Manganese(I) selenoether chemistry: synthesis, multinuclear NMR studies and the structures of $[MnCl(CO)_3(MeSeCH_2CH_2SeMe)]$, $[MnCl(CO)_3(MeSeCH_2CH_2CH_2SeMe)]$ and $[MnBr(CO)_3\{C_6H_4(SeMe)_2-o\}]$



Julie Connolly, Maxwell K. Davies and Gillian Reid*

Department of Chemistry, University of Southampton, Highfield, Southampton, UK SO17 1BJ

Received 3rd September 1998, Accepted 5th October 1998

Reaction of $[Mn(CO)_5X]$ (X = Cl, Br or I) with RSe(CH₂)_nSeR (R = Me or Ph, n = 2; R = Me, n = 3) or C₆H₄(SeMe)₂-o in refluxing CHCl₃ yielded the neutral manganese(II) complexes $[MnX(CO)_3 \{RSe(CH_2)_nSeR\}]$ or $[MnX(CO)_3 \{C_6H_4(SeMe)_2-o\}]$ as yellow or orange solids. Infrared spectroscopic studies confirmed the *fac*-tricarbonyl arrangement. Proton, ¹³C-{¹H}, ⁷⁷Se-{¹H} and ⁵⁵Mn NMR spectroscopy has been used to probe the solution behaviour, and show that only those compounds involving PhSeCH₂CH₂SePh are undergoing rapid pyramidal inversion on the NMR timescale, with the individual NMR distinguishable invertomers observed for the other complexes (*meso*-1, *meso*-2 and DL) in varying ratios. X-Ray crystallographic analyses on three examples confirmed a *fac*-tricarbonyl arrangement, with the diselenoether ligand chelating: $[MnCl(CO)_3(MeSeCH_2CH_2SeMe)]$ and $[MnCl(CO)_3(MeSeCH_2CH_2SeMe)]$ adopt the DL arrangement, while $[MnBr(CO)_3\{C_6H_4(SeMe)_2-o\}]$ is in the *meso*-2 form in the solid state. These are the first structure determinations on selenoether complexes of manganese(I) carbonyl halides. Most importantly, ⁵⁵Mn NMR spectroscopic studies show that $\delta(^{55}Mn)$ is to low frequency of those of the corresponding thioether compounds, lying in the range $\delta - 175$ to -702, the lowest frequencies occurring for the iodo derivatives. The CO stretching frequencies and ⁵⁵Mn NMR shifts show that, for a given X, the manganese(I) centre in $[MnX(CO)_3(diselenoether)]$ is more shielded than in $[MnX(CO)_3(dithioether)]$, possibly indicating increased σ donation in the former.

Introduction

Recently we have been investigating the co-ordination chemistry of macrocyclic thio- and seleno-ethers with early transition metal centres, with particular interest in the hard (metal)-soft (ligand) interactions in these species. In the course of this work we have reported a range of complexes of hard chromium(III) and vanadium(III) ions, including [CrX2- $([16]aneSe_4)]PF_6$ (X = Cl or Br; [16]aneSe_4 = 1,5,9,13-tetraselenacyclohexadecane), cis-[CrX₂([14]aneS₄)]PF₆ ([14]aneS₄ = 1,4,8,11-tetrathiacyclotetradecane) and $[VX_3([9]aneS_3)]$ $(X = Cl, Br \text{ or } I; [9]aneS_3 = 1,4,7-trithiacyclononane). EXAFS$ measurements and single crystal X-ray analyses have enabled us to establish the metal-thioether and -selenoether distances in these compounds.¹⁻³ More recently we have initiated a systematic investigation of manganese(I) carbonyl complexes incorporating Group 15 and 16 donor ligands. The lower electronegativity of Se over S leads one to anticipate that R₂Se might be a better ligand than R₂S to transition metal ions. However, experimental data to substantiate this are very limited. We proposed that 55 Mn NMR and IR [ν (CO)] data on a series of manganese(I) carbonyl halide adducts involving thioether and selenoether ligands might provide experimental evidence to support or refute this suggestion; ⁵⁵Mn is 100% I = 5/2 ($\Xi = 24.840$ MHz) and possesses a moderately high quadrupole moment (0.55×10^{-28} m²), thus for complexes with less than $O_{\rm h}$ or $T_{\rm d}$ symmetry a considerable electric field gradient is expected to result in substantial broadening of the resonances. However, 55Mn NMR studies have been conducted on several different classes of compound, including for example [Mn₂(CO)₁₀] and derivatives, [Mn(CO)₅X], [MnCp(CO)₃],

