

A Density Functional Theory Investigation on Intramolecular Hydrogen Transfer of the $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$ Cluster

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The reduction of $[Os_3(CO)_{11}P(OMe)_3]$ and subsequent ionic coupling of the reduced species with $[Ru(\eta^5-C_5H_5)(CH_3CN)_3]^+$ resulted in the formation of $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$ which can be converted to spiked tetrahedral cluster, $[HOs_3(CO)_{11}P(OMe)_3Ru_2(\eta^5-C_5H_5)(C_5H_4)]$ via the intramolecular hydrogen transfer. Due to the unavailability of a suitable single crystal, the PW91/SDD and LDA/SDD density functional methods were used to predict possible structures and the available spectroscopic information (IR, NMR) of $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$. The most probable geometry found by constrained search is the isomer (**a2**) in which the phosphite, $P(OMe)_3$, occupies an axial position on one of the two osmium atoms that is edge bridged by the $Ru(CO)_2(\eta^5-C_5H_5)$ unit. By using the most probably geometry, the predicted infrared frequencies and 1H , ^{13}C and ^{31}P NMR chemical shifts of the compound are in the same range as the experimental values. For this type of complex, the LDA/SDD method is appropriate for IR predictions whereas the OPBE/IGLO-II method is appropriate for NMR predictions. The activation energy and reaction energy of the intramolecular hydrogen transfer coupled with the structural change of the transition metal framework were estimated at the PW91/SDD level to be 110.32 and -0.14 kcal/mol respectively.

Key Words: DFT, Osmium-carbonyl cluster, Intramolecular hydrogen transfer

Introduction

A transition metal cluster is a collection of bound transition metal atoms that exhibits distinguish physico-chemical properties from those of bulk or isolated atoms.¹ It can exist in various forms and nuclearities depending on its composition, stability and the synthetic pathway. Research in transition-metal cluster is now in focus again due to the recent interest in nanostructured materials for a much wider range of practical applications, including catalysis.²⁻⁴ Building large carbonyl clusters and exploring their physico-chemical properties has long been an active research area for many synthetic inorganic chemists including our research on building a transition metal cluster with mixed metal atoms with high nuclearity.⁵⁻⁸

An efficient pathway to build nuclearity clusters is the reduction of the parent cluster with subsequent ionic coupling between the parent cluster anion and a suitable cationic capping complex. The cationic capping species such as $Ru(\eta^5-C_5H_5)(CH_3CN)_3$ ²⁺,⁹ $[Ru(\eta^6-C_6H_6)(CH_3CN)_3]$ ^{2+,10} $[Os(\eta^6-C_6H_6)(CH_3CN)_3]$ ^{2+,11} and $[Ru(\eta^5-C_5H_5)(CH_3CN)_3]^+$ ¹², which are isolobal to the $[M(CO)_3]$ fragment,¹³ have been reacted effectively with anionic cluster compounds to build the higher nuclearity clusters. The reaction between $[Os_3(CO)_{11}P(OMe)_3]^{2-}$ with $[Ru(\eta^5-C_5H_5)(CH_3CN)_3]^+$ resulted in the formation of $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$.⁸ By using the empirical 18 electron rule, the structure of $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$ was predicted to be square-base pyramidal from the total electron count of

74. But there exists many possible isomers due to several three-dimensional atomic arrangements in this structure. This compound underwent intramolecular hydrogen transfer upon heating in toluene under reflux condition to yield $[HOs_3(CO)_{11}P(OMe)_3Ru_2(\eta^5-C_5H_5)(C_5H_4)]$.^{8,14} Although spectroscopic characterizations (MS, IR and NMR) were carried out on $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$ and $[HOs_3(CO)_{11}P(OMe)_3Ru_2(\eta^5-C_5H_5)(C_5H_4)]$ to determine their structures, the unavailability of suitable single crystal of $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$ for an X-ray diffraction experiment makes the other structural determination less conclusive. We chose to investigate possible geometries of these complexes by using computational techniques. The spectroscopic parameters were calculated and compared with available experimental data to validate our predicted structures.

Nowadays computational techniques have gained wider acceptance as a useful complementary tool to experimental techniques. The Density Functional Theory (DFT) method is now a method of choice for electronic structure calculation due to its accuracy at low computational cost. By specifying the exchange-correlation functional, the DFT method takes into account the electron correlation, which is crucial for transition metal complexes, while keeping the computational cost as low as that of the Hartree-Fock method.¹⁵ Common exchange-correlation functionals are the local density approximation (LDA), Perdew-Wang gradient corrected functional (PW91) and Becke's three parameters hybrid functional (B3LYP). Many literatures

use DFT calculations to help elucidating the geometry and spectroscopic information.^{3,16-18}

