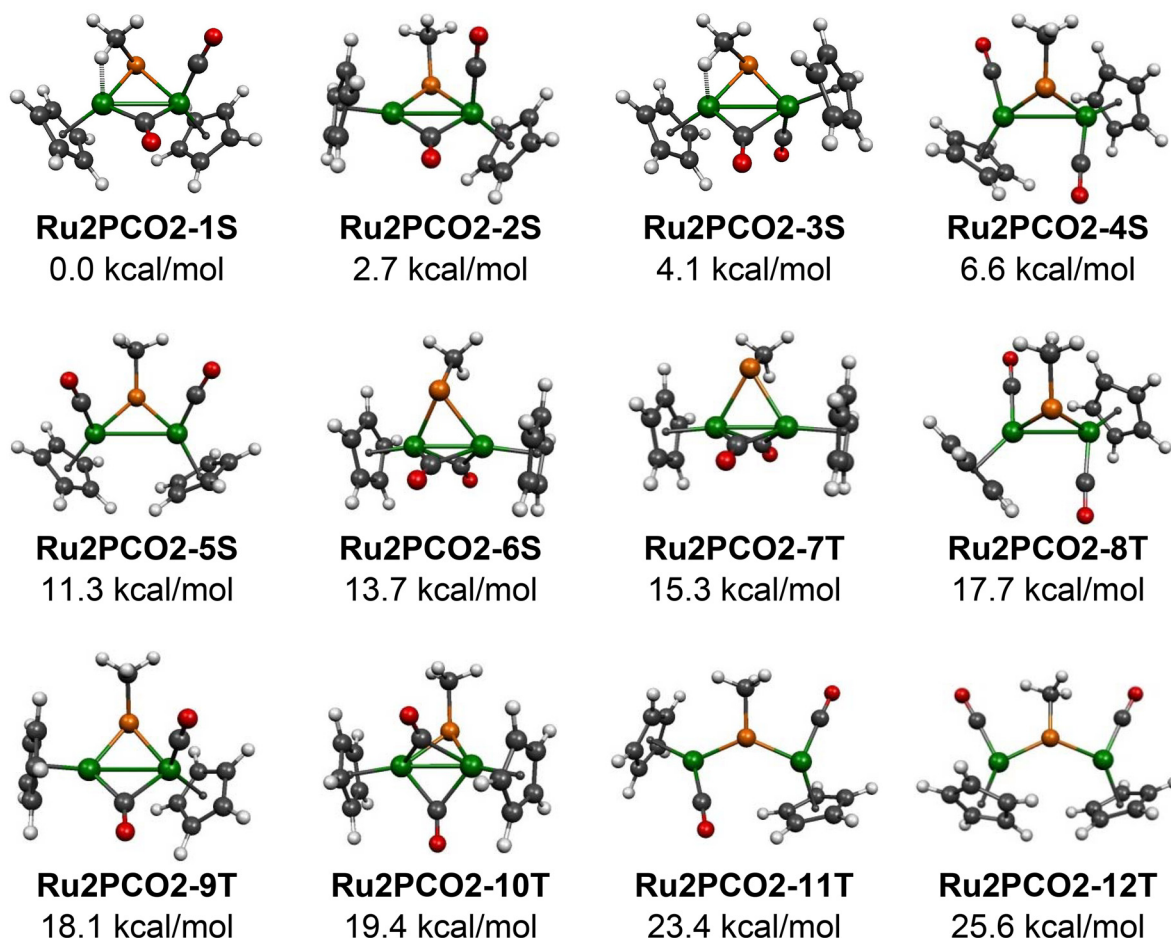


Fig. 8 The 12 lowest energy [MePRu₂(CO)₂Cp₂] structures lying within 30 kcal mol^{−1} of the lowest energy structure. The C–H–Ru agostic interactions in Ru2PCO2-1S and Ru2PCO2-3S are marked with dash lines.