[Mn(CO)₃X(SbPh₃)₂] and [Mn(CO)₄X(η¹-Ph₂SbCH₂SbPh₂)]⁴⁻⁹ and we have reported ¹⁰ the preparation and characterisation of [MnX(CO)₃(L-L)] [L-L = MeSCH₂CH₂SMe, MeSCH₂-CH₂CH₂SMe, PhSCH₂CH₂SPh or C₆H₄(SMe)₂-o]. These species adopt *fac*-tricarbonyl arrangements with the dithioether chelating. The ⁵⁵Mn NMR studies showed that where pyramidal inversion was slow on the NMR timescale the resonances for the individual invertomers were generally well resolved, giving δ (⁵⁵Mn) in the range +67 to -537. All of these species are significantly deshielded with respect to phosphine complexes such as [MnX(CO)₃(dppe)] (X = Cl, δ -1141; X = Br, -1254).¹¹

In this paper we describe the first detailed preparations and spectroscopic characterisation, including ${}^{1}H$, ${}^{13}C-{}^{1}H$ }, ^{55}Mn and $^{77}Se-\{^{1}H\}$ NMR spectroscopy, of the manganese(1) diselenoether complexes $[MnX(CO)_3(L-L)]$ [X = Cl, Br or I; $L-L = MeSeCH_2CH_2SeMe$, $MeSeCH_2CH_2CH_2SeMe$, PhSe-CH₂CH₂SePh or C₆H₄(SMe)₂-o] and compare the results with those for related thioether compounds. The crystal structures of [MnCl(CO)₃(MeSeCH₂CH₂SeMe)], [MnCl(CO)₃(MeSeCH₂- CH_2CH_2SeMe)] and $[MnBr(CO)_3\{C_6H_4(SeMe)_2-o\}]$ are also described. Prior to this work there were very few known examples of selenoether complexes of Mn^I. The complex [MnBr(CO)₃(MeSeCH₂CH₂SeMe)] was first described by Abel and Hutson¹² some 30 years ago. We have resynthesized this in order to obtain multinuclear NMR data to allow comparisons with the other selenoether compounds and with the thioether analogue. The only other manganese(I) selenoether compounds reported are [Mn(NO)₃(Ph₂Se)₂]¹³ and the dinuclear $[{Mn(C_5H_4Me)(CO)_2}_2(SeMe_2)]$ for which a crystal structure shows a bridging Me₂Se ligand.¹