Maruo *et al.* synthesized tricarbonyl rhenium complexes by several synthetic pathways.³ Three complexes were formed and characterized by several techniques including DFT and time-dependent DFT methods. The calculations agree well with the experimental results and allow the authors to understand the stability of different complexes. Hunstock *et al.* compared the nonbridged (D_{3h}) and doubly bridged (C_{2v}) structures of tri-nuclear transition metal carbonyl clusters $M_3(CO)_{12}$, $M=Fe$, Ru and Os, calculated by using several methods.¹⁸ They found that DFT was capable to reproduce the energetic preference for the cluster structures with nonbridged carbonyl in osmium and ruthenium clusters. Lokbani-Azzouz *et al.* performed DFT calculations with LDA and BP functionals on several mixed transition metal-main group metal carbonyl clusters $M_4E_2(CO)_{12}$, $M=Fe$, Ru and Os, and $E=Bi$, As, P, N, CH and SiH, which exhibit an octahedral core.¹⁹ Two skeletal structures of M_4 in a square or butterfly form are possible. The LDA predicted the correct skeletal topologies of M_4 as a square for ruthenium clusters but the carbonyl ligands arrangement is not well predicted. This is acceptable since it is known that a transition from terminal to bridging position required a small amount of energy.¹ Two skeletal structures of osmium clusters are predicted to be close in energy at the LDA level. This suggests that the energy difference between both skeletal isomers decrease when M becomes heavier.

Being guided by the success of DFT in structural and spectroscopic prediction in the literature, we performed Density Functional Theory (DFT) calculations to predict the most probable structure of $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$ and its spectroscopic information. This method was also applied to $[HOs_3(CO)_{11}P(OMe)_3Ru_2(\eta^5-C_5H_5)(C_5H_4)]$ as a validity test. The calculation were verified with all available experimental data.

Methods

The synthesis and spectroscopic analysis of $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$ and $[HOs_3(CO)_{11}P(OMe)_3Ru_2((\eta^5-C_5H_5)(C_5H_4))]$ have been reported previously as has the structure of the latter cluster.⁸

The Density Functional Theory method (DFT) was used to predict the most probable structural isomer of $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$ which is the precursor to $[HOs_3(CO)_{11}P(OMe)_3Ru_2((\eta^5-C_5H_5)(C_5H_4))]$. The Local Density Approximation (LDA) and the Perdew-Wang 91 (PW91) exchange-correlation functionals were employed in combination with the relativistic Stuttgart-Dresden effective core potential (SDD) basis set. The charge density fitting method was used to reduce the computational time. Initially the performance of the chosen methods was verified by comparing the calculated $[HOs_3(CO)_{11}P(OMe)_3Ru_2((\eta^5-C_5H_5)(C_5H_4))]$ geometry with the available data from single crystal X-ray diffraction experiment.⁸

Due to a large number of possible isomers of $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$, the constrained search for the most probable structural isomer was carried out in two steps as following:

- Initially, the $P(OMe)_3$ ligand was initially replaced by a

CO ligand. The geometries of six possible arrangements of an edge-bridging tetrahedron of $[Os_3(CO)_{12}(Ru(\eta^5-C_5H_5))_2]$ as suggested in Figure 1 were optimized. The lowest energy isomer was selected.

II) A terminal CO ligand in the lowest energy isomer from previous step was replaced by the $P(OMe)_3$ ligand, hence yielding $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$. The steric consideration of the $P(OMe)_3$ ligand was taken into account during this step. The geometry optimization was performed on all generated structures. The most probable isomer is the lowest energy structure.

For all considered structures, the harmonic vibrational frequency was calculated to identify that the isomers are located at the minima on the potential energy surface. The NMR calculations was performed on the most stable isomer by using the IGLO-II basis set which is developed for the ESR and NMR calculations.²⁰ The calculated 1H and ^{13}C chemical shielding from tetramethylsilane, TMS, were chosen as a standard reference for 1H and ^{13}C chemical shift whereas the ^{31}P chemical shielding from $^{31}P(OMe)_3$ was chosen as a standard reference for ^{31}P chemical shift.

The activation and reaction energy of the intramolecular hydrogen transfer process were then estimated. The transition state structure connecting the $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$ and $[HOs_3(CO)_{11}P(OMe)_3Ru_2((\eta^5-C_5H_5)(C_5H_4))]$ was located by the Synchronous Transit-Guided Quasi-Newton (STQN) method developed by Schlegel.^{21,22}

All calculations were performed by using the GAUSSIAN 03 package.²³ The Avogadro software was used to render molecular structures.²⁴

Results and Discussion

Tables 1 and 2 reported some selected bond lengths and bond angles of $[HOs_3(CO)_{11}P(OMe)_3Ru_2((\eta^5-C_5H_5)(C_5H_4))]$ with the available data from single crystal X-ray diffraction experiment (see Figure 2 for numbering scheme).⁸ This complex has a special bonding arising from the carbon atom in C_5H_4 ring that has hydrogen transferred. This carbon atom acts as a bridging