Table 4 The 12 [MePRu₂(CO)₂Cp₂] structures lying within 30 kcal mol^{−1} of the lowest energy structure

Structure (symmetry)	ΔE	Ru–Ru interaction		Ru–P interactions		<i>P</i> distance (Å)	ΔE Fe
		Distance	WBI	Distances	WBI	Above Ru ₂ C plane	
Ru2PCO2-1S (<i>C</i> ₁)	0.0	2.752	0.37	2.29, 2.34	0.79, 0.82	1.29	
Ru2PCO2-2S (<i>C</i> ₁)	2.7	2.731	0.36	2.26, 2.34	1.03, 0.84	0.88	9.7
Ru2PCO2-3S (<i>C</i> ₁)	4.1	2.772	0.33	2.29, 2.36	0.80, 0.81	1.37	14.5
Ru2PCO2-4S (<i>C</i> ₁)	6.6	3.022	0.38	2.18, 2.19	1.15, 1.13	0.49	
Ru2PCO2-5S (<i>C</i> _s)	11.3	3.088	0.38	2.17, 2.18	1.08, 1.20	0.44	
Ru2PCO2-6S (<i>C</i> _s)	13.7	2.506	0.57	2.38, 2.38	0.78, 0.78	1.10	
Ru2PCO2-7T (<i>C</i> _s)	15.3	2.496	0.44	2.37, 2.37	0.76, 0.76	1.11	7.0
Ru2PCO2-8T (<i>C</i> _s)	17.7	2.751	0.46	2.23, 2.30	0.77, 0.97	0.78	
Ru2PCO2-9T (<i>C</i> _s)	18.1	2.748	0.33	2.24, 2.28	0.94, 0.89	0.78	0.0
Ru2PCO2-10T (<i>C</i> _{2v})	19.4	2.624	0.34	2.28, 2.28	0.88, 0.88	0.78	7.0
Ru2PCO2-11T (<i>C</i> _s)	23.4	4.053	0.09	2.21, 2.22	1.07, 1.13	0.08	8.6
Ru2PCO2-12T (<i>C</i> ₂)	25.6	4.031	0.10	2.22, 2.22	1.11, 1.09	0.01	6.9

rings and a *cis* orientation of the CO groups. Structure **Ru2PCO2-2S** does not have a similar C–H–Ru agostic interaction and both of its CO groups are the usual two-electron donor CO groups. Therefore the ruthenium atom in **Ru2PCO2-2S** not bearing the terminal CO group has only a 16-electron configuration.

The next two singlet [MePRu₂(CO)₂Cp₂] structures, namely **Ru2PCO2-4S** and **Ru2PCO2-5S**, lying 6.6 and 11.3 kcal mol^{−1} can be interpreted as having formal Ru–Ru single bonds and four-electron donor bridging MeP groups thereby giving each ruthenium atom the favored 18-electron configuration (Fig. 8 and Table 4). The lengths of the Ru–Ru single bonds in



Table 5 Harmonic $\nu(\text{CO})$ frequencies of the 12 $[\text{MePRu}_2(\text{CO})_2\text{Cp}_2]$ structures (scaled values in cm^{-1} and IR intensities in km mol^{-1} in parentheses) with the bridging $\nu(\text{CO})$ frequency in *italics*

Structure	$\nu(\text{CO})$ frequencies, cm^{-1}
Ru2PCO2-1S	1810(563), 1977(960)
Ru2PCO2-2S	1801(584), 1987(771)
Ru2PCO2-3S	1807(588), 1972(730)
Ru2PCO2-4S	1952(1692), 1964(63)
Ru2PCO2-5S	1955(715), 1981(1373)
Ru2PCO2-6S	1804(789), 1880(541)
Ru2PCO2-7T	1827(1027), 1857(256)
Ru2PCO2-8T	1944(1485), 1958(128)
Ru2PCO2-9T	1811(607), 1987(846)
Ru2PCO2-10T	1812(919), 1847(259)
Ru2PCO2-11T	1955(1268), 1968(670)
Ru2PCO2-12T	1959(1302), 1976(1280)

Ru2PCO2-4S and **Ru2PCO2-5S** are 3.022 and 3.088 Å, respectively, corresponding to WBI values of 0.38. These Ru–Ru single bond distances in **Ru2PCO2-4S** and **Ru2PCO2-5S** without a bridging CO group are ~ 0.3 Å longer than the Ru–Ru distances in **Ru2PCO2-1S**, **Ru2PCO2-2S**, and **Ru2PCO2-3S** with a bridging CO group. The P=Ru double bonds from the four-electron donor bridging MeP group to the ruthenium atom of lengths 2.18 Å are clearly shorter than the P–Ru single bonds of lengths ~ 2.3 Å in several of the structures discussed above. The WBIs of these P=Ru double bonds ranging from 1.08 to 1.20 are somewhat higher than the WBIs of the P–Ru single bonds in the carbonyl-rich $[\text{MePRu}_2(\text{CO})_n\text{Cp}_2]$ ($n = 4, 3$) structures discussed above.

