Annulation With and Without CO-Insertion: Striking a Balance in Reactivity of Chromium Carbene Complexes with Chalcogen-Stabilized Iron Cluster Appendage

Pradeep Mathur, a,* Sundargopal Ghosh, a Amitabha Sarkar, b Arnold L. Rheingold c and Ilia A. Guzei c

aChemistry Department, Indian Institute of Technology, Powai, Bombay 400076, India
bDivision of Organic Chemistry (Synthesis), National Chemical Laboratory, Pune 411008, India
cDepartment of Chemistry, University of Delaware, Newark, DE 19716, USA

Abstract—Two new sets of products [(CO)6Fe2EE’{μ-μ(indanone)}] (2a: E, E’=Se; 2b: E=S, E’=Se) and [(CO)6Fe2EE’{μ-(2-ethoxy, α-naphthol)}] (3a: E, E’=Se; 3b: E=S, E’=Se; 3c: E=S, E’=Te) have been isolated from the thermolysis of the cluster-bound Fischer carbene complex, [(CO)6Fe2EE’{μ-C(Ph)–C–(OEt)}=Cr(CO)5}], 1a–c (1a: E, E’=Se; 1b: E=S, E’=Se; 1c: E=S, E’=Te) in THF. The products are formed by annulation pathway with and without the intermediacy of CO insertion step, from the same substrate. They have been characterized by IR and 1H, 13C, 77Se and 125Te NMR spectroscopy and the structural types were unequivocally established by crystallographic analysis of compounds 2a and 3c.

Introduction

Some of the earliest information on the reactivity of the chalcogen-stabilized iron carbonyl clusters, [Fe2(CO)6(μ-EE’)] (where E, E’=S, Se, Te) originated in studies of the reduction of carbon–carbon triple bond in terminal alkyynes by the reactive chalcogen–chalcogen bonds present in [Fe2(CO)6(μ-EE’)] and these iron carbonyl clusters of the same structure with compositions differing only in the identity of the main group element are very rare.1 The chalcogen-stabilized diiron compounds [Fe2(CO)6(μ-EE’)] add readily to the triple bond of the carbene complexes [(CO)6M=C(OEt)(C=CPh)] (M=Cr, W) to form the adducts, [(CO)6Fe2EE’{μ-PhC=CC(OEt)}M(CO)5].2 In those cases where E≠E’, such additions are highly regio-selective.2a The Fischer carbene moiety in these adducts is a potential organic functional group, as demonstrated by its facile transformation to an amino carbene group, an ester, an orthoester 3b or an enol ether derivative.3

The competitive formation of benzannulation and cyclopentannulation products from the same reaction is well preceded and can be tuned e.g. by N-acylation.4 Our preliminary investigations indicate3 that product formation upon thermolysis is strongly influenced by the nature of chalcogen atoms present in the trimetallic adducts [(CO)6Fe2EE’{μ-C(Ph)=CC(OEt)}Cr(CO)5] and such studies, including those aimed towards cyclopentannulation continue to be of significant interest as they show some exciting physical properties e.g. non linear optical activity, NLO.5 Thus, when thermolysis of [(CO)6Fe2STe{μ-PhC=CC(OEt)}Cr(CO)5] was carried out, no cyclopentannulation was observed; two compounds were isolated and identified as [(CO)6Fe2{μ-SC(Ph)=C(Te)=C(H)(OEt)}]2 and [(CO)6Fe2{μ-SC(H)(Ph)=C(Te)=C(H)(OEt)}].2b Since the amount of isolated products did not provide a satisfactory material balance, the reaction mixture was closely examined for identification of more polar components and here we report the results of an investigation of the thermolysis of [(CO)6Fe2EE’{μ-C(Ph)=CC(OEt)}=Cr(CO)5}], 1a–c (1a: E, E’=Se; 1b: E=S, E’=Se; 1c: E=S, E’=Te) with a focus on the isolation of more polar compounds.