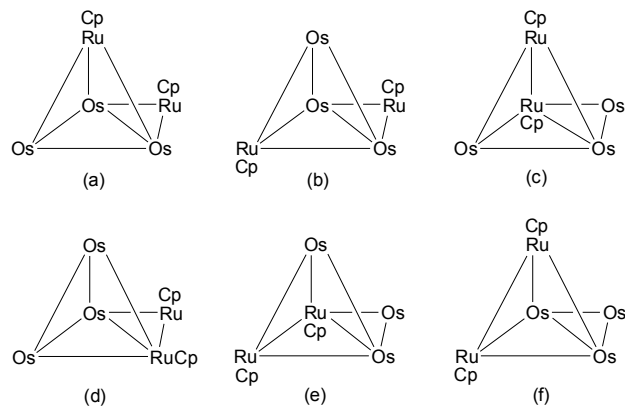


Figure 1. Six possible arrangements of ruthenium-cyclopentadienyl and osmium carbonyl fragments on edge-bridging tetrahedral geometry of $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$. The carbonyl and phosphite ligands were omitted for clarity; their positions were a subject of this study.

Table 1. Selected calculated bond lengths of $\text{HOs}_3(\text{CO})_{11}\text{P}(\text{OMe})_3\text{Ru}_2(\eta^5\text{-C}_5\text{H}_5)(\text{C}_5\text{H}_4)$ molecule compared to the data from single crystal X-ray experiment. The SDD basis set was used in calculations. The root-mean-square error (RMSE) was evaluated using the experimental value as the reference. See Figure 2 for the numbering scheme.

Bond length (Å)	PW91	LDA	Expt.
Os(1)-Os(2)	3.039	2.960	2.955
Os(1)-Os(3)	2.842	2.813	2.748
Os(2)-Os(3)	2.885	2.806	2.791
Os(1)-Ru(4)	2.790	2.737	2.877
Os(2)-Ru(4)	2.928	2.789	2.927
Os(3)-Ru(4)	2.772	2.737	2.893
Os(2)-Ru(5)	2.844	2.802	2.817
Os(2)-C(5')	2.446	2.571	2.432
Ru(5)-C(5')	2.104	2.037	2.116
Os(2)-C(52)	2.080	2.053	2.052
Ru(5)-C(52)	2.035	2.001	2.030
Os(1)-H _{transfer}	1.850	1.827	n.a.
Os(2)-H _{transfer}	1.830	1.841	n.a.
RMSE	0.067	0.092	

Table 2. Selected calculated bond angles of $\text{HOs}_3(\text{CO})_{11}\text{P}(\text{OMe})_3\text{Ru}_2(\eta^5\text{-C}_5\text{H}_5)(\text{C}_5\text{H}_4)$ molecule compared to the data from single crystal X-ray experiment. The SDD basis set was used in calculations. The root-mean-square error (RMSE) was evaluated using the experimental value as the reference. See Figure 2 for the numbering scheme.

Bond angle (°)	PW91	LDA	Expt.
C(1')-C(2')-Ru(4)	72.77	69.08	71.90
C(1')-C(5')-Ru(4)	68.10	66.31	69.40
C(2')-C(1')-Ru(4)	70.71	72.53	70.60
C(2')-C(3')-Ru(4)	70.60	70.21	71.00
C(3')-C(2')-Ru(4)	72.26	72.53	72.10
C(3')-C(4')-Ru(4)	71.10	71.29	72.10
C(4')-C(3')-Ru(4)	72.75	71.47	71.80
C(4')-C(5')-Ru(4)	68.69	69.45	68.50
C(5')-C(1')-Ru(4)	75.32	75.66	74.50
C(5')-C(4')-Ru(4)	74.65	72.94	73.90
O(21)-C(21)-Os(2)	179.41	179.05	179.00
O(22)-C(22)-Os(2)	172.44	172.22	172.10
O(51)-C(51)-Ru(5)	176.10	178.50	176.40
O(52)-C(52)-Os(2)	137.78	138.68	137.00
O(52)-C(52)-Ru(5)	134.73	133.71	135.70
Os(1)-H _{transfer} -Os(2)	111.3	107.6	n.a.
RMSE	0.72	1.57	

Table 3. The energy of six isomers (**a**)-(**f**) of $\text{Os}_3(\text{CO})_{12}(\text{Ru}(\eta^5\text{-C}_5\text{H}_5))_2$ relative to that of the lowest energy isomer (**a**)

$\text{Os}_3(\text{CO})_{12}(\text{Ru}(\eta^5\text{-C}_5\text{H}_5))_2$ isomers	Relative energy (kcal/mol)	
	PW91	LDA
(a)	0.00	0.00
(b)	17.81	17.30
(c)	29.01	26.44
(d)	25.21	26.81
(e)	28.80	27.70
(f)	9.32	8.91