Similarly to what is seen for **Ru2PCO3-1S**, the lowest unoccupied molecular orbitals of **Ru2PCO2-1S** (Fig. 9) include an empty σ^* Ru–Ru antibonding component (LUMO) consistent with a net order of 1 with respect to the Ru–Ru interaction, and four LUMO's located on the metals consistent with a formal description of di-Ru(II) and implicitly a di-anionic phosphinidene.

The $[\text{MePRu}_2(\text{CO})_2\text{Cp}_2]$ structure **Ru2PCO2-6S**, lying 13.7 kcal mol^{-1} in energy above **Ru2PCO2-1S**, is the lowest energy structure clearly having the formal Ru=Ru double bond required to give each ruthenium atom the favored 18-electron configuration in a structure with a two-electron donor bridging MeP group (Fig. 8 and Table 4). In **Ru2PCO2-6S** both CO groups bridge the relatively short Ru=Ru double

bond of length 2.596 Å with a WBI of 0.57 significantly higher than the WBI of the Ru–Ru single bonds found in other structures. The Ru–P single bonds in **Ru2PCO2-6S** have typical lengths of 2.38 Å and WBI values of 0.78 with the phosphorus atom 1.10 Å outside the $\text{Ru}_2\text{C}(\text{methyl})$ plane.

The next six $[\text{MePRu}_2(\text{CO})_2\text{Cp}_2]$ structures from **Ru2PCO2-7T** to **Ru2PCO2-12T** in energy above the six singlet structures **Ru2PCO2-1S** to **Ru2PCO2-6S** are triplet structures ranging in energy from 15.3 to 25.6 kcal mol^{-1} above **Ru2PCO2-1S** (Fig. 8 and Table 3). Five of these six triplet $[\text{MePRu}_2(\text{CO})_2\text{Cp}_2]$ structures have counterparts in low-energy structures in the analogous iron system $[\text{MePFe}_2(\text{CO})_2\text{Cp}_2]$.¹⁸ Structure **Ru2PCO2-7T** is the triplet counterpart of the singlet structure **Ru2PCO2-6S** with a Ru=Ru double bond of length 2.496 Å with a WBI of 0.44. This gives each ruthenium atom in **Ru2PCO2-7T** the favored 18-electron configuration leaving the two unpaired electrons in a Ru=Ru double bond of the $\sigma + 2/2\pi$ type analogous to the O=O double bond in dioxygen. Structure **Ru2PCO2-7T** can be related to the experimentally known^{10–12} triplet binuclear cyclopentadienyliron carbonyl derivative $[\text{Cp}_2\text{Fe}_2(\mu\text{-CO})_3]$ by replacement of iron with ruthenium and replacement of one of the bridging CO groups with a two-electron donor bridging MeP group. In addition **Ru2PCO2-7T** can be contrasted with **Ru2PCO2-10T**, also with a bridging two-electron donor bridging MeP group and two bridging $\mu\text{-CO}$ groups, but with a formal Ru–Ru single bond of length 2.624 Å having a WBI of 0.34. Thus in **Ru2PCO2-10T** the triplet spin state arises from a 17-electron configuration of each ruthenium atom rather than from the $\sigma + 2/2\pi$ Ru=Ru double bond in **Ru2PCO2-7T**.

