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New complexes with M-Si-O or M-Si-S linkages (M = Fe or Co).

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Abstract

 $Ph_2XSiFe(CO)_2Cp [X = p-tolylS (1a), MeO (1b)]$ and $Ph[2-MeOC_6H_4]XSiFe(CO)_2Cp [X = Cl (2a), OMe (2b)]$ have been fully characterised, including X-ray crystal structure determinations for 1a, 1b and 2a. None of the examples showed any tendency for migration of the X groups from silicon to iron, with elimination of silylene. However very ready loss of the X groups was seen in the electrospray mass spectra, suggesting formation of the cationic silylene-iron complex ions is favoured. This was especially so for 2a and 2b, where intramolecular stabilisation of the silicon centre from the 2-OMe group is possible. The stable siloxane $O[SiPh_2{Co(CO)_4}]_2$ was also characterised; the X-ray crystal structure analysis shows a Si-O-Si bond angle of 153°.

1. Introduction

The 1,2-migration of a group X from silicon to a transition metal in compounds containing M-SiR₂X units has been frequently invoked to explain rearrangement processes,

especially those where metal-silylenes are probable intermediates [1-3]. Only recently however has this been directly observed, by Mitchell and Tilley in the reversible interconversion of $(Et_3P)_2Pt(H)Si(SBu^t)_2OTf$ and $[(Et_3P)_2Pt(\mu-SBu^t)Si(H)(SBu^t)]^+$ via H-migration [4]. More commonly the systems in which 1,2-migrations are indirectly most strongly implicated are those involving disilanes, where migration of the β -silyl group to the metal generates silyl/silylene intermediates [5]. Examples of this type are known for many transition metals [1].

So far there appear to be no definite examples in which a -OR or -SR group transfers from silicon to a transition metal. Migration involving a Si-O bond may be implicated in the metal-catalysed disproportionation of $O(SiMe_2H)_2$ into Me_2SiH_2 and polysiloxanes, though other mechanisms can be proposed [6]. A related example is the reaction of $O(SiH_3)_2$ with $Co_2(CO)_8$ to give products such as $Si[Co_2(CO)_7]_2$ without Si-O bonds [7]. Transfer of -SR is perhaps even more likely, given the 1,2-migration of an H-group described above, since Si-H and Si-S bonds have similar strengths (e.g. Si-H 320-340 kJ mol⁻¹, Si-S 314 kJ mol⁻¹ [8]).

There are now several examples of species with M-Si-O linkages described in the literature, with varying properties. Greene and Curtis tentatively characterised the siloxane $O[SiMe_2\{Co(CO)_4\}]_2$, but found it to be extremely unstable [9]. In contrast, species such as $(MeO)_3SiCo(CO)_4$ are apparently readily handled [10]. Braunstein [11] has developed extensive chemistry of bi-metallic compounds based on $(MeO)_3Si$ groups acting as bridging ligands. More recently, stable metallosilanol and metallosiloxane compounds have been reported, including derivatives $(Me_5C_5)(PMe_3)(OC)_2MSiR_2(OH)$ (M = Cr, Mo, W) from Malisch's group [12], and osmium examples from Roper's laboratory [13]. Both mono- and octa-Co(CO)₄ derivatives of octasilasesquioxane have been characterised [14], following earlier work with smaller cyclosiloxanes [15].

This present paper describes some new complexes with L_nM -Si-OR or L_nM -Si-SR bonds. The fragment L_nM chosen was mainly $Cp(CO)_2Fe$ since this is the best studied for silvl group migration. The supporting groups on the silicon were Ph since these stabilise silvlene intermediates better than Me. One example incorporating Co(CO)₄ was also synthesised since this group appears to encourage silvl migration under very mild conditions [3].

2. Experimental Section

2.1 General

All manipulations were carried out in an oxygen-free N₂ atmosphere with carefully dried solvents in standard Schlenk equipment. Ph(2-MeOC₆H₄)SiCl₂ was prepared by coupling ortholithiated anisole with Ph₂SiCl₂, and (Ph₂HSi)₂O by hydrolysis of Ph₂HSiCl. Co₂(CO)₈ was freshly sublimed before use while [Fe(CO)₂Cp]₂ was used as received. "TolS" refers specifically to p-tolyl-thio group.

2.2 Instrumentation.

Infrared spectra were recorded on a Digilab FTS-40 FTIR spectrophotometer. NMR spectroscopy was performed using a Bruker AC300P Multinuclear FT spectrometer. Routine electrospray mass spectra (ESMS) were obtained on a VG Platform II spectrometer operating under standard conditions. EDESI mass spectra were collected using a Micromass Quattro LC instrument, in positive-ion mode. Samples were introduced directly to the source at 4 μ L min⁻¹ via a syringe pump as acetonitrile/methanol solutions. Data collection was carried out in continuum mode. The cone voltage was initially set at 0 V and was increased by increments of 1 V after every scan up to a maximum of 200 V.