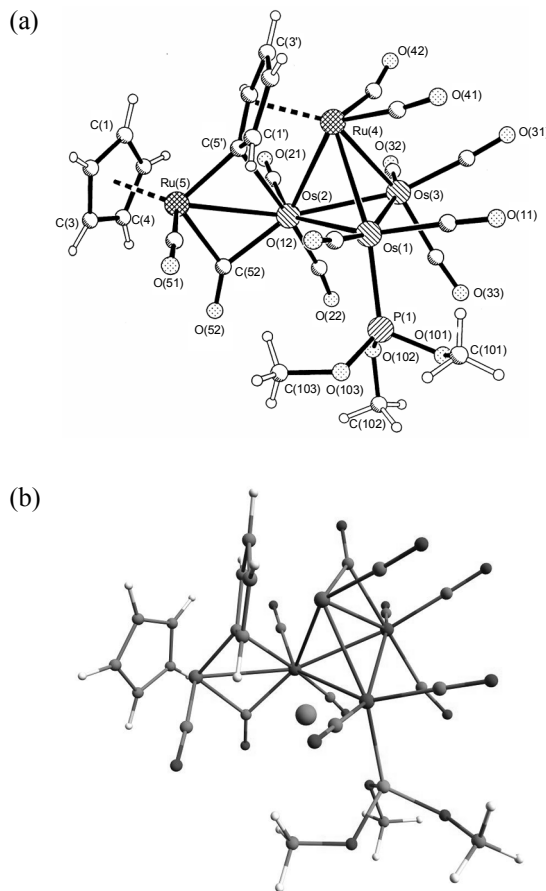


Figure 2. The structure of $\text{HOs}_3(\text{CO})_{11}\text{P}(\text{OMe})_3\text{Ru}_2(\eta^5\text{-C}_5\text{H}_5)(\text{C}_5\text{H}_4)$ from (a) X-ray and (b) PW91/SDD calculation with the numbering as used in Tables 1 and 2. The transferred hydrogen is highlighted.

ligand donating its electron to Ru(5) and Os(2) atoms. The optimized geometries from both functionals are in very good agreement with the experiment. The PW91/SDD performs slightly better than the LDA/SDD as a result of the generalized gradient approximation. As it is well known that X-ray diffraction experiment can not detect the hydrogen position, the calculation shows clear advantage in locating the transferred hydrogen in the complex. It situates in a bridging position between Os(1) and Os(2) atom with almost equal distances around 1.8 Å.

Six possible arrangements of $[\text{Os}_3(\text{CO})_{12}(\text{Ru}(\eta^5\text{-C}_5\text{H}_5))_2]$ as suggested in Figure 1 were optimized. Figure 3 displayed the PW91/SDD optimized structures of all isomers. The isomer (**a**) with one $\text{Ru}(\eta^5\text{-C}_5\text{H}_5)$ fragment at the edge-bridging position is the lowest energy isomer. Table 3 reports the energy of $[\text{Os}_3(\text{CO})_{12}(\text{Ru}(\eta^5\text{-C}_5\text{H}_5))_2]$ isomers relative to that of isomer (**a**). The greater energy of other isomers rules out the possibility of co-existence with the lowest-energy isomer. The metal arrangement of isomer (**a**) will be used in step II of constrained search. Both LDA and PW91 functionals yield the isomeric energy in the same order. The structure of isomers (**c**), (**d**) and (**e**) are less favorable due to the steric clash between the cyclopentadienyl rings on adjacent ruthenium atoms. The metal-metal bond distances of isomer (**a**) of $[\text{Os}_3(\text{CO})_{12}(\text{Ru}(\eta^5\text{-C}_5\text{H}_5))_2]$ range from 2.751 Å (Os(1)-Os(2)) to 2.934 Å (Os(2)-Os(3)) at the PW91/SDD level and from 2.690 Å (Os(1)-Os(2)) to 2.861 Å