The next two triplet $[\text{MePRu}_2(\text{CO})_2\text{Cp}_2]$ structures **Ru2PCO2-11T** and **Ru2PCO2-12T**, lying 23.4 and 25.6 kcal mol^{-1} , respectively, in energy above **Ru2PCO2-1S**, are a pair of stereoisomers with *trans* and *cis* orientations, respectively, of the single terminal CO group on each ruthenium atom (Fig. 8 and Table 4). The long Ru...Ru distances of ~ 4.04 Å in **Ru2PCO2-11T** and **Ru2PCO2-12T** suggest the lack of a formal ruthenium–ruthenium bond. The presence of the phosphorus atom less than 0.1 Å out of the $\text{Ru}_2\text{C}(\text{methyl})$ plane suggests a four-electron donor bridging MeP group. The P=Ru distances of 2.22 Å with WBI values ranging from 1.07 to 1.13 can be interpreted as double bonds from the phosphorus atom to each ruthenium atom. The combination of these double bonds, a terminal CO group on each ruthenium atom, and the

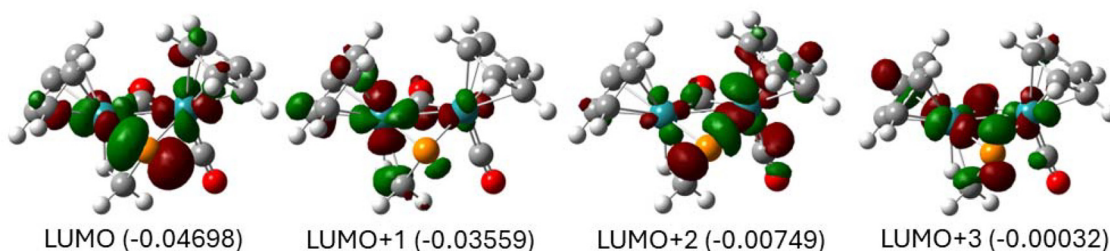


Fig. 9 Frontier molecular orbitals of **Ru2PCO2-1S**.



Cp ligand gives each ruthenium atom in **Ru2PCO2-11T** and **Ru2PCO2-12T** a 17-electron configuration consistent with a binuclear triplet.

3.4. Structures of the monocarbonyl [MePRu₂(CO)Cp₂]

The potential energy surface of the monocarbonyl [MePRu₂(CO)Cp₂] is remarkably simple since a single structure **Ru2PCO-1S** is found to lie a full 24 kcal mol⁻¹ below the next lowest energy structure (Fig. 10). The short P=Ru distances of 2.17 Å with WBI values of 1.2 suggest the possibility of formal double bonds corresponding to a four-electron donor bridging MeP group. Interpreting the Ru=Ru distance of 2.73 Å with a

WBI of 0.49 somewhat higher than the WBI values below 0.4 for Ru–Ru single bonds in singlet structures as a formal double bond gives each ruthenium atom in **Ru2PCO-1S** the favored 18-electron configuration. The single CO group in **Ru2PCO-1S** exhibits a $\nu(\text{CO})$ frequency of 1783 cm⁻¹ consistent with its bridging position. The iron analogue of **Ru2PCO-1S** in the [MePFe₂(CO)Cp] system is the lowest energy singlet structure but lies 16.7 kcal mol⁻¹ above the quintet spin state global minimum. This is another example of the effect of the larger ligand field strengths in ruthenium complexes relative to analogous iron complexes.

Of the frontier molecular orbitals of **Ru2PCO-1S** (Fig. 11), the four lowest-energy LUMOs are located on the ruthenium atoms consistent with two d⁶ centers. This formal di-Ru(II) description implies the phosphinidene ligand is acting as a dianion. The orbitals in Fig. 11 also indicate that the phosphorus atom is essentially non-hybridized, as expected.²¹ The spatial distribution of LUMO+1 suggests a two-electron three-center interaction of the two metals with a phosphorus p lone pair. In this interaction, the LUMO+1 is the π^* component, while the other two orbitals, HOMO–2 and HOMO–6, display π Ru–Ru bonding character. On the other hand, orbitals HOMO–1 and HOMO–5 indicate that the bond order for the σ interaction between the two metals is zero, as the antibonding HOMO–1 is doubly occupied. Thus, the net bond order for the Ru–Ru interaction is 1, arising from a π bond and with no σ bonding.

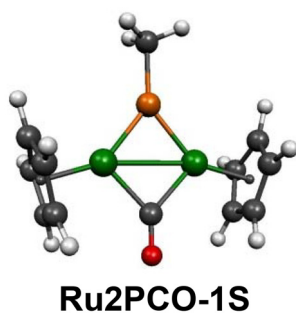


Fig. 10 The [MePRu₂(CO)Cp₂] structure lying ~24 kcal mol⁻¹ below the next lowest energy structure.