Elemental analysis was performed by the Campbell Microanalytical Laboratory, University of Otago. Melting points were measured on a Reichart Thermopan melting point apparatus and are uncorrected.

2.3 Reactions.

2.3.1 Preparation of Ph₂ClSiFe(CO)₂Cp.

This known compound was prepared in 60% yield from Na[Fe(CO)₂Cp] and Ph₂SiCl₂ in thf. Mp 88-89°C (Lit. 89-94°C [16] or 95-96°C [17]), IR: v CO (petroleum spirits) 2015 (s), 1965 (s) cm⁻¹; ¹H NMR: δ 7.68, 7.35 (m, 10H, C₆H₅), δ 4.80 (s, 5H, C₅H₅); ¹³C NMR: δ 214.5 (CO), δ 142.3 (*ipso*), 133.6 (*ortho*), 129.1 (*para*), 127.8 (*meta*) (C₆H₅), δ 85.5 (C₅H₅); ESMS: (MeCN/H₂O), m/z 418 (100%) [M-Cl+MeCN+H₂O]⁺; 400 (45%) [M-Cl+MeCN]⁺; 377 (35%) [M-Cl+H₂O]⁺; 359 (40%) [M-Cl]⁺.

2.3.2 Preparation of Ph₂(TolS)SiFe(CO)₂Cp (1a).

To a stirred THF solution (30 mL) of Ph₂ClSiFe(CO)₂Cp (0.683 g, 1.73 mmol) was added Na[STol] (0.253 g, 1.73 mmol). After 2 days stirring the IR spectrum of the reaction solution

showed complete conversion to a new species. The solvent was evaporated and replaced with ether (30 ml). The solution was filtered, petroleum spirits (10 ml) was added and the mixture was concentrated to ca 5 mL, and stored at -20°C. Yellow crystals of Ph₂(TolS)SiFe(CO)₂Cp were formed (0.400 g, 48%). Mp 133°C, found C: 63.59, H: 4.61 %, C₂₆H₂₂O₂SiSFe requires C: 64.73, H: 4.60 %; HRMS: found 482.046, required 482.0459, IR: v CO (THF) 2005 (s), 1953 (s) cm⁻¹; ¹H NMR: δ 7.67, 7.30 (m, 10H, C₆H₅), δ 6.99 (d, *J* = 7.8 Hz, 2H, C₆H₄), δ 6.79 (d, *J* = 8.1 Hz, 2H, C₆H₄), δ 4.63 (s, 5H, C₅H₅), δ 2.19 (s, 3H, CH₃); ¹³C NMR: δ 214.3 (CO), 141.4 - 127.6 (m, *C*₆H₅, *C*₆H₄), δ 85.0 (*C*₅H₅), δ 21.0 (*C*H₃); FABMS: m/z: 482 (7%) [M+H]⁺, 454 (68%) [M+H-CO]⁺, 426 (100%) [M+H-2CO]⁺, 359 (74%) [M-STol]⁺, 304 (33%) [M-STol-2CO]⁺. Attempted purification by chromatography on silica led to extensive decomposition, the only product isolated being [Fe₂(µ-STol)₂(CO)₂Cp₂]⁺ (identified spectroscopically: ESMS m/z 544 [M⁺]; IR v(CO) (thf) 1972 cm⁻¹; ESR: g_{iso} 2.004, Δ p-p 10G, cf ref 18) with an unidentified anion.

2.3.3 Preparation of Ph₂(MeO)SiFe(CO)₂Cp (1b).

A THF solution (10 ml) of Ph₂ClSiFe(CO)₂Cp (0.501 g, 1.27 mmol) was treated with a methanolic solution of NaOMe (0.82 ml, containing 1.25 mmol NaOMe). The reaction was left to stir overnight. The solvent was replaced with ether, the solution was filtered and left to crystallise at 4°C. Small, thin yellow crystals of Ph₂(MeO)SiFe(CO)₂Cp formed (0.399 g, 81%). Mp 99°C ; found C: 61.57, H: 4.71 %, C₂₀H₁₈O₂SiFe requires C: 61.55, H: 4.65 %; IR: v CO (THF) 1998 (s), 1943 (s) cm⁻¹; ¹H NMR: δ 7.60, 7.34 (m, 5H, C₆H₅), δ 4.65 (s, 5H, C₅H₅), δ 3.48 (s, 3H, OCH₃); ¹³C NMR: δ 214.9 (CO), δ 142.6 (*ipso*), 133.9 (*ortho*), 128.7 (*para*), 127.7 (*meta*) (SiC₆H₅), δ 84.5 (C₅H₅), δ 52.2 (OCH₃); ²⁹Si NMR: δ 57.2, ESMS (MeCN/H₂O): Cone 20 V: m/z 408 (100%) [M + NH₄]⁺, 391 (40%) [M + H]⁺; Cone 40 V: m/z 400 [M-OMe+MeCN]⁺. More detailed ESMS information about fragmentation processes for this compound is presented in the Results and Discussion section below.