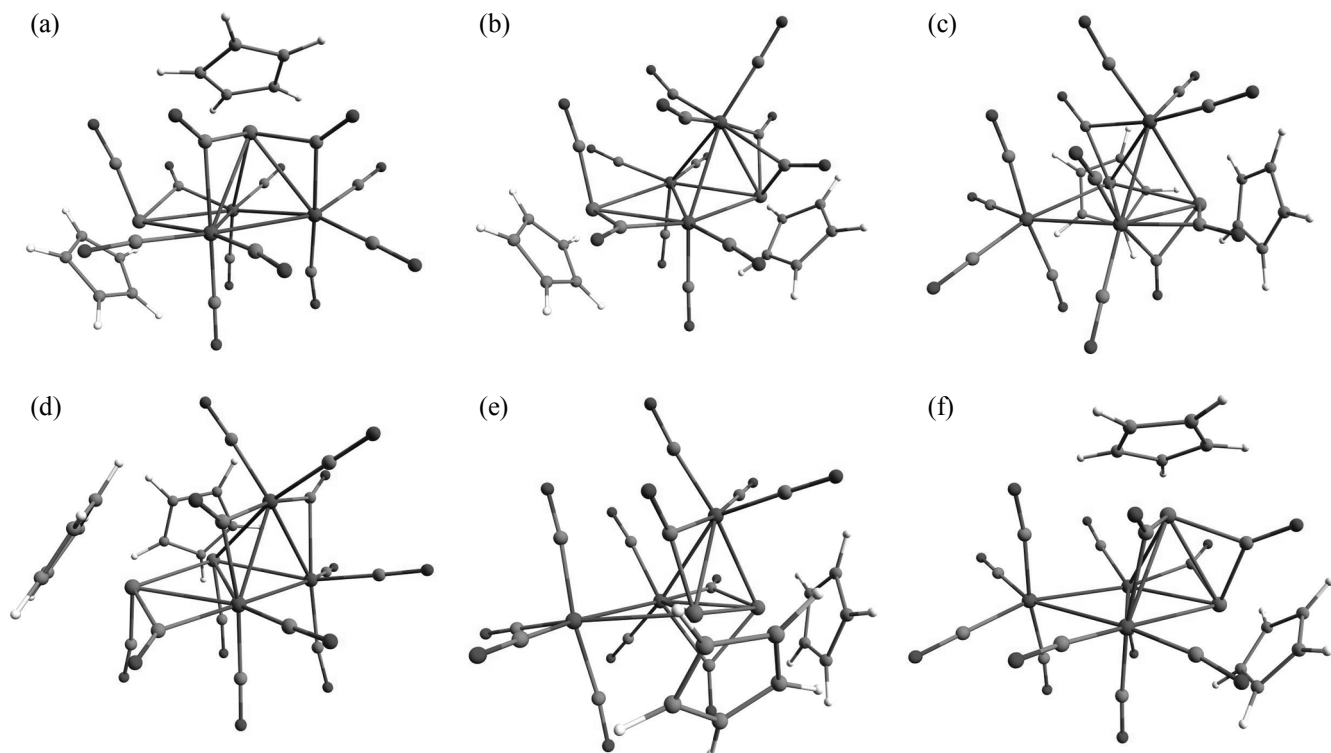


Figure 3. The PW91/SDD optimized geometries of six possible isomers (a)-(f) of $\text{Os}_3(\text{CO})_{12}(\text{Ru}(\eta^5\text{-C}_5\text{H}_5))_2$.

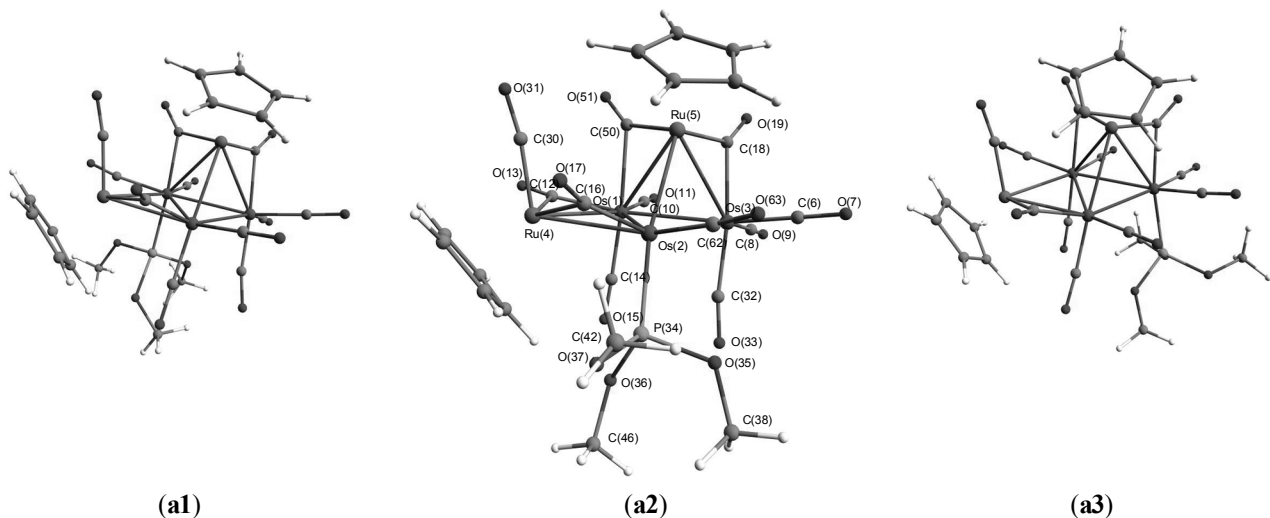


Figure 4. The PW91/SDD optimized geometries of three possible isomers (a1)-(a3) of $\text{Os}_3(\text{CO})_{11}\text{P}(\text{OMe})_3(\text{Ru}(\eta^5\text{-C}_5\text{H}_5))_2$. The numbering as used in Tables 5 and 6 is shown in the isomer (a2).