Results and Discussion

Thermolysis of complexes [(CO)6Fe2EE’{μ-C(Ph)=CC(OEt)}=Cr(CO)5}], 1a–c (1a: E, E’=Se; 1b: E=S, E’=Se; 1c: E=S, E’=Te) was carried out under previously reported condition (THF reflux, 3 h). Chromatographic work-up using a polar solvent system of CH2Cl2/hexane (40:60 v/v) for elution afforded two new complexes: [(CO)6Fe2EE’{μ-μ(indanone)}] (2a: E, E’=Se; 2b: E=S, E’=Se) and [(CO)6Fe2EE’{μ-(2-ethoxy, α-naphthol)}] (3a: E, E’=Se; 2b: E=S, E’=Se).
These products were characterized by their spectral characteristics and crystal structure determination of representative complexes 2a and 3c. When the thermolysis reaction of complexes 1a−c was carried out in toluene at 85°C, formation of α-naphthol derivatives, 3a−c predominates. The thermolysis of 1a gives better yield of the α-naphthol derivative, 3a (70%) than the thermolysis of 1b and 1c (3b: 34%; 3c: 38%) (Scheme 1).

The IR spectra of compounds 2a and 2b showed signals corresponding to the presence of the terminally bonded metal carbonyl groups and a band at 1727 cm⁻¹ due to a ketonic carbonyl function. The high value of the ketone carbonyl absorption suggested the presence of a five-membered ketone unit in this molecule. The 1H NMR spectra showed typical sets of doublets associated with Se satellites, i.e. the carbon bearing these protons are attached to a selenium atom. The 13C NMR spectra displayed a downfield signal for the Fe−CO groups at 209 ppm. Two peaks were observed for the saturated carbon atoms in complexes 2a and 2b; the deshielded signals are assigned to the carbons adjacent to the ketone whose presence was confirmed by the signal at δ 199.3 (2a) and 198.1 (2b) ppm. For complex 2a, the 77Se NMR spectrum showed two signals. The downfield signal is a doublet with J_{Se−H} = 25.9 Hz. One doublet signal was observed with J_{Se−H} of 18.3 Hz for complex 2b.

Yellow crystals of 2a were grown from hexane solvent at −4°C and a single crystal X-ray diffraction analysis was carried out. Its molecular structure is shown in Fig. 1.

The structure can best be described as consisting of an open Fe₂(CO)₆Se₂ unit with an indanone moiety formally inserted between the two Se atoms. The distance between the two carbon atoms [C(7) and C(15)] which are attached to the selenium atoms, is slightly longer (1.502(6) Å) than the analogous C−C bond distance of 1.48(1) Å, observed in a previously reported double butterfly cluster [{(CO)₆Fe₂(μ−Se)₂}₂C(Ph)−C(H)] and the analogous C−C bond distance of 1.47(2) Å in the enol ether derivative [{(CO)₆Fe₂(μ−SC(Ph)–C(Se))}]. The two C−Se bond distances in compound 2a are nearly equal.

The IR spectra of compound 3a−c showed typical peaks for terminally bonded metal-CO groups. The 1H NMR spectra of compounds 3a−c featured a sharp singlet in the region δ 6.1−6.2 ppm, which was exchangeable with D₂O and consequently was assigned to the phenolic hydroxyl group. The 13C NMR spectra displayed six aromatic quaternary carbon peaks and four aromatic carbon atoms bearing a hydrogen each between δ 113.2 and 143.2 ppm.

Figure 1. [(CO)₆Fe₂Se₂[μ-indanone]] (2a).
The $^{77}$Se NMR spectrum showed two expected singlets for $3a$ and one singlet for $3b$. Similarly, the $^{125}$Te NMR spectrum of compound $3c$ featured a single peak. An unambiguous assignment of these two isostructural complexes was made on the basis of crystal structure determination of complex $3c$.