2.3.4 Preparation of $Ph(2-MeOC_6H_4)ClSiFe(CO)_2Cp$ (2a).

 $[Fe(CO)_2Cp]_2$ (0.71 g, 2.00 mmol) in THF (20 ml) was reduced with sodium amalgam (1%) to Na[Fe(CO)_2Cp]. The resulting solution was transferred by syringe into a Schlenk flask at 0°C containing Ph(2-MeOC_6H_4)SiCl_2 (1.3 g, 4.6 mmol) in ether (10 ml). The stirred solution was allowed to return to room temperature and left overnight. The solvent was removed under

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vacuum and replaced with ether (30 ml). The solution was passed through a filter stick to remove NaCl and the solvent removed once more. Chromatography (silica, CH₂Cl₂/petroleum spirit 1:5) provided a bright yellow band which was collected and recrystallised from ether/petroleum spirit to provide pale yellow crystals of Ph(2-MeOC₆H₄)ClSiFe(CO)₂Cp. (0.54 g, 32%). Mp 108-110°C; found C: 56.37, H: 4.02 %; C₂₀H₁₇O₃SiClFe requires C: 56.56, H: 4.03 %; IR: v CO (petroleum spirit) 2016 (s), 1967 (s) cm⁻¹. ¹H NMR: δ 7.9 - 6.8 (m, 9H, C₆H₅, C₆H₄), δ 4.72 (s, 5H, C₅H₅), δ 3.68 (s, 3H, OCH₃); ¹³C NMR: δ 214.0, 213.9 (2CO), δ 142.5 - 109.9 (m, C₆H₅ and C₆H₄), δ 85.5 (C₅H₅), δ 60.3 (CH₃); ESMS: (MeCN/H₂O) m/z 429 [M - Cl + OH + Na]⁺, 425, [M - Cl + 2H₂O], 407 [M - Cl + H₂O]⁺

2.3.5. Preparation of Ph[2-(MeO)C₆H₄](MeO)SiFe(CO)₂Cp (**2b**)

A THF solution (10 ml) of Ph[2-(MeO)C₆H₅]ClSiFe(CO)₂Cp (0.174 g, 0.410 mmol) was treated with a methanolic solution of NaOMe (3.6 ml, containing 0.45 mmol NaOMe). After 14 h the solvent was evaporated and the residue was extracted with ether. After filtration the solution was stored at -25°C to give ragged yellow crystals of Ph[2-(MeO)C₆H₄](MeO)SiFe(CO)₂Cp (0.077 g, 45%). Mp 176-179°C; IR: v CO (THF) 2001 (s), 1946 (s) cm⁻¹; ¹H NMR: δ 7.58, 7.27 (m, 5H, C₆H₅), δ 6.9 (m, 4H, C₆H₄), δ 4.65 (s, 5H, C₅H₅), δ 3.66 (s, 3H, OCH₃), δ 3.44 (s, 3H, SiOCH₃); ESMS: (MeCN/H₂O); m/z 421 [M + H]⁺.

2.3.6 Reaction of $(Ph_2HSi)_2O$ with $Co_2(CO)_8$ (3).

To a Schlenk flask was added (PhH₂Si)₂O (0.562 g, 1.47 mmol), Co₂(CO)₈ (0.500 g, 1.46 mmol) and petroleum spirits (25 mL). The mixture was stirred overnight at room temperature to produce a pale brown solution with some white precipitate. The solvent was removed under vacuum, and the residue was washed with small quantities of petroleum spirit to remove unreacted Co₂(CO)₈. The crude product was extracted from small amounts of Co₄(CO)₁₂ with diethyl ether, and this solution was evaporated slowly to yield large, colourless crystals of O[SiPh₂{Co(CO)₄}]₂ (0.874 g, 83%). IR v(CO): (petroleum spirits, cm⁻¹) 2097(m), 2096 (m), 2036(m), 2035 (m), 2014 (vs), 2005(s), 1998(s). The compound was further characterised by an X-ray crystal structure determination.