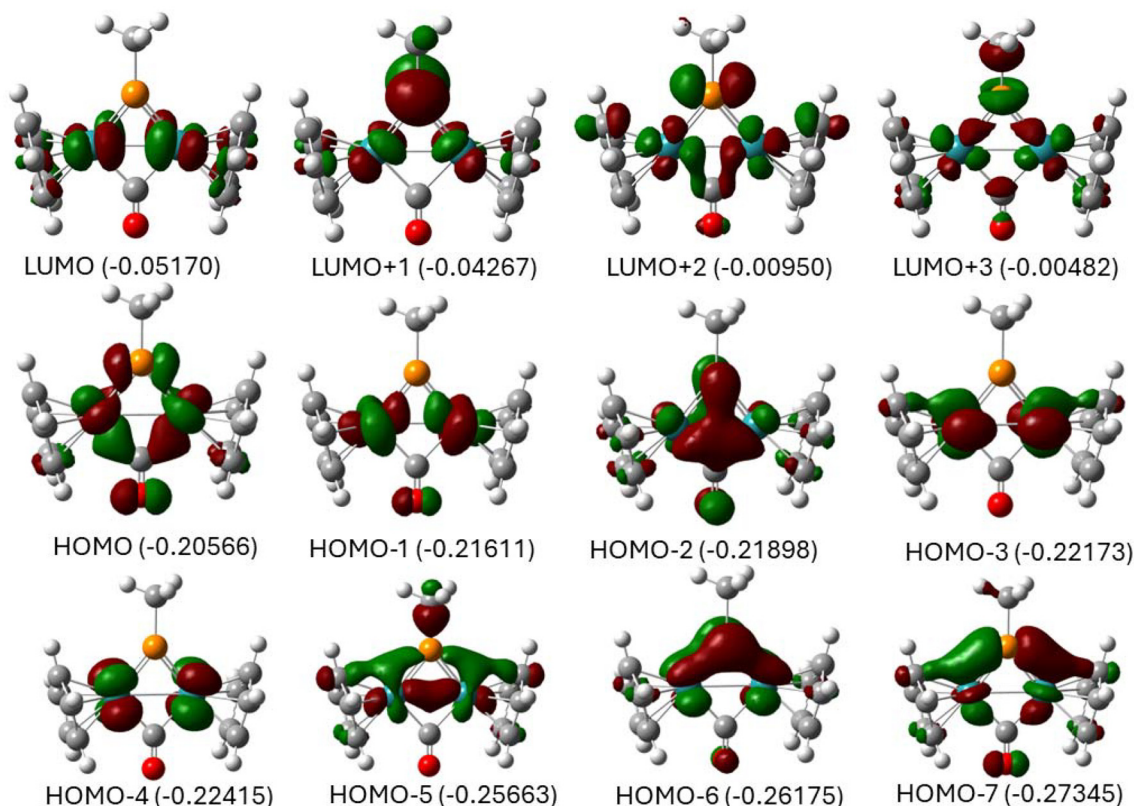


Fig. 11 Frontier molecular orbitals of **Ru2PCO-1S**.



Table 6 Carbonyl dissociation and disproportionation energies of the $[\text{MePRu}_2(\text{CO})_n\text{Cp}_2]$ derivatives (kcal mol^{-1}) considering the lowest energy structures

Reaction	ΔH	ΔG
$[\text{MePRu}_2(\text{CO})_4\text{Cp}_2]$ (Ru2PCO4-1S) $\rightarrow$ $[\text{MePRu}_2(\text{CO})_3\text{Cp}_2]$ (Ru2PCO3-1S) + CO	13.1	3.8
$[\text{MePRu}_2(\text{CO})_3\text{Cp}_2]$ (Ru2PCO3-1S) $\rightarrow$ $[\text{MePRu}_2(\text{CO})_2\text{Cp}_2]$ (Ru2PCO2-1S) + CO	42.9	32.7
$[\text{MePRu}_2(\text{CO})_2\text{Cp}_2]$ (Ru2PCO2-1S) $\rightarrow$ $[\text{MePRu}_2(\text{CO})\text{Cp}_2]$ (Ru2PCO-1S) + CO	27.5	14.1
$2 [\text{MePRu}_2(\text{CO})_2\text{Cp}_2] \rightarrow [\text{MePRu}_2(\text{CO})_3\text{Cp}_2] + [\text{MePRu}_2(\text{CO})\text{Cp}_2]$	−15.4	−18.6

3.5. Thermochemistry

Table 6 lists the carbonyl dissociation energies (ΔH and ΔG) for carbonyl dissociation processes of the type $[\text{MePRu}_2(\text{CO})_n\text{Cp}_2] \rightarrow [\text{MePRu}_2(\text{CO})_{n-1}\text{Cp}_2] + \text{CO}$ considering the lowest-energy structures, all of which are singlets. Such carbonyl dissociation processes are seen to be endothermic in all cases. However, carbonyl dissociation from the tetracarbonyl $[\text{MePRu}_2(\text{CO})_4\text{Cp}_2]$ to give the tricarbonyl $[\text{MePRu}_2(\text{CO})_3\text{Cp}_2]$ thereby forming a Ru–Ru bond is only weakly endothermic suggesting that this process can occur relatively easily. More significantly, combining the carbonyl dissociation energies for the tricarbonyl $[\text{MePRu}_2(\text{CO})_3\text{Cp}_2]$ and the dicarbonyl $[\text{MePRu}_2(\text{CO})_2\text{Cp}_2]$ shows that the disproportionation of the dicarbonyl into the tricarbonyl and monocarbonyl is significantly exothermic. This suggests that the decarbonylation of the stable tricarbonyl, such as the experimentally known³ $[\text{PhPRu}_2(\mu\text{-CO})(\text{CO})_2\text{Cp}]$, by photochemical or thermal methods is likely to lead directly to the monocarbonyl $[\text{MePRu}_2(\text{CO})\text{Cp}_2]$ for which structure **Ru2PCO-1S** is clearly a thermodynamic sink lying more than 20 kcal mol^{-1} below any of its isomers.

4. Conclusion

The potential energy surfaces for the tetracarbonyls $[\text{MePM}_2(\text{CO})_4\text{Cp}_2]$ and the tricarbonyls $[\text{MePM}_2(\text{CO})_3\text{Cp}_2]$ ($\text{M} = \text{Fe}, \text{Ru}$) are very similar for the iron and ruthenium systems. The energetically favored structures for both metals have a two-electron donor bridging MeP group without a direct metal–metal bond for the tetracarbonyls and a metal–metal single bond for the tricarbonyls thereby leading to the favored 18-electron metal configurations in all cases. The tetracarbonyls and tricarbonyls are thus saturated systems where the favored structures are not likely to be affected by the difference in ligand field strengths between analogous iron and ruthenium derivatives.

The situation is very different for the dicarbonyls and monocarbonyls $[\text{MePM}_2(\text{CO})_n\text{Cp}_2]$ ($n = 2, 1$; $\text{M} = \text{Fe}, \text{Ru}$). For the iron dicarbonyl system $[\text{MePFe}_2(\text{CO})_2\text{Cp}_2]$ the 11 lowest-energy structures are all triplet or quintet structures with the lowest-energy singlet isomer lying $\sim 10 \text{ kcal mol}^{-1}$ above the overall lowest-energy structure. However, for the ruthenium dicarbonyl system $[\text{MePRu}_2(\text{CO})_2\text{Cp}_2]$ the six lowest energy structures are all singlets with the lowest energy triplet isomer lying $\sim 15 \text{ kcal mol}^{-1}$ above the lowest energy structure. This

major difference between the preferred structure types for the iron and ruthenium derivatives of the type $[\text{MePM}_2(\text{CO})_n\text{Cp}_2]$ is clearly a manifestation of the effect of the higher ligand field strengths in ruthenium complexes relative to iron complexes.