(Os(2)-Os(3)) at the LDA/SDD level. These distances are comparable to those found in other simple polynuclear osmium-ruthenium carbonyl clusters.²⁵ The PW91/SDD predicts bond lengths slightly greater than those from LDA/SDD method but both calculations yield reasonable bond lengths.

Taking isomer (a) as starting point for step II of constrained search, a carbonyl ligand was replaced by a bulky phosphite, $\text{P}(\text{OMe})_3$, ligand taking the steric effect into account. From the 107° cone angle of the phosphite ligand⁹ and the study of similar $\text{Os}_5(\text{CO})_n(\text{PMe}_3)$ compounds by Wang *et al.*,²⁶ the ligand replacement was limited to the three possible axial positions.

Figure 4 shows the optimized geometries of three isomers identified as (a1)-(a3).

Both the PW91/SDD and LDA/SDD methods indicated that the isomer (a2) is the most probable geometry of $[\text{Os}_3(\text{CO})_{11}\text{P}(\text{OMe})_3(\text{Ru}(\eta^5\text{-C}_5\text{H}_5))_2]$. Table 4 reports the energy of the three proposed isomers of $[\text{Os}_3(\text{CO})_{11}\text{P}(\text{OMe})_3(\text{Ru}(\eta^5\text{-C}_5\text{H}_5))_2]$ relative to that of isomer (a2). The energetic difference among these three isomers due to the relocation of phosphite ligand is much less than the energetic difference among six possible transition-metal arrangements in step I of constrained search. Hence we believe that a constrained search is justified.

Table 4. The energy of three isomers (**a1**)-(**a3**) of $\text{Os}_3(\text{CO})_{11}\text{P}(\text{OMe})_3(\text{Ru}(\eta^5\text{-C}_5\text{H}_5)_2$ relative to that of the lowest energy isomer (**a2**)

Isomer	Relative energy (kcal/mol)	
	PW91	LDA
(a1)	1.87	1.03
(a2)	0.00	0.00
(a3)	5.36	1.32

Table 5. The selected bond lengths of isomer (**a2**) of $\text{Os}_3(\text{CO})_{11}\text{P}(\text{OMe})_3(\text{Ru}(\eta^5\text{-C}_5\text{H}_5)_2$. See Figure 4 for the numbering scheme.

$\text{Os}_3(\text{CO})_{11}\text{P}(\text{OMe})_3(\text{Ru}(\eta^5\text{-C}_5\text{H}_5)_2$	Bond length (Å)	
	PW91	LDA
Os(1)-Os(2)	2.723	2.747
Os(1)-Os(3)	2.810	2.706
Os(2)-Os(3)	2.915	2.897
Os(1)-Ru(4)	2.872	2.753
Os(1)-Ru(5)	2.835	2.815
Os(2)-Ru(4)	2.903	2.812
Os(2)-Ru(5)	2.845	2.802
Os(3)-Ru(5)	2.779	2.665
Os(1)-C(12)	1.987	2.058
Ru(4)-C(12)	2.167	2.000
Os(2)-C(18)	2.200	2.162
Ru(5)-C(18)	1.961	1.938
Os(3)-C(20)	2.215	2.141
Ru(5)-C(20)	1.962	1.964
Ru(4)-C(38)	2.274	2.227
Ru(5)-C(26)	2.292	2.230
Os(2)-P(46)	2.367	2.316

Table 6. The selected bond angles of isomer (**a2**) of $\text{Os}_3(\text{CO})_{11}\text{P}(\text{OMe})_3(\text{Ru}(\eta^5\text{-C}_5\text{H}_5)_2$. See Figure 4 for the numbering scheme.

$\text{Os}_3(\text{CO})_{11}\text{P}(\text{OMe})_3(\text{Ru}(\eta^5\text{-C}_5\text{H}_5)_2$	Bond angle (°)	
	PW91	LDA
Os(1)-Ru(4)-Os(2)	56.27	59.14
Os(1)-C(10)-O(11)	177.76	176.06
Os(1)-C(12)-O(13)	145.99	139.03
Os(1)-C(14)-O(15)	176.20	176.10
Os(2)-C(16)-O(17)	155.91	173.78
Os(2)-C(62)-O(63)	176.40	174.45
Os(1)-Os(2)-P(46)	94.91	89.19
Os(3)-C(6)-O(7)	177.59	177.60
Os(3)-C(8)-O(9)	177.79	176.93
Os(3)-C(44)-O(45)	168.72	153.45
Ru(4)-C(42)-O(43)	171.08	170.83
Ru(5)-C(18)-O(19)	141.17	140.85
Ru(5)-C(20)-O(21)	141.70	140.53
Os(1)-Ru(4)-C(38)	135.48	140.74
Ru(4)-C(38)-C(36)	70.38	69.95
Os(2)-Ru(5)-C(26)	138.27	137.52
Ru(5)-C(26)-C(28)	72.14	71.47