2.4 X-ray crystallography.

For compounds **1a** and **2a** unit cell parameters and intensity data were collected using a Siemens SMART CCD diffractometer, using standard collection procedures, with monochromatic Mo-Ka X-rays (0.71073 Å). Corrections for absorption and other effects were carried out with SADABS [19]. For **1b** and **3** data were collected on a Nicolet R3 four-circle diffractometer and corrected for absorption by a ϕ scan method. All other calculations used the SHELX97 programs [20]. The structures were solved by direct methods, and developed routinely with refinement based on F². All non-hydrogen atoms were assigned anisotropic temperature factors, and hydrogen atoms were included in calculated positions. Selected bond parameters are in the captions to Figures 1, 3-5.

2.4.1 Crystal data for Ph₂(TolS)SiFe(CO)₂Cp (1a)

 $C_{26}H_{22}FeO_2SSi$, $M_r 482.44$, triclinic, $P\bar{1}$, a = 8.5754(2), b = 9.9558(2), c = 14.7730(3) Å, $\alpha = 81.066(1)$, $\beta = 75.044(1)$, $\gamma = 68.367(1)^\circ$, V = 1130.15(4) Å³, $D_{calc} = 1.418$ g cm⁻³, Z = 2, F(000) = 500, μ (Mo-K α) 0.833 mm⁻¹, T_{max} 1.000, T_{min} 0.7217, crystal size 0.44 x 0.37 x 0.09 mm³. T 148 K.

A total of 13606 reflections, 3751 unique ($R_{int} 0.0378$) was collected $2^{\circ} < \theta < 25^{\circ}$. Final R1 0.0575 (data with I > 2 σ (I)), 0.0606 (all data), wR2 0.1722, GoF 1.030, final $\Delta e +1.20/-0.51$. 2.4.2 Crystal data for Ph₂(MeO)SiFe(CO)₂Cp (**1b**)

 $C_{20}H_{18}FeO_3Si$, M_r 390.28, monoclinic, C2/c, a = 31.67(3), b = 7.609(4), c = 17.35(2) Å, β = 122.85(5)°, V = 3512(5) Å³, D_{calc} = 1.476 g cm⁻³, Z = 8, F(000) = 1616, μ (Mo-K α) 0.91 mm⁻¹, T_{max} 0.86, T_{min} 0.71, crystal size 0.82 x 0.53 x 0.12 mm³. T 141 K.

A total of 2357 reflections, 2097 unique (R_{int} 0.0497) was collected $2^{\circ} < \theta < 22.5^{\circ}$. Final R₁ 0.0532 (data with I > 2 σ (I)), wR₂ 0.1256, GoF 1.036, final Δe +0.96/-0.37.

2.4.3 Crystal data for $Ph(2-MeOC_6H_4)ClSiFe(CO)_2Cp$ (2a)

 $C_{20}H_{17}ClFeO_3Si$, M_r 424.73, Monoclinic, P_{21}/n , a = 9.0901(1), b = 14.6719(2), c = 15.0469(2)

Å, $\beta = 107.38(1)$, V = 1915.20(4) Å³, D_{calc} = 1.473 g cm⁻³, Z = 4, F(000) = 872, μ (Mo-K α)

 1.006 mm^{-1} , T_{max} 0.8496, T_{min} 0.7853, crystal size 0.42 x 0.28 x 0.22 mm³. T 203 K.

A total of 19366 reflections, 4462 unique ($R_{int} 0.0335$) was collected $2^{\circ} < \theta < 28^{\circ}$. Final $R_1 0.0371$ (data with I > $2\sigma(I)$), 0.0446(all data), w $R_2 0.0889$, GoF 1.063, final $\Delta e +1.09/-0.47$. 2.4.4 Crystal data for $O[SiPh_2{Co(CO)_4}]_2$ (**3**)

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 $C_{32}H_{20}Co_2O_9Si_2$, M_r 722.52, triclinic, P1, a = 10.362(3), b = 10.383(5), c = 17.331(5) Å, $\alpha = 74.42(2)$, $\beta = 80.87(5)$, $\gamma = 61.34(2)^\circ$, V = 1575.1(10) Å³, $D_{calc} = 1.523$ g cm⁻³, Z = 2, F(000) = 732, μ (Mo-K α) 1.183 mm⁻¹, T_{max} 0.897, T_{min} 0.809, crystal size 0.72 x 0.36 x 0.18 mm³. T 141 K.

A total of 5234 reflections, 4921 unique (R_{int} 0.0376) was collected $2^{\circ} < \theta < 24^{\circ}$. Final R₁ 0.0481 (data with I > 2 σ (I)), wR₂ 0.0978, GoF 1.029, final Δe +0.38/-0.36.