Table 1 Infrared spectroscopic data (CO region) and FAB mass spectrometric data

Compound	$\tilde{v}(CO)^{a}/cm^{-1}$	m/z ^b
[MnCl(CO) ₃ (MeSeCH ₂ CH ₂ SeMe)] [MnBr(CO) ₂ (MeSeCH ₂ CH ₃ SeMe)]	2034, 1959, 1917 2031, 1955, 1917	392, $[MnCl(CO)_3(L-L)]^+$; 357, $[Mn(CO)_3(L-L)]^+$; 308, $[MnCl(L-L)]^+$ 436, $[MnBr(CO)_3(L-L)]^+$; 357, $[Mn(CO)_3(L-L)]^+$; 352, $[MnBr(L-L)]^+$
[MnI(CO) ₃ (MeSeCH ₂ CH ₂ SeMe)]	2027, 1955, 1918	484, [MnI(CO) ₃ (L–L)] ⁺ ; 400, [MnI(L–L)] ⁺ ; 357, [Mn(CO) ₃ (L–L)] ⁺
[MnCl(CO) ₃ (MeSeCH ₂ CH ₂ CH ₂ SeMe)]	2032, 1955, 1917	$371, [Mn(CO)_3(L-L)]^+; 322, [MnCl(L-L)]^+$
$[MnBr(CO)_{3}(MeSeCH_{2}CH_{2}CH_{2}SeMe)]$	2030, 1954, 1918	$371, [Mn(CO)_3(L-L)]^+; 366, [MnBr(L-L)]^+$
$[MnI(CO)_{3}(MeSeCH_{2}CH_{2}CH_{2}SeMe)]$	2025, 1951, 1918	414, $[MnI(L-L)]^+$; 371, $[Mn(CO)_3(L-L)]^+$
[MnCl(CO) ₃ (PhSeCH ₂ CH ₂ SePh)]	2034, 1956, 1925	481, $[Mn(CO)_3(L-L)]^+$; 432, $[MnCl(L-L)]^+$
$[MnBr(CO)_{3}(PhSeCH_{2}CH_{2}SePh)]$	2034, 1960, 1927	481, $[Mn(CO)_3(L-L)]^+$; 476, $[MnBr(L-L)]^+$
[MnI(CO) ₃ (PhSeCH ₂ CH ₂ SePh)]	2029, 1957, 1928	524, $[MnI(L-L)]^{+}$; 481, $[Mn(CO)_{3}(L-L)]^{+}$
$[MnCl(CO)_3 \{C_6H_4(SeMe)_2 - o\}]$	2037, 1964, 1924	405, $[Mn(CO)_3(L-L)]^+$; 356, $[MnCl(L-L)]^+$
$[MnBr(CO)_3 \{C_6H_4(SeMe)_2 - o\}]$	2035, 1963, 1924	484, $[MnBr(CO)_3(L-L)]^+$; 405, $[Mn(CO)_3(L-L)]^+$; 400, $[MnBr(L-L)]^+$
$[MnI(CO)_3\{C_6H_4(SeMe)_2-o\}]$	2030, 1960, 1924	532, $[MnI(CO)_3(L-L)]^+$; 448, $[MnI(L-L)]^+$; 405, $[Mn(CO)_3(L-L)]^+$
^{<i>a</i>} Solutions in CHCl ₃ , all CO bands were stro	ng. ^b Recorded using 3-ni	trobenzyl alcohol matrix; major peaks only.

Results and discussion

Syntheses

Treatment of $[MnX(CO)_5]$ (X = Cl, Br or I) with L–L [MeSeCH₂CH₂SeMe, MeSeCH₂CH₂CH₂SeMe, PhSeCH₂CH₂-SePh or C₆H₄(SeMe)₂-o] in gently refluxing CHCl₃ affords the neutral species $[MnX(CO)_3(L-L)]$ as yellow or orange solids. In all cases the reaction flasks were wrapped with foil to protect the manganese species from visible light. The isolated compounds are very soluble in chlorocarbons, but poorly soluble in hydrocarbons. The solids are reasonably stable, although solutions of the compounds decompose rapidly when exposed to oxygen and/or sunlight.

FAB Mass spectrometric data for these compounds are given in Table 1. Generally, peaks with the correct isotopic distributions consistent with $[MnX(L-L)]^+$ and $[Mn(CO)_3(L-L)]^+$ are observed, although the parent molecule and further peaks attributed to loss of X are also seen in a number of cases. Solution IR spectroscopic studies show three CO stretching vibrations, consistent with formation of *fac*-[MnX(CO)_3(L-L)]; $\nu(CO)$ shifts to low frequency according to the series X = Cl > Br > I, although it is virtually insensitive to changes in the diselenoether ligand L. Importantly however, the values of $\nu(CO)$ for these manganese selenoether compounds are consistently slightly to low frequency of those observed for [MnX(CO)_3(dithioether)], consistent with increased occupation of the CO π^* orbitals in the former.¹⁰

NMR Spectroscopy

We have recorded ¹H, ¹³C-{¹H}, ⁷⁷Se-{¹H} and ⁵⁵Mn NMR data for the compounds in this work. The combination of observable NMR nuclei, particularly 13C-{1H}, 55Mn and ⁷⁷Se-{¹H}, provides a very powerful way in which to monitor the species present in solution. Four isomers are possible for fac-[MnX(CO)₃(diselenoether)]; meso-1, meso-2 and a pair of NMR indistinguishable DL enantiomers, with interconversion occurring via pyramidal inversion at the co-ordinated selenoether donors. Abel et al.¹⁵ have conducted detailed, quantitative NMR spectroscopic studies on a wide range of transition metal complexes with Group 16 donor ligands to probe inversion processes, and, the donor atom type, the nature of the interdonor linkage and the choice of transition metal have all been shown to be important parameters. In this work we are concerned with monitoring the trends in δ ⁽⁵⁵Mn) and δ ⁽⁷⁷Se) to probe the Se-Mn co-ordination rather than with the inversion barriers. Indeed Abel et al. have commented that the substantial quadrupole moment associated with ⁵⁵Mn is expected to lead to extensive broadening and hence these compounds are not suited to detailed dynamic ¹H NMR studies; we have shown that this is also true for [MnX(CO)₃(dithioether)], where broad resonances were observed at room temperature even for compounds where inversion is slow on the NMR timescale; cooling the solutions sharpened the signals slightly.¹⁰