A ruthenium derivative of the type $[\text{MePRu}_2(\text{CO})_2\text{Cp}]$ with a two-electron donor bridging MeP group requires a formal Ru= Ru double bond for each ruthenium atom to have the favored 18-electron configuration. An example of such a structure is **Ru2PCO2-6S** (Fig. 8 and Table 4). However, this structure lies 13.7 kcal mol^{-1} above its lowest energy isomer **Ru2PCO2-1S**. Both **Ru2PCO2-1S** and the slightly higher energy **Ru2PCO2-3S** have only a Ru–Ru single bond. However, in both of these structures there is an agostic C–H–Ru interaction between one of the methyl hydrogens and the ruthenium atom. The resulting donation of the two electrons from a methyl C–H bond to a ruthenium atom combined with a Ru–Ru single bond can give both ruthenium atoms the favored 18-electron configuration. The ruthenium atoms in the $[\text{MePRu}_2(\text{CO})_2\text{Cp}]$ isomers **Ru2PCO2-4S** and **Ru2CO2-5S** acquire the favored 18-electron configuration in a still different manner, namely by having four-electron donor bridging MeP groups rather than two-electron donor MeP groups combined with an Ru–Ru single bond. The four-electron donor bridging MeP groups in **Ru2PCO2-4S** and **Ru2CO2-5S** exhibit relatively short P= Ru distances of $\sim 2.18 \text{ \AA}$ suggesting formal double bonds.

The effect of the higher ligand field strength of ruthenium complexes relative to analogous iron complexes is also exhibited in the monocarbonyl systems $[\text{MePM}_2(\text{CO})\text{Cp}_2]$ ($\text{M} = \text{Fe}, \text{Ru}$). For the iron system $[\text{MePFe}_2(\text{CO})\text{Cp}_2]$ the eight lowest energy structures are triplet and quintet spin state structures with the lowest energy singlet structure lying 16.7 kcal mol^{-1} above the lowest energy isomer. The same type of singlet $[\text{MePRu}_2(\text{CO})\text{Cp}_2]$ structure in the analogous ruthenium system, namely **Ru2PCO-1S** with bridging CO and MeP groups, is not only the lowest energy such structure but also lies more than 20 kcal mol^{-1} below the next lowest energy isomer. The ruthenium atoms in **Ru2PCO-1S** attain the favored 18-electron configuration through a formal Ru= Ru double bond and its MeP bridge being a four-electron donor with P= Ru double bond distances of $\sim 2.15 \text{ \AA}$. Thermochemical information on CO dissociation energies predict the $[\text{RPRu}_2(\text{CO})\text{Cp}_2]$ structures of the type **Ru2PCO-1S** to be the preferred product from the decarbonylation of $[\text{RPRu}_2(\text{CO})_3\text{Cp}_2]$ derivatives by thermal or photochemical methods.



Data availability

The optimized coordinates of the structures presented in this manuscript are provided as a separate standard xyz-file which can be visualized with any visualiser program – we recommend using Mercury (from CCDC) which is a free program. The ESI† also contains the distance matrix between all the atoms, as provided by the Gaussian program. All of the Gaussian log files of all the structures presented in this manuscript are available upon request.

Conflicts of interest

The authors have no conflicts of interest to declare.