3. Results and Discussion

3.1.1 Ph₂(TolS)Fe(CO)₂Cp (**1a**)

This compound was readily prepared from $ClPh_2SiFe(CO)_2Cp$, on reaction with a stoichiometric amount of Na[STol] in thf. The yellow crystalline material is essentially air-stable but attempted chromatography on silica led to rapid decomposition, giving among other products the known radical cation $[Fe_2(CO)_2Cp_2(STol)_2]^+$, characterised by its ESMS, its v(CO) spectrum and its characteristic ESR spectrum [18]. While this formally involves a transfer of the thiol group from silicon to iron, the reaction was non-specific so mechanistic discussion is unwarranted.

[line diagrams 1-5 near here]

In an attempt to induce a migration reaction, $Ph_2(TolS)SiFe(CO)_2Cp$ was thermolysed or photolysed, in the presence of HMPA to trap any silvlene intermediates. However, only slow decomposition to intractable products was observed, suggesting that the deoligomerisation reaction observed for disilyl complexes of Fe(CO)_2Cp does not have a parallel with Si-S bonds.

It is known that Pt(II) thiolate complexes form stable adducts via a bridging S atom to mercury(II) halides, as in 4 [21]. It was therefore of interest to see if the S atom of $Ph_2(TolS)SiFe(CO)_2Cp$ behaved similarly, to give the complex **5**. When $Ph_2(TolS)SiFe(CO)_2Cp$ and HgI_2 were mixed in d⁶-DMSO in an NMR tube, the ¹H signals from the Cp protons shifted by +0.36 ppm, and the tolyl-group protons also shifted markedly. These changes are consistent with the formation of an adduct of the type **5**. However all attempts to isolate the adduct were thwarted. A yellow solid was formed which contained Hg, I and tolyl groups, and probably contained (TolS)HgI (or the corresponding symmetrised equivalents), while the solution showed carbonyl-region infrared peaks which were consistent with Ph₂ISiFe(CO)₂Cp. All this points to

the initial formation of **5**, but further reaction exchanges the Si-S bond (possibly via a silylene) for a Si-I one. However the system did not provide anything other than this tentative information.

The X-ray structure determination of $Ph_2(TolS)SiFe(CO)_2Cp$ was undertaken since the only previous examples related to this were $(TolS)_{3-n}(Tf)_nSiRu(PMe_3)_2Cp^*$, $[(TolS)_2(MeCN)SiRu(PMe_3)_2Cp^*]^+$ and $[(TolS)(phen)SiRu(PMe_3)_2Cp^*]^{2+}$ from Tilley's group [22], and $Ph_2(PhS)SiRhH_2(PMe_3)_3$ [23]. The structure is illustrated in Fig 1, which shows the expected arrangement about both the Fe and Si atoms.

[Figure 1 near here]

The orientation of the tolyl group is towards the metal fragment, which appears unusual but is presumably the result of crystal packing interactions on a flexible bond. The closest intramolecular approach between the two fragments, O(1)...C(41) (3.39 Å) is essentially the sum of the Van der Waals radii for the two atoms, so there is no barrier to the observed orientation. The Fe-Si bond length is 2.296(2) Å, which can be compared to those in the only other two molecules of the type Ph₂XSiFe(CO)₂Cp to have been determined, X = F 2.278(1) Å [24] and X = OMe 2.292(2) Å (see below). The trend in the order STol > OMe > F is that expected on electronegativity grounds. The Si-S bond length of 2.180(2), and the Si-S-C bond angle of 102.8(2)° are both slightly larger than the corresponding parameters in Ph₃SiSPh (2.156 Å and 99.5° respectively [25]), but the differences are not chemically significant.

*3.3.2 Ph*₂(*MeO*)*SiFe*(*CO*)₂*Cp* (**1b**)

This compound was readily prepared in good yield from Na[OMe] and

Ph₂ClSiFe(CO)₂Cp in thf, showing no tendency to undergo base cleavage of the Fe-Si bond [26]. The spectroscopic characterisation given in the Experimental section is as expected and only the electrospray mass spectra merit further discussion. When run in MeCN/H₂O at low cone voltages the main peaks observed corresponded to $[M+H]^+$ and $[M+NH_4]^+$ arising from chemical ionisation by attachment of H⁺ or NH₄⁺, presumably at the oxygen atom of the methoxy group. More interesting is the behaviour at higher cone voltages, where the fragmentation patterns can yield some revealing information. At low cone voltage settings, fragmentation of the complexes is minimised and the electrospray ionisation mass spectra consist of ions derived from the intact

parent molecule, sometimes also incorporating a molecule of the solvent used as a mobile phase. The appearance of $[M + H + solvent]^+$ ions is strongly dependent on how good a donor the solvent is, so acetonitrile, pyridine and dimethylsulfoxide (DMSO) are common adducts whereas water, methanol and dichloromethane are less frequently observed. These adducts are the first to disappear as the fragmentation energy (cone voltage) is increased. Further fragmentation generally consists of the loss of neutral fragments from the pseudo-molecular ion. These features can all be observed in the energy-dependent electrospray ionisation mass spectrum (EDESI-MS) [27] of Ph₂(MeO)SiFe(CO)₂Cp (Figure 2).

[Figure 2 near here]

This data presentation technique provides a two-dimensional map of cone voltage vs. m/z ratio, and allows visualisation of the entire fragmentation pattern in a convenient format. Ph₂(MeO)SiFe(CO)₂Cp was analysed in an acetonitrile/methanol solution, with trace amounts of DMSO present, and the fragmentation processes are summarised in Scheme 1

At the lowest cone voltage, only the ion $[M + H + DMSO]^+$ is observed (A). The DMSO adduct is observed rather than MeCN or MeOH, because DMSO is a superior donor. As the cone voltage is increased, the ion $[M + H]^+$ (**D**) makes an appearance, following loss of the loosely attached solvent. At ~15 V, three more ions appear, all silylene species resulting from elimination of methanol from the parent ion. The three are $[Ph_2Si=Fe(CO)_2Cp]^+$ (**E**) at 359 *m/z*, with the base-stabilised species $[(MeCN)Ph_2Si=Fe(CO)_2Cp]^+$ (**C**) at 400 *m/z* and

 $[(DMSO)Ph_2Si=Fe(CO)_2Cp]^+$ (**B**) at 437 *m/z*. The last two result from collisions with gas-phase solvent molecules subsequent to methanol elimination. **C** is much more intense than **B** now, because there is no time for an equilibrium to be set up which the better donor can dominate. EDESI-MS/MS [27] experiments show that **B** does not derive directly from **A**; if **A** is selected and subjected to fragmentation in an argon-filled collision cell, **B** does not appear at all. At ~40

V, both CO ligands are lost simultaneously to form $[Ph_2Si=FeCp]^+(G)$. A proportion of these

ions react with MeCN to form $[Ph_2Si=FeCp(MeCN)]^+$ (**F**), with the MeCN presumably attached to the coordinatively unsaturated iron rather than the silicon atom.

[Scheme 1 near here]

The crystal structure of the complex **1b** was determined for comparison with that of $Ph_2(TolS)SiFe(CO)_2Cp$ (**1a**) discussed above. This is shown in Figure 3.

[Figure 3 near here]

The two molecules are similar overall, other than the relative conformation of the STol/OMe groups in the respective examples. The Fe-Si bond lengths are not significantly different despite the more electronegative substituent in the OMe example, a feature that usually leads to bond shortening. The Si-O bond in **1b** is 1.656(5) Å which is shorter than expected by comparison with the Si-S bond of **1a** (2.180(2)Å) and the difference in covalent radii of O and S (0.3-0.4 Å), but this is not unusual for bonds to Si where Si-O bonds are shortened relative to Si-S by stronger π bonding. The wider Si-O-C bond of 125.7(4)° in **1b** compared to the Si-S-C bond of 102.8(2)° in **1a** is also usual for silyl ethers.

3.3.3 $Ph(2-MeOC_6H_4)XSiFe(CO)_2Cp, X = Cl, OMe.$

These compounds were prepared because they have an *ortho*-OMe group on one of the aryl rings that is positioned so that it can interact with the silicon atom, stabilising a silylene centre if one is formed. Ogino has shown that *ortho*- CH_2NMe_2 groups can stabilise iron-silylenes using this approach [23].

The chloro example $Ph(2-MeOC_6H_4)CISiFe(CO)_2Cp$ was prepared analogously to $Ph_2CISiFe(CO)_2Cp$. It could be chromatographed on silica which suggested that the Si-Cl bond is relatively unreactive, and this was confirmed by the lack of any reaction at the Si-Cl bond with Na[STol]. However it did react with [OMe]⁻ to give the corresponding methoxy-silyl species $Ph(2-MeOC_6H_4)(MeO)SiFe(CO)_2Cp$.