For the selenoether species ¹H NMR spectra were recorded at 300 K in CDCl₃ solution. These were not very informative since the broad ligand resonances made assignment of the isomers impossible. ¹³C-{¹H} NMR spectroscopy is rather more useful, providing evidence that pyramidal inversion is rapid on the NMR timescale at 300 K only for the compounds involving PhSeCH₂CH₂SePh. For these systems a single resonance is observed for the CH₂ groups and this is to high frequency of that for the 'free' ligand (δ 27.3). For the other compounds several resonances are observed for the CH₂ units and up to four for the Me groups (one for each of the meso-1 and meso-2 and two for the DL isomer), also to high frequency of the corresponding resonance of the 'free' ligand, confirming the presence of up to three NMR distinguishable invertomers. As for the dithioether species, the CO resonances are very broad even for the complexes where inversion is slow on the NMR timescale, due to rapid quadrupolar relaxation from the directly bonded ⁵⁵Mn nucleus and hence the individual CO resonances are not resolved.

The ⁷⁷Se-{¹H} and ⁵⁵Mn NMR data for [MnX(CO)₃(L-L)] are presented in Table 2. One 77Se resonance is expected for each of the meso isomers and two for the DL isomer. The results from the ⁷⁷Se-{¹H} NMR studies are in accord with those from the ${}^{13}C-{}^{1}H$ NMR studies. The rapidly inverting [MnX(CO)₃-(PhSeCH₂CH₂SePh)] species each show a single resonance approximately 100 ppm to high frequency of that of free PhSeCH₂CH₂SePh, confirming formation of a five-membered chelate ring.¹⁶ Complexes involving the other diselenoether ligands each show the presence of either all three NMR distinguishable isomers (or two of the three); in cases where we only observe three $\delta(^{77}Se)$ resonances we conclude that either only two isomers are present in appreciable quantity (meso-2 and DL) or the resonances are coincidental. The co-ordination shifts $[\delta(\text{complex}) - \delta(\text{'free' ligand})]$ for MeSeCH₂CH₂SeMe and C₆H₄(SeMe)₂-o are approximately 120 to 170 and 170 to 200 ppm respectively. These large shifts are consistent with the formation of a five-membered chelate ring, whereas δ (⁷⁷Se- $\{^{1}H\}$) for the complexes of MeSeCH₂CH₂CH₂SeMe, which involve a six-membered chelate ring, are close to those of the 'free' ligand.

Compound	$\delta(^{55}Mn) (w_{1/2}/Hz)^{b}$	$\delta(^{77}\text{Se-}\{^1\text{H}\})^c$	Approximate ratio <i>meso-1:meso-2:</i> DL
[MnCl(CO) ₃ (MeSeCH ₂ CH ₂ SeMe)]	-277^{d} (2000), -342 (2000), -410 (2900)	256, 269, 282, 290	1:3:6
[MnBr(CO) ₃ (MeSeCH ₂ CH ₂ SeMe)]	$-345^{d}(900), -428(1500), -506(2500)$	259 (meso-2); 281, 289 (DL)	1:5:6
[MnI(CO) ₃ (MeSeCH ₂ CH ₂ SeMe)]	$-500^{d} (\approx 1340), -607 (1220), -698 (2170)$	283 (meso-2); 239, 276 (DL)	1:5:10
[MnCl(CO) ₃ (MeSeCH ₂ CH ₂ CH ₂ SeMe)]	-175 (2080), -190 (sh), -219 (sh)	75, 80, 92, 118	1:2:4
[MnBr(CO) ₃ (MeSeCH ₂ CH ₂ CH ₂ SeMe)]	-257(2000), -317(2000)	62, 65, 81, 109	1:2:4
[MnI(CO) ₃ (MeSeCH ₂ CH ₂ CH ₂ SeMe)]	-402^{d} (sh), -426 (1550), -499 (1760)	39, 66, 73, 92	1:2:7
[MnCl(CO) ₃ (PhSeCH ₂ CH ