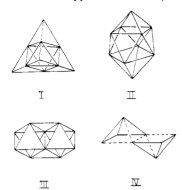
ligands appears at δ 457, a position that appears to be characteristic of Ru₆C cores.¹²

In comparison with the four previously characterized decanuclear clusters, Ru₁₀C₂(CO)₂₄²⁻ is unique in both its structure and its electron count (138 valence electrons). The framework of Os₁₀C(CO)₂₄²⁻¹³ is a tetracapped octahedron (see I) and its 134



valence electrons can be explained by Wade–Mingos rules. ^14,15 In contrast $Rh_{10}S(CO)_{22}^{2-,16}$ $Rh_{10}P(CO)_{22}^{3-,17}$ and $Rh_{10}As-(CO)_{22}^{3-18}$ each displays a bicapped square antiprism of metal atoms (see II) surrounding the non-metal atom and each has the 142 electrons predicted by Lauher. Using a similar treatment, Ciani and Sironi²⁰ predicted 134 electrons for a ten-atom D_{2h} framework formed by two edge-fused octahedra plus apical-apical bonds (see III). The distorted framework observed for Ru₁₀C₂-(CO)₂₄²⁻ suggests that the "extra" four electrons present in the real compound may occupy metal-metal antibonding orbitals largely localized on the apical ruthenium atoms, thereby affecting the apical-apical interactions primarily and other bonds to the apical atoms to a lesser extent.

The electron count displayed by Ru₁₀C₂(CO)₂₄²⁻ may be rationalized in the following way, which depends on the fact that a 74-electron square pyramid (e.g., Ru₅C(CO)₁₅) has 18 electrons per metal atom, if localized M-M bonds are assumed, but an 86-electron octahedron (e.g., Ru₆C(CO)₁₇) does not. ^{15,19} Removing two Ru(CO)₂ caps (maintaining inversion symmetry) leaves two square pyramids sharing a basal edge (see IV). This structure may be viewed as two discrete $Ru_4C(CO)_{10}$ "butterfly" units connected in a slipped fashion. The connection involves *five* Ru-Ru contacts between the subunits, which satisfies the 5electron deficiency of each Ru₄C(CO)₁₀ moiety (cf. Fe₄C- $(CO)_{12}^{2-}$. Thus, the $Ru_8C_2(CO)_{20}^{2-}$ framework (114 electrons) is electron precise and adding the two capping Ru(CO)₂ units (each 12 electrons and providing no extra framework pairs¹⁵) gives the observed formulation.22

(12) The value observed for $(CH_3CN)_2Cu_2Ru_6C(CO)_{16}$ is δ 458.¹⁰ The shift reported⁸ for $Ru_6C(CO)_{16}^{-2}$ has been redetermined as δ 459: Bradley, J. S., personal communication.

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Dicarbide clusters have been observed previously only for cobalt and rhodium; two general classes have been characterized. In one class, represented by $Co_{13}C_2(CO)_{24}^{4-23}$ and $Rh_{15}C_2(CO)_{23}^{-24}$ the carbon atoms are well separated and occupy two distinct cavities (trigonal prismatic for Co, octahedral for Rh) within the cluster framework. The second class, consisting of $Co_{11}(C_2)$ - $(CO)_{22}^{3-25}$ and $Rh_{12}(C_2)(CO)_{25}$, 26 populate a single cavity with a C_2 unit. Although $Ru_{10}C_2(CO)_{24}^{2-2}$ is a member of the first class, its more condensed structure brings the two carbon atoms in closer proximity. This suggests the possibility of conversion to the second class under appropriate conditions.

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Supplementary Material Available: Listings of atomic coordinates, bond lenghts, and bond angles (13 pages). Ordering information is given on any current masthead page.

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Reactions of Nucleophiles with α -Halo Ketones¹

Glen A. Russell* and Francisco Ros²

Department of Chemistry, Iowa State University Ames, Iowa 50011

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Nucleophilic substitutions occurring by electron-transfer radical chain processes (S_{RN}1) at the α -C atoms of α -nitro ketones and esters³ or of α -halomercury ketones⁴ are recognized. Since the radical anion of α -bromo-p-nitroacetophenone rapidly loses bromide ion, $^5 \alpha$ -halo ketones are also candidates for $S_{RN}1$ processes.