In the ESMS of Ph(2-MeOC₆H₄)(OMe)SiFe(CO)₂Cp in MeCN/H₂O, the main peak at low cone voltages was the [M+H]⁺ ion, formed presumably by attachment at the OMe oxygen atom, together with some solvated [M+H+H₂O]⁺. However as the cone voltage was increased, loss of MeOH readily occurred to generate a strong signal from the silylene cation [Ph(2-MeOC₆H₄)Si=Fe(CO)₂Cp]⁺, together with increasing amounts of [Ph(2-MeOC₆H₄)Si=Fe(CO)_nCp]⁺ (n = 1, 0) under more forcing conditions. In this case EDESI-MS studies of Ph(2-MeOC₆H₄)(OMe)SiFe(CO)₂Cp showed that the pendant OMe group prevents coordination of donors including acetonitrile, pyridine and DMSO after the elimination of methanol. This observation is strong evidence that the pendant OMe group provides a degree of intramolecular stabilisation by coordination to the silicon centre, certainly enough to prevent association of other bases.

In an attempt to reproduce the ESMS results on a preparative scale, $Ph(2-MeOC_6H_4)(MeO)SiFe(CO)_2Cp$, was sealed in an NMR tube with $[Me_3O]BF_4$ in CDCl₃, in the hope that the MeO group on silicon would be eliminated as Me₂O. However no reaction took place, even after several days at room temperature.

The X-ray crystal structure of $Ph(2-MeOC_6H_4)CISiFe(CO)_2Cp$ (**2a**) was undertaken to see if there was any tendency for the silicon atom to become five-coordinate. Figure 4 shows that the *ortho*-OMe group is orientated so that the oxygen atom is pointing towards the silicon atom, and there are indications of a weak bonding intra-molecular interaction between these atoms.

[Figure 4 near here]

The Si...O distance of 2.991 Å is shorter than the sum of the van der Waals radii for Si and O at 3.6 Å, though much longer than the sum of the covalent radii of about 1.84 Å. The C(21)-C(26)-O(3) angle is 115.5° , suggesting the methoxy group is drawn towards the silicon to shorten the O..Si distance. The Cl on the Si atom lies opposite to the site where the O...Si interaction is developing. Comparison with the structure of MeC(O)N(SiMe₂Cl)₂ which has Si-Cl bonds to both four- and five-coordinate silicon atoms [28] shows that the value of 2.115 Å for the Si-Cl distance in Ph(2-MeOC₆H₄)ClSiFe(CO)₂Cp is longer than the Si-Cl bond to the tetrahedral silicon atom (2.050 Å) though much shorter than the Si-Cl *trans* to O on the five-coordinate Si (2.348 Å). This is significant, since a major factor determining the tendency for five-coordination

at silicon is the capacity for a Si-X bond to be stretched, which is facile for X = CI [29]. Finally, the sum of the angles about the silicon atom for the potentially equatorial groups [Fe-Si-C(21), Fe-Si-C(31) and C(21)-Si-C(31)] is 342.2°, midway between tetrahedral (328°) and planar (360°). All of this points to a weak bonding O...Si interaction, poised to stabilise the silicon centre once the *trans* group is lost in the mass spectrometer.

The Fe-Si bond in Ph(2-MeOC₆H₄)ClSiFe(CO)₂Cp is 2.2698(6) Å, surprisingly shorter than even Ph₂FSiFe(CO)₂Cp (2.278(1) Å [24]). Based on electronegativity grounds alone a bond length comparable to that of Ph₂(MeO)SiFe(CO)₂Cp (2.292(2) Å) might have been expected. This may be related to the small distortion towards trigonal bipyramidal geometry, with the Fe in a pseudo-equatorial site, induced by the *ortho*-OMe group interaction.

$3.3.4 O[SiPh_2{Co(CO)_4}]_2$

The early report of $O[SiMe_2\{Co(CO)_4\}]_2$ described it as extremely unstable, decomposing slowly even at -78°C [9]. Since 1,2 silyl-migration reactions of $R_5Si_2Co(CO)_4$ have been shown to be extremely facile [3], it was of interest to synthesise a stable example of a heteroatom-substituted silyl-cobalt compound. Accordingly, we reacted $O(SiPh_2H)_2$ with $Co_2(CO)_8$ and obtained good yields of $O[SiPh_2\{Co(CO)_4\}]_2$. This showed none of the instability of the methyl analogue, since it was readily isolated. The solid was even stable for considerable periods in air, although in solution it proved to be moderately air-sensitive. The reasons for the remarkable difference in behaviour between the methyl and phenyl examples are not obvious, but are presumably attributable to the steric protection provided by the phenyl groups. The molecule is very crowded and perhaps is prevented from adopting a conformation where a migration reaction is possible, if this is involved in the decomposition process.

The carbonyl region infrared spectra of $O[SiPh_2\{Co(CO)_4\}]_2$ showed doubling of the normal pattern found for terminal $Co(CO)_4$ groups. This is unlikely to arise from coupling between the two ends of the molecule and suggests perhaps that two conformers exist in solution. To determine the solid state structure an X-ray study was carried out. The crystal contains discrete molecules, as shown in Figure 5.