Red crystals of $3c$ were grown from hexane/CH$_2$Cl$_2$ at 0°C and a single crystal X-ray diffraction analysis was carried out. Its molecular structure is shown in Fig. 2. A naphthol ring attached to the chalcogen atoms of Fe$_2$STe butterfly core was observed. The olefinic bond distance of 1.346(11) Å in $3c$ [C(7) and C(8)] attached with the chalcogen atoms, is comparable to the olefinic bond distance of 1.36 Å in the uncoordinated naphthol ring. No appreciable deviation from planarity was observed with its S and Te substituents (dihedral angles: Te–C(7)–C(8)–C(9)=8.7°, S–C(8)–C(7)–C(16)=2.2° and Te–C(7)–C(16)–O(8)=6.3°). This angularly fused polycyclic structure can be viewed as an unusual, organometallic analog of phenanthrene with a sulfur atom in the ‘bay’ region. The position of the sulfur atom also suggests that, with appropriate metal ion, it can lead to an ortho-metallated complex involving the C(10) center.$^7$

Formation of these products, $2a$–$b$ and $3a$–$c$ can be explained by involving steps well-precedented in metal-carbene chemistry$^{10}$ where the cis-olefinic bond makes the annulation reactions readily attainable. Two product sets, $2$ and $3$ probably result from two alternative pathways—one involves CO insertion step (leading to $3a$–$c$) while the other does not (leading to $2a$ and $2b$). It is interesting to note that the same substrate can react via both these pathways at the same time albeit with different rates. This suggests that the electron density on the chromium metal in complexes $1a$–$c$ lies somewhere in between usual alkoxy and amino carbene complexes of chromium.$^{11}$ It is indeed a rare instance that the same substrate can react via both these pathways at the same time albeit with different rates. This suggests that the electron density on the chromium metal in complexes $1a$–$c$ lies somewhere in between usual alkoxy and amino carbene complexes of chromium.$^{11}$ It is indeed a rare instance that the same carbene complex gives rise to two different products in the same reaction.$^{12}$

Moreover, the transformation from $1a$–$c$ to $3a$–$c$ belongs to a unique class of reactions where an annulation occurs from a cis-cinnamyl chromium carbene derivative without the requirement of an added alkyne, and the product is a vicinally dioxygenated product. Examples of such transformation are rare and the only example known is a photo-induced one.$^{13}$ It is significant that the irradiation of substrate $1a$ with 400 W medium pressure mercury lamp with a Pyrex filter for 6 h failed to produce a noticeable change in the starting complex, though the thermal reaction is comparatively facile, and in this respect, curiously enough, our reaction differs from that reported by Merlic.$^{13}$

**Experimental**

**General**

All reactions and other manipulations were carried out under an argon or nitrogen atmosphere, using standard Schlenk techniques. Solvents were deoxygenated immediately prior to use. Reactions were monitored by FT-IR spectroscopy and thin-layer chromatography. Infrared spectra were recorded on a Nicolet-Impact 400 FT-IR spectrometer as n-hexane solutions in sodium chloride cell at 0.1 mm path length.

Photolysis reaction was done in 400 W medium pressure mercury vapor lamp using Pyrex filter. Elemental analyses were performed using a Carlo Erba 1106 automatic analyzer. $^1$H, $^{13}$C, $^{77}$Se and $^{125}$Te NMR spectra were recorded on a Varian VXR 300S spectrometer in CDCl$_3$ at 25°C. The operating frequency for $^{77}$Se NMR was 57.23 MHz with a pulse width of 15 μs and a delay of 1.0 s and operating frequency for $^{125}$Te was 94.70 MHz with pulse width of 9.5 μs and a delay of 1 s. $^{77}$Se NMR spectra were referenced to Me$_2$Se (δ=0 ppm) and $^{125}$Te NMR spectra were referenced to Me$_2$Te (δ=0 ppm).