We have examined the reactions of sterically hindered α haloisobutyrophenones (1) with Me₂C=NO₂⁻ and found that with p-nitro or p-cyano substituents competing ionic and free-radical substitution processes lead to different products. However, since the free radical process is not observed in the reaction of a variety

⁽²²⁾ If the Ru₁₀ cluster is divided into two Ru₅C(CO)₁ subunits, each subunit is five electrons deficient, but there are seven Ru-Ru contacts. However, an 86-electron octahedral complex has formally only 11 M-M bonds (108 - 86 = 22) even though there are 12 M-M contacts in the octahedral frame.¹⁵ (Note that the equivalent geometry for 86 electrons, a capped square pyramid, clearly has only 11 M-M bonds.¹⁹) Thus, if in joining the Ru₅C subunits, one of the Ru-Ru contacts per octahedron formed is not counted as bonding, the five-electron deficiency per subunit is satisfied by five Ru-Ru bonds.

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Table I. Reaction of p-YC₆H₅COCMe₂X (1) with Me₂C=NO₂"M* in Me₂SO^a

	Y	M ⁺	conditions ^b	time, min	yield, % ^c			
X					2	3	4	1
Cl	NO,	Li ⁺	hv	80	38	7	36	14
	-	K ⁺	$h\nu$	80	41 (36)	16 (6)	32 (26)	6
		K ⁺	dark	80	29	19	36	7
		K ⁺	hv, 10 mol % p-DNB	45	28	16	31	20
		K ⁺	$h\nu$, 10 mol % (Me ₃ C) ₂ NO·	80	0	18	69	6
		K+, 18-crown-6	hv	80	77	10	10	0
Br	NO,	Li ⁺	$h\nu$	45	19	45	24	0
	•	K ⁺	$h\nu$	15	8 (6)	62 (45)	4(1)	0
		K*	$h\nu$, 20 mol % (Me ₃ C), NO·	15	0	61	5	6
		K ⁺ , 18-crown-6	hν	45	51	28	4	0
		K+, 18-crown-6	dark	45	40	28	11	4
		K ⁺ , 18-crown-6	$h\nu$, 20 mol % p -DNB	45	27	48	5	6
		K+, 18-crown-6	$h\nu$, 20 mol % (Me ₃ C) ₂ NO	45	0	78	6	4
		K ⁺	$h\nu$	60	51 (40)	12	18	0
		K ⁺	dark	60	9	19	48	5
		K ⁺	hv, 15 mol % p-DNB	60	7	18	32	23
Cl	CN	K ⁺	$h\nu$, 15 mol % (Me ₃ C) ₂ NO·	60	0	21 (14)	49 (39)	13
		K+, 18-crown-6	$h\nu$	60	69	5	12	0

^a Standard conditions: the α-halo ketone (1 mmol) in Me₂SO was added with stirring under N₂ to Me₂C=NO₂M in Me₂SO, prepared in situ from Me₃COM, or Me₃COK/18-crown-6 (1/1) (1.05 mmol) and Me₂CHNO₂ (1.05 mmol) to give solution with [Li⁺] = [K⁺, 18-C-6] = 0.1 M; [K⁺] = 0.3 M (Me₂C=NO₂K was initially insoluble in Me₂SO). ^b hν: irradiation with a 300-W sunlamp at ca. 50 cm. Dark: flask was wrapped with aluminum foil. ^c Crude yields by ¹H NMR analysis; numbers in parentheses represent yields of pure isolated products by TLC (silica gel; C₆H₆-ethyl acetate); all products gave satisfactory elemental analyses and ¹H NMR, IR, and mass spectral data.