[Figure 5 near here]

As expected the backbone of the molecule consists of two silicon atoms, both tetrahedrally coordinated, joined in a Si-O-Si linkage. Each silicon atom is further bonded to two phenyl rings and to a cobalt atom which has trigonal bipyramidal coordination, the silicon occupying an axial site. One parameter of interest is the Si-O-Si angle of $153.4(2)^{\circ}$, which is surprisingly bent given the fact that other bulky disiloxanes (e.g. $O(SiPh_3)_2$ [30], $O[SiMeF{Fe(CO)_2Cp}]_2$ [31], $O[Si{OBu^t}_{3]_2}$ [32] and $O[SiCl_2{C(SiMe_3)_3}]_2$ [33] all show linear Si-O-Si bonds. It is generally assumed that the linear arrangement is stabilised by Si..O π -bonding, and consistent with these ideas the Si-O bond length in $O[SiPh_2{Co(CO)_4}]_2$ of 1.630(3) Å is closer to that in $O(SiH_3)_2$ (Si-O-Si 142.2(3)°, Si-O 1.634 Å [34]) than to $O[SiMeF{Fe(CO)_2Cp}]_2$ (Si-O-Si 180°, Si-O 1.603 Å [31]).

The Co-Si bonds (2.348 Å average) are relatively long, with previous values for simple $R_3SiCo(CO)_4$ ranging from 2.254 Å to 2.38 Å [35]. This is presumably because of the bulk of the substituents in **3**.

In the solid state, molecules of $O[SiPh_2{Co(CO)_4}]_2$ adopt a *gauche* configuration for the two cobalt-containing groups, with interlocking of CO groups and Ph substituents. A possible explanation for the complex IR spectrum in solution is that there is also an *anti* configuration which interchanges slowly on the IR time scale, giving rise to doubled peaks. This seems more likely than coupling of vibrational modes across the two halves of the molecule.

4. Supplementary material.

Crystallographic data for the structural analyses have been deposited with the Cambridge Crystallographic Data Centre, CCDC no 172420-172423 for **1a**, **1b**, **2a** and **3** respectively. Copies of this information may be obtained free of charge from the Director, CCDC, 12 Union Rd., Cambridge CB2 1EZ, UK (Fax: +44-1223-336033; e-mail: deposit@ccdc.cam.ac.uk or www: http://www.ccdc.cam.ac.uk).

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Captions to Figures

Figure 1. The structure of Ph₂(TolS)SiFe(CO)₂Cp. Bond parameters include: Fe(1)-Si(1) 2.296(2), Si(1)-S(1) 2.180(2), Fe(1)-C_{cp} (av) 2.097(6) Å; Fe(1)-Si(1)-S(1) 113.68°, Si(1)-S(1)-C(41) 102.8(2), Fe(1)-Si(1)-C(21) 118.1(2)°, Fe(1)-Si(1)-C(31) 112.0(2)°.

Figure 2. Positive-ion EDESI-MS map [27] of Ph₂(MeO)SiFe(CO)₂Cp, run in a mobile phase of MeCN/MeOH with traces of DMSO present.

Figure 3. The structure of Ph₂(MeO)SiFe(CO)₂Cp. Bond parameters include: Fe(1)-Si(1) 2.293(3), Si(1)-O(3) 1.656(5), C(3)-O(3) 1.421(8), Fe(1)-C_{cp} (av) 2.093(7) Å; Fe(1)-Si(1)-O(3) 112.3(2)°, Si(1)-O(3)-C(3) 125.7(4), Fe(1)-Si(1)-C(11) 116.1(2)°, Fe(1)-Si(1)-C(21) 114.9(2)°.

Figure 4. The structure of Ph(2-MeOC₆H₄)ClSiFe(CO)₂Cp. Bond parameters include: Fe(1)-Si(1) 2.2698(2), Si(1)-Cl(1) 2.1151(7), Fe(1)-C_{cp} (av) 2.093(2) Å, Si(1)...O(3) 2.991 Å; Fe(1)-Si(1)-Cl(1) 108.09(3)°, C(26)-O(3)-C(3) 118.8(2), C(21)-C(26)-O(3) 115.5(2)°.

Figure 5. The structure of O[SiPh₂(Co(CO)₄]₂. Bond parameters include: Co(1)-Si(1) 2.341(2), Co(2)-Si(2) 2.354(2), Si(1)-O(9) 1.630(3), Si(2)-O(9) 1.629(3) Å; Si(1)-O(9)-Si(2) 153.4(2)°.