Chromium hexacarbonyl, phenylacetylene and THF-d8 were purchased from Aldrich Chemical Co., and these
were used without further purification. The homo- and mixed-chalcogenide iron carbonyl clusters Fe₂(µ-Se₂)(CO)₆, 14 Fe₂(µ-SeSe)(CO)₆,15 and Fe₂(µ-TeTe)(CO)₆,16 α, β unsaturated mixed chalcogenide carbene complexes [(CO)₂Fe(µ-CPh)(=C=O)(Et)=Cr(CO)₅], (1a: E, 1b: E=Se; 1c: E=S, E=Te) and the alkynyl Fischer carbene complexes [(CO)₂M(=C(OEt)(C=Ph)]17 (M=Cr, W) were prepared as previously reported.

**General procedure for thermolysis of [(CO)ₓFe₂(CO)ₓ][{µ-C(PH)=CC(OE)(–OEt)=Cr(CO)₅}] (1a: E, E=Se; 1b: E=S, E=Se; 1c: E=S, E=Te)

A solution of [(CO)ₓFe₂(CO)ₓ][{µ-C(PH)=CC(OE)(–OEt)=Cr(CO)₅}], (1a: E, E=Se; 1b: E=S, E=Se; 1c: E=S, E=Te) in THF (10 mL) was heated at 85°C for 3 h. The reaction was monitored by TLC, and stopped when all the starting material was consumed. The solution was filtered through Celite to remove the insoluble materials. After removal of solvent from the filtrate, the residue was subjected to chromatographic work-up using silica gel TLC plates. Elution with hexane/CH₂Cl₂ (40:60) mixture afforded two polar complexes, in order of polarity: yellow [(CO)ₓFe₂(µ-C(Ph)=CC(OE)(–OEt)=Cr(CO)₅)] (2a: E, E=Se; 2b: E=S, E=Se) and red [(CO)ₓFe₂(µ-C(2-ethoxy, α-naphthyl)) (3a: E, E=Se; 3b: E=S, E=Se; 3c: E=S, E=Te).

2a: yield 15%. IR: 2072 (s), 2036 (vs), 2003 (s), 1998 (s), 1984 (w), 1728 (m); ¹H NMR: δ 4.06 (1H, d, Jₛₑ₋₁₇=6.2 Hz, Jₛₑ₋₁₇=19.8 Hz, (COCH₃), 4.95 (1H, d, Jₛₑ₋₁₇=6.1 Hz, Jₛₑ₋₁₇=25.8 Hz, CCH), 7.4–7.7 (4H, m, C₆H₅), 13C NMR: δ 48.3 (SeCH(CH₃), Jₛₑ₋₁₇=154.9 Hz), 54.5 (SeCH(CH₃), Jₛₑ₋₁₇=150.2 Hz), 124.4, 126.4, 129.7, 136.6 (4CH, in indanone ring), 133.4, 152.8 (quat C in indanone ring), 199 (CO, in indanone ring), 208.6 (Fe(CO)₅); ³⁷/³⁹Se NMR: δ 618 (dd, Jₛₑ₋₁₇=25.9 and 2.3 Hz, SeC(H)C), 636 (d, Jₛₑ₋₁₇=19.1 Hz, SeC(H)(CO); mp: 162–164°C Anal. calcd (Found) for Fe₂Se₂C₁₅O₇H₁₀: C, 31.7 (31.9), H, 1.05 (1.09).

3a: yield 18%. IR: 2071 (s), 2035 (vs), 2001 (s), 1985 (w), 1951 (w); ¹H NMR: δ 1.55 (3H, t, J₇=6.9 Hz, CH₃), 4.12 (2H, q, J₇=1.1 Hz, OCH₂), 6.09 (1H, s, –OH), 7.79–7.98 (2H, two sets of doublets, naphthol ring), 7.44–7.32 (3H, m, naphthol ring); ¹³C NMR: δ 15.5 (Jₛₑ₋₁₇=127.4 Hz, CH₃), 71.6 (Jₛₑ₋₁₇=140 Hz, OCH₂), 122.6, 125.9, 126.9, 127.2 (CH carbons in naphthol ring), 113.2, 123.6, 130.3, 135.8, 140.4, 143.1 (quat carbons in naphthol ring), 208.7 (Fe(CO)₅); ⁷⁷/⁷⁹Se NMR: δ 353 and 358 ppm; mp: 126–128°C Anal. calcd (Found) for Fe₂Se₂C₁₅O₇H₁₀: C, 34.6 (34.9), H, 1.60 (1.79).

3b: yield 11%. IR: 2073 (s), 2038 (vs), 2005 (s), 1987 (s), 1727 (m); ¹H NMR: δ 4.01 (1H, d, J₇=6 Hz, Jₛₑ₋₁₇=18.4 Hz, SeCH₂), 4.63 (1H, d, J₇=6.1 Hz, SCH), 7.44–7.75 (4H, m, C₆H₅), 13C NMR: δ 52.6 (Jₛₑ₋₁₇=152.9 Hz, SeCH₂(Ph)), 57.2 (Jₛₑ₋₁₇=149.1 Hz, SCH(Ph)), 124.4, 126.5, 130.1, 136.6 (4CH, in indanone ring), 134.4, 152.1 (quat C in indanone ring), 198.2 (CO in indanone ring), 208.2 (Fe(CO)₅); ⁷⁷/⁷⁹Se NMR: δ 712 (d, Jₛₑ₋₁₇=18.3 Hz; mp: 156–158°C Anal. calcd (Found) for Fe₂Se₂C₁₅O₇H₁₀: C, 34.5 (34.9), H, 1.15 (1.29).

3c: yield 23%. IR: 2073 (s), 2039 (vs), 2001 (s), 1955 (w); ¹H NMR: δ 1.43 (3H, t, J₇=6.9 Hz, CH₃), 4.11 (2H, q, J₇=7.1 Hz, OCH₂), 6.18 (1H, s, –OH), 7.29–7.42 (5H, m, C₆H₅), 7.87–7.95 (2H, two sets of doublets, naphthol ring); ¹³C NMR: δ 15.6 (Jₛₑ₋₁₇=126.2 Hz, CH₃), 71.6 (Jₛₑ₋₁₇=141.3 Hz, OCH₂), 122.5, 125.1, 126.1, 127.1 (CH carbons in naphthol ring), 123.6, 130.5, 134.2, 138.1, 139.9, 142.1 (quat carbon atoms in naphthol ring), 208.1 (Fe(CO)₅); ⁷⁷/⁷⁹Se NMR: δ 458 ppm; mp 122–124°C. Anal. calcd (Found) for Fe₂Se₂C₁₅O₇H₁₀: C, 37.5 (37.7), H, 1.73 (1.86).

**General procedure for thermolysis of [(CO)ₓFe₂(CO)ₓ][{µ-C(PH)=CC(OE)(–OEt)=Cr(CO)₅}] (1a–c) in toluene

A solution of [(CO)ₓFe₂(CO)ₓ][{µ-C(PH)=CC(OE)(–OEt)=Cr(CO)₅}], (1a: E, E=Se; 1b: E=S, E=Se; 1c: E=S, E=Te) in toluene (15 mL) was refluxed for 3 h and the solution was filtered through Celite to remove the insoluble materials. After removal of solvent from the filtrate, the residue was subjected to chromatographic work-up using silica gel TLC plates. Elution with hexane/CH₂Cl₂ (40:60) mixture afforded two polar complexes, in order of polarity: yellow [(CO)ₓFe₂(µ-C(Ph)=C=O)(Et)=Cr(CO)₅)] (2a: E, E=Se; 2b: E=S, E=Se) and red [(CO)ₓFe₂(µ-C(2-ethoxy, α-naphthyl)] (3a: E, E=Se; 3b: E=S, E=Se; 3c: E=S, E=Te).
Photolysis of \([\text{CO})_6\text{Fe}_2\text{Se}_2\{\mu-\text{C(Ph)}=\text{CC(OEt)}=\text{Cr(CO)}_3\}\] (1a) in THF-\(d_8\)

A solution of 1a (0.005 g, 0.006 mmol) in THF-\(d_8\) (0.7 mL), was photolysed in a NMR tube using 400 W medium pressure mercury lamp for 6 h. After photolysis, \(^1\text{H} \text{NMR spectrum of the reaction mixture was recorded.}

No signal other than that of 1a was observed.

Crystal structure determination of 2a and 3c

Red crystals of 2a and 3c were selected and mounted with epoxy cement to glass fibres. Single crystal X-ray data were collected on Siemens P4 diffractometer by using MoK\(\alpha\) radiation. The unit-cell parameters were obtained by the least-squares refinement of the angular settings of 24 reflections (20° ≤ 2θ ≤ 25°). Pertinent crystallographic data of 2a and 3c are summarized in Table 1. The systematic absences in the diffraction data for 2a, 3c are uniquely consistent for the reported space groups. The structures were solved using direct methods, completed by subsequent difference Fourier syntheses and refined by full-matrix least-squares procedures. Empirical absorption corrections for 2a was applied by using program \textsc{difabs}.\(^{18}\) All non-hydrogen atoms were refined with anisotropic displacement coefficients and hydrogen atoms were treated as idealized contributions.

All software and sources of the scattering factors are contained in the \textsc{shelxtl} (5.3) program library (G. Sheldrick, Siemens XRD, Madison, WI). Selected bond lengths and bond angles for 2a and 3c are listed in Tables 2 and 3.

Conclusion

In Fischer carbene chemistry, the reactivity of the complexes is dominated by, among other things, the metal atom and its electron-richness or deficiency. While aryl or vinyl alkoxy carbene complexes of chromium undergo Do¨tz reaction (a benzannulation process that involves a CO-insertion step) under thermal conditions, CO-insertion is not observed for corresponding complexes of tungsten or amino carbene analogs of chromium. In this paper, we have reported an interesting variant where the vinyl group of a chromium carbene complex is supported on a chalcogen-stabilized diiron cluster, and such substrate exhibits simultaneous operation of both the pathways—with \textit{and} without the CO-insertion step during the same reaction. It is not clear at this point whether the electronic influence is restricted to the contribution by the chalcogen atoms alone, and this point needs further investigation.

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\textbf{References}


\begin{table}
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\begin{tabular}{|c|c|c|}
\hline
& & \\
$\text{Fe}$(1)&$\text{Fe}$(2)&$\text{Te}$
\hline
Fe(1)&2.550(2)&Te(1)&C(7)
\hline
Fe(1)&2.518(2)&S&C(8)
\hline
Fe(2)&2.243(2)&C(11)&C(12)
\hline
Fe(1)&2.267(3)&C(9)&C(14)
\hline
Fe(2)&2.550(2)&C(7)&C(8)
\hline
Fe(2)&55.13(7)&Fe(2)&C(2)&C(7)
\hline
Fe(2)&56.03(7)&C(16)&C(7)&Te(1)
\hline
Fe(2)&68.84(8)&Fe(1)&C(11)&C(7)
\hline
Fe(2)&60.41(5)&C(8)&C(7)&Te(1)
\hline
Fe(2)&60.19(5)&C(9)&C(8)&S
\hline
Fe(2)&60.40(5)&C(7)&C(8)&S
\hline
Fe(2)&82.34(7)&C(8)&S&Fe(1)
\hline
\end{tabular}
\caption{Selected bond distances [Å] and bond angles [°] for 2a}
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