Scheme I

1a,
$$X = Cl$$
; $Y = NO_2$
a', $X = Rr$; $Y = NO_2$
b, $X = Cl$; $Y = NO_2$
c, $X = Cl$; $Y = H$

2a, $Y = NO_2$
b, $Y = CN$

2a, $Y = NO_2$
b, $Y = CN$

3a, $Y = NO_2$
b, $Y = CN$
c, $Y = H$

4a, $Y = NO_2$
b, $Y = CN$
c, $Y = H$

of nucleophiles with the unsubstituted $\alpha\text{-chloroisobutyrophenone,}$ we conclude that substitution in $\alpha\text{-halo}$ ketones by an electron-transfer mechanism is a process of limited scope even when S_N2 substitution is sterically hindered.

The reactions of 1a, 1a', or 1b with $Me_2C=NO_2^-$ in Me_2SO gave the β -nitro ketones 2, the oxiranes 3, and the α -hydroxy ketones 4 (Scheme I), while 1c gave only 3c (60%) and 4c (12%). Table I summarizes the products from the reactions of 1a,b as a function of irradiation, the presence of free-radical scavengers, and the nature of the cation. The yields of the β -nitro ketones are higher with sunlamp irradiation than in the dark while 10–20 mol % of β -dinitrobenzene (β -DNB) or (β -Bu)2NO caused partial or complete inhibition of their formation. The yields of the ox-

iranes 3 and the α -hydroxy ketones 4 were not decreased in the dark or in the presence of the scavengers, their total yields increasing at the expense of 2. Complexation of the counterion of $Me_2C=NO_2^-M^+$ increased the yield of the β -nitro ketones. For example, the yield of 2a was increased from 41% to 77% by the presence of 18-crown-6 (K⁺, Me₂SO).

presence of 18-crown-6 (K^+ , Me_2SO).

The formation of ${\bf 2a}$ and ${\bf 2b}$ is thus formulated as an $S_{RN}1$ reaction (Scheme II).⁶ The formation of ${\bf 3}$ apparently involves

Scheme II

$$S_{RN}1$$
 mechanism (Y = O_2N , CN)
 $1^- \cdot \rightarrow p\text{-YC}_6H_4COCMe_2 \cdot + X^-$
 $p\text{-YC}_6H_4COCMe_2 \cdot + Me_2C=NO_2^- \rightarrow 2^-$
 $2^- \cdot + 1 \rightarrow 2 + 1^-$

nucleophilic attack by the carbon atom of $Me_2C=NO_2^-$ at the carbonyl carbon of 1 followed by S_Ni displacement of halogen by the carbonyl oxygen atom. The α -hydroxy ketones (4) may arise by O-attack of $Me_2C=NO_2^-$ at either the α or carbonyl carbon atoms of 1 to form nitronic esters, which decomposed during the reaction or upon workup.⁷

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Other anions that react with 1a by the $S_{RN}1$ process are $(EtO_2C)_2CR^-$ (R=H, Me), PhS⁻, and PhSO₂⁻. With the malonate anions in Me₂SO only the $S_{RN}1$ process was observed to yield 5a (49%) and 5b (62%), whose formation was completely inhibited by 10 mol % $(t\text{-Bu})_2NO$. With 1c alkylation was not observed with $(EtO_2C)_2CMe^-$ and 4c was the major product (62%). PhS⁻ and PhSO₂⁻ with 1a gave 5c and 5d by competing $S_{RN}1$ and ionic processes (the nitroxide retarded the photostimulated but not the dark reactions). Only ionic substitution was observed with 1c to give 5e and 5f. PhC=CLi or $n\text{-}C_3H_7C$ =CLi reacted (THF, -60 °C) with 1a or 1c to give the oxiranes 6 in nearly quantitative yields, whereas $(EtO)_2PO^-$ or $(EtO)_2PS^-$ reacted with 1a or 1c in a process unaffected by irradiation or free-radical scavengers to give the known enol phosphates⁸ and thiophosphates in a Perkow-type reaction. 9.10

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Photoinduced Reductive Elimination of Iron Atoms and Methane from CH_3FeH

Geoffrey A. Ozin* and John G. McCaffrey

Lash Miller Chemical Laboratories, University of Toronto Toronto, Ontario, Canada M5S 1A1

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In this communication we report on a novel photoreversible reductive elimination/oxidative addition reaction (1)

$$CH_3FeH \xrightarrow{420 \text{ nm}} Fe + CH_4$$
 (1)

in the Fe/CH₄ system operating on a single iron atom site at 10-12 K and induced by 420- and 300-nm narrow-band irradiation, respectively.

Narrow-band irradiation into the intense 300-nm $(3d^74p^1, ^5D_4)$ atomic resonance line of Fe atoms¹ in CH₄ under high dispersion $(1/10^4)$ conditions at 12 K caused rapid bleaching of all Fe atom bands with concomitant growth of a weak, broad absorption around 415-420 nm (Figure 1A-C). The corresponding infrared experiments (Figure 2) clearly demonstrated the production of the CH₃FeH insertion product absorbing strongly at 2921, 2888, 2869, 1650, 1148, 1145, 547, 544, 519, 300, 293 cm⁻¹, which aside from slightly better resolution (see later) is in accord with the original observations of Billups et al.² The presence of a *single* intense ν_{FeH} and *three* ν_{CH} modes in the

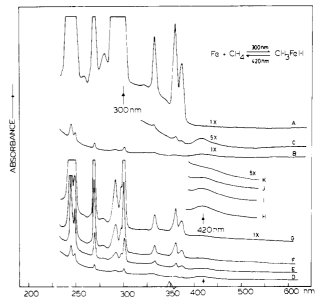


Figure 1. UV-visible spectra: (A) of Fe atoms isolated in solid CH₄ (1/10⁴) at 10-12 K; (B) following 30 min of 300-nm photolysis (Oriel 450-W Xe lamp, Oriel monochromator 20-nm band pass, 10-cm water cell, intensity at the sample 85 μ W cm⁻²): (C) 5× ordinate expansion of B in the 300-600-nm region; (D-G) samples similar to B at 0, 2, 7, and 12 min of 420-nm photolysis (intensity at the sample 175 μ W cm⁻²); (H-K) 5× ordinate expansion of D-G in the 370-540-nm region.

iron-hydrogen and -methyl stretching regions, respectively, together argue in favor of a CH₃FeH rather than a CH₂FeH₂ formulation for the Fe/CH₄ 300-nm photoproduct.

The thermal reactivity of Fe atoms with respect to CH₄ was also examined in the accessible cryogenic range 10-50 K. Up to the temperature that the methane actually sublimed away from the sample window (around 50 K), no new infrared or optical bands, ascribable to an Fe atom-CH₄ reaction product, were ever observed.

Let us now focus attention on the 415–420-nm photoreactivity of the CH₃FeH insertion product, generated from 300-nm excitation of Fe atoms under rigorously monatomic conditions (Figure 1B,C). The outcome of these irradiations were probed by UV-visible and infrared spectroscopy. Both experiments showed the monotonic bleaching of the CH₃FeH absorptions with 420-nm irradiation time (Figure 1D-K). Especially noteworthy was the concurrent and rapid generation of atomic iron, clearly seen by the steady growth of the atomic resonance lines in the optical spectrum (Figure 1D-G). Infrared bands characteristic of new photoproducts were not observed at any time during the photoannihilation of CH₃FeH. The 420-nm photoproduction of Fe atoms from CH₃FeH is found to be highly efficient, and essentially quantitative, in terms of the ability to fully recover the Fe atoms consumed in the original 300-nm photogeneration of CH₃FeH. Furthermore, the atomic Fe produced in this photofragmentation process is identical in form (Figure 1D-G) with the originally deposited Fe atoms and moreover can be readily back-converted to CH₃FeH by further 300-nm excitation. These observations confirm that the photogenerated Fe atoms are not trapped in a special matrix site and/or in a different electronic state following their ejection from CH₃FeH.

Some information pertaining to the geometry of the ground electronic state of CH₃FeH can be derived from infrared spectroscopy. For CH₃FeH generated by 300-nm photoexcitation of Fe atoms in solid CH₄, one finds (Figure 2B) that the vibrational modes of CH₃FeH observed in the range 4000-250 cm⁻¹ are comprised of two types, those arising from symmetrical stretching and bending motions of CH, FeH, and FeC bonds and those

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