Table 7. The CO stretching frequencies of $\text{Os}_3(\text{CO})_{11}\text{P}(\text{OMe})_3(\text{Ru}(\eta^5\text{-C}_5\text{H}_5)_2$

	v(CO) (cm ⁻¹)
PW91	1724, 1749, 1859, 1863, 1878, 1899, 1941
LDA	1738, 1785, 1925, 1935, 1956, 1966, 1989
Expt.	1716, 1772, 1933, 1976, 1991, 2000, 2039

The edge-bridging tetrahedral structure of $[\text{Os}_3(\text{CO})_{11}\text{P}(\text{OMe})_3(\text{Ru}(\eta^5\text{-C}_5\text{H}_5)_2]$ has the phosphite ligand attached to the Os(2) atom (see Figure 4). The next most likely phosphate position is at Os(1) atom. By simple consideration, the (**a1**) and (**a2**) should be equivalent by symmetry but the calculation however indicated a slight energy difference. This might be due to a small distortion in the optimized geometry. The (**a3**) isomer is the least probable (highest energy) among the three isomers. Although the PW91/SDD and LDA/SDD results provide a similar qualitative trend, the preferential position of phosphate ligand can be observed clearly in the PW91/SDD level which is a better approximation than the LDA/SDD level.

Tables 5 and 6 reported selected bond lengths and bond angles of the isomer (**a2**) respectively (see Figure 4 for numbering scheme). The Os(2)-P(46) bond length is 2.367 Å for the PW91/SDD level and 2.316 Å for the LDA/SDD level, respectively. There are seven terminal carbonyl ligands and four bridging carbonyl ligands spanning the Os(1)-Ru(4), Os(2)-Ru(4), Os(2)-Ru(5), and Os(3)-Ru(5) edges. Overall, as expected, the metal-carbon bonds connecting bridging carbonyl ligands are longer than those connecting terminal carbonyl ligands. The metal-metal bond distances are in the range of 2.723 Å (Os(1)-Os(2)) to 2.915 Å (Os(2)-Os(3)) at PW91/SDD level and of 2.665 Å (Os(3)-Ru(5)) to 2.897 Å (Os(2)-Os(3)) at LDA/SDD level. Upon replacing one CO by P(OMe)₃, the Os(1)-Ru(4)-Os(2) bond angles change slightly and the Os(2)-Ru(4) bond distances change only marginally (0.001 Å at PW91/SDD level and 0.004 Å at LDA/SDD level). The metal-metal bond lengths from both levels exhibit the same trend and lie in the expected range when compared to the experimental data of the other simple polynuclear osmium-ruthenium carbonyl clusters.²⁵

To verify our predicted geometry, the vibrational spectra and the NMR chemical shift were calculated and verified against available experimental data. The vibrational frequencies of $[\text{Os}_3(\text{CO})_{11}\text{P}(\text{OMe})_3(\text{Ru}(\eta^5\text{-C}_5\text{H}_5)_2]$ were calculated at both PW91/SDD and LDA/SDD levels. The calculated carbonyl stretching vibrational frequencies are summarized in Table 7. Both functionals yield the frequencies in the same range as the experimental values although they are not in a perfect quantitative agreement. It seems that LDA/SDD method is more appropriate for the IR calculation than the PW91/SDD method. From calculations, the stretching frequencies fall into two groups. The lower stretching frequency modes were identified to be those of bridging carbonyl ligands while the higher frequency modes correspond to those of terminal carbonyl ligands. Both methods reproduce the CO stretching of bridging carbonyl ligands well while those of terminal ones are underestimated. This might be due to the neglect of solvent effect in our calculation. We expect

Table 8. The predicted NMR chemical shift of $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$. The IGLO-II basis set was used in all calculations. To compare with the room-temperature experimental data, the calculated results were averaged among all equivalent nuclei.

Chemical Shift (ppm)	PW91	LDA	OPBE	Expt.
1H	4.18	4.53	3.90	3.58
1H	5.36	5.47	5.08	5.39
^{13}C	102.77	105.23	93.99	88.80
^{13}C	104.97	105.15	96.79	90.68
^{13}C	64.9	64.16	60.74	53.65
^{31}P	-16.56	-41.78	-39.78	-36.06

that, upon including solvent effect, the calculated stretching frequency will be red-shifted and match the experimental data better.

Table 8 reports the 1H , ^{13}C , and ^{31}P NMR chemical shifts of $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$ calculated by using the PW91, LDA and OPBE exchange-correlation functionals with the IGLO-II basis set. The PW91 and LDA functionals did not perform well when compared to the experimental chemical shift. The agreement with the experiment improves when the OPBE functional was used.²⁷ Given that it is much more difficult to compute the chemical shifts, our agreement from OPBE/IGLO-II method is acceptable. The calculation can be used to characterize and explain the experimental spectrum. The experimental 1H shifts at 3.58 and 5.39 ppm correspond to the proton from the methyl groups of the phosphite ligand and the cyclopentadiene ring, respectively. The ^{13}C shifts at 88.80 ppm and 90.68 ppm are from two non-identical cyclopentadienyl rings bonded to ruthenium atoms. The ^{13}C shift at 53.65 ppm is from the methyl carbon of the phosphite ligand. The experimental ^{31}P shift at -36.06 ppm of phosphite ligand was predicted by the OPBE/IGLO-II level to be -39.78 ppm which is in acceptable agreement.