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References

- 1 M. E. García, D. García-Vivó, A. Ramos and M. A. Ruiz, Phosphinidene-bridged binuclear complexes, *Coord. Chem. Rev.*, 2017, **330**, 1–36.
- 2 C. M. Alvarez, M. G. Alvarez, M. E. García, R. González, M. A. Ruiz, H. Hamidov and J. C. Jeffery, High-yield synthesis and reactivity of stable diiron complexes with bent-phosphinidene bridges, *Organometallics*, 2005, **24**, 5503–5505.
- 3 S. Takemoto, Y. Kimura, K. Kammikawa and H. Matsuzaka, P–H bond addition to a dinuclear ruthenium imido complex: Synthesis and reactivity of an amido phosphide complex, *Organometallics*, 2008, **27**, 1780–1785.
- 4 O. S. Mills, The crystal structure of dicyclopentadienyl-diiron tetracarbonyl, *Acta Crystallogr.*, 1958, **11**, 620–623.
- 5 R. F. Bryan and P. T. Greene, Crystal structure of trans-di- μ -carbonyldicarbonyl-di- π -cyclopentadienyl-di-iron (Fe–Fe), a redetermination, *J. Chem. Soc. A*, 1970, 3064–3068.
- 6 A. Mitschler, B. Rees and M. S. Lehmann, Electron density in bis(dicarbonyl- π -cyclopentadienyliron) at liquid-nitrogen temperature by X-ray and ue, *J. Am. Chem. Soc.*, 1978, **100**, 3390–3397.
- 7 E. O. Fischer and A. Vogler, Dimeric cyclopentadienylrutheniumdicarbonyl, *Z. Naturforsch.*, 1962, **17b**, 421–422.
- 8 O. S. Mills and J. P. Nice, Structure of bis(cyclopentadienyl-dicarbonylruthenium), *J. Organomet. Chem.*, 1967, **9**, 339–344.
- 9 A. M. Arif, A. H. Cowley, N. C. Norman, A. G. Orpen and M. Pakulski, Transition-metal phosphinidene complexes: Syntheses, structures, and bonding in dinuclear phosphinidene complexes containing 14- and 15-electron metal fragments, *Organometallics*, 1988, **7**, 309–318.
- 10 J. V. Caspar and T. J. Meyer, Mechanistic aspects of the photochemistry of metal-metal bonds—evidence for the intervention of two different primary photoproducts in the photochemistry of $(\eta^5\text{-C}_5\text{H}_5)_2\text{Fe}_2(\text{CO})_4$, *J. Am. Chem. Soc.*, 1980, **102**, 7794–7795.
- 11 R. H. Hooker, K. A. Mahmoud and A. J. Rest, Spectroscopic evidence for the formation of tri- μ -carbonyl-bis $\{(\eta^5\text{-cyclopentadienyl})\text{iron}\}$ on photolysis of bis $\{[\text{dicarbonyl}](\eta^5\text{-cyclopentadienyl})\text{iron}\}$ in low-temperature matrices, *J. Chem. Soc., Chem. Commun.*, 1983, 1022–1024.
- 12 A. F. Hepp, J. P. Blaha, C. Lewis and M. S. Wrighton, Photochemistry of $(\eta^5\text{-C}_5\text{H}_5)_2\text{Fe}_2(\text{CO})_4$ and related complexes in rigid matrices at low temperature—loss of carbon monoxide from the trans isomer to yield triply CO-bridged species, *Organometallics*, 1984, **3**, 174–177.
- 13 R. B. King, Polynuclear cyclopentadienyl metal carbonyls of iron and cobalt, *Inorg. Chem.*, 1966, **5**, 2227.
- 14 F. A. Kvietok and B. E. Bursten, Stepwise photochemical CO loss from $\text{Cp}_2\text{Fe}_2(\text{CO})_2(\mu\text{-CO})_2$ in low-temperature matrices—evidence for an unsupported Fe $\equiv$ Fe triple bond, *J. Am. Chem. Soc.*, 1994, **116**, 9807–9808.
- 15 M. Vitale, M. E. Archer and B. E. Bursten, Theoretical and experimental studies of the unprecedented spin-dependent structures of $\text{Cp}_2\text{Fe}_2(\text{CO})_2$, the double-CO-loss product of $\text{Cp}_2\text{Fe}_2(\text{CO})_4$, *Chem. Commun.*, 1998, 179–180.
- 16 H. Y. Wang, Y. Xie, R. B. King and H. F. Schaefer, Unsaturation in binuclear cyclopentadienyliron carbonyls, *Inorg. Chem.*, 2006, **45**, 3384–3392.
- 17 M. A. Alvarez, M. E. García, R. González, A. Ramos and M. Ruiz, Synthesis and decarbonylation reactions of diiron cyclopentadienyl complexes with bent-phosphinidene bridges, *Organometallics*, 2011, **30**, 1102–1115.
- 18 O. Rudenco, A. Lupan, R. Silaghi-Dumitrescu and R. B. King, Decarbonylation products of binuclear methylphosphinidene complexes of cyclopentadienyliron carbonyls: triplet and quintet structures are favored energetically over singlet structures with iron-iron multiple bonding, *ACS Omega*, 2024, **9**, 12125–12134.
- 19 C. Adamo and V. Barone, Toward reliable density functional methods without adjustable parameters: The PBE0 model, *J. Chem. Phys.*, 1999, **110**, 6158–6169.
- 20 Gaussian09 (Revision E.01), Gaussian, Inc., Wallingford, CT, 2009. The complete reference is given in the ESI.†
- 21 R. Silaghi-Dumitrescu and F. J. Carrascoza-Mayen, A twist in the anomeric effect, *Stud. Univ. Babeş-Bolyai, Chem.*, 2014, **LIX**(3), 95–101.