Furthermore, the isomer (**a2**) has the structure that correlates best with the available single crystal X-ray structure of $[HOs_3(CO)_{11}P(OMe)_3Ru_2(\eta^5-C_5H_5)(C_5H_4)]$ which is the product of thermal intramolecular hydrogen transfer (see Figure 2). This experimental evidence helps justifying the employed DFT method which, in general case, might not be adequate to distinguish between isomers with small energy differences.^{28,29} With all available data, we therefore conclude that the isomer (**a2**) in which the phosphite, $P(OMe)_3$, occupies an axial position on one of the two osmium atoms that is edge bridged by the $Ru(CO)_2(\eta^5-C_5H_5)$ unit should be the most probable structure of $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$.

The intramolecular hydrogen transfer converts $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$ to $[HOs_3(CO)_{11}P(OMe)_3Ru_2((\eta^5-C_5H_5)(C_5H_4))]$. This involves breaking of the hydrogen-carbon bond from one cyclopentadienyl ring and making the hydrogen-osmium bond. This process caused a square-based tetrahedral framework to become a spiked tetrahedral framework by elongating one Os-edge bridged Ru bond. A corresponding transition state structure was located with a large imaginary frequency of 944.81 cm^{-1} corresponding to the hydrogen transfer motion from cyclopentadienyl ring to the bridging position of Os(1)-

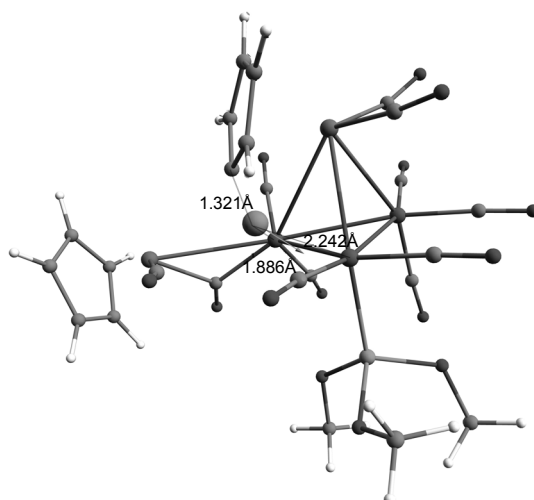


Figure 5. The PW91/SDD optimized transition state structure of intramolecular hydrogen transfer step with the transferred hydrogen being highlighted. The corresponding reaction coordinate is represented by the displacement arrow vectors.

Os(2) bond (see Figure 5). The intrinsic reaction coordinate calculations confirms that this transition state indeed links the correct reactant and product. This transition state structure has an elongated C(58)-H(63) bond length of 1.321 Å in comparison to other C-H bond lengths. The activation and reaction energy of intramolecular hydrogen transfer are 110.32 kcal/mol and 110.32 and -0.14 kcal/mol. They were estimated only at the PW91/SDD method. The LDA/SDD level was not used because the LDA functional was known to overestimate the energy involving in the breaking and forming of the chemical bond. The activation energy of this process is relatively high while the reaction energy change is very small. The almost zero value of reaction energy could possibly imply that reactant and product are stable to the same extent. The reason that both two structures does not immediately convert to each other might be explained from a very high activation energy. This is in agreement with what observed in the experiment that one needs to reflux the reactant at high temperature in order to convert it to become the product. A detailed investigation on the reaction mechanism of this important intramolecular hydrogen transfer will be discussed in separate communication.

Conclusion

By using a constrained search, the prediction of possible isomeric structure of $[Os_3(CO)_{11}P(OMe)_3(Ru(\eta^5-C_5H_5))_2]$ was carried out in two steps. This reduces the computational cost of exploring many possible isomers. The most possible structure is the isomer (**a2**) in which $P(OMe)_3$ occupies an axial position on one of the two osmium atoms that is edge bridged by the $Ru(CO)_2(\eta^5-C_5H_5)$ unit. This is verified by calculating the IR and NMR data and comparing them with the available experimental data. The calculated method must be carefully selected for the IR and NMR predictions while the structural prediction is less susceptible to the used method. The isomer (**a2**) also correlates well with $[HOs_3(CO)_{11}P(OMe)_3Ru_2((\eta^5-C_5H_5)(C_5H_4))]$

which is the product of intramolecular hydrogen transfer process. The high activation and almost zero reaction energy might explain the observation that two complexes are stable and can convert from one form to another only by reflux in toluene.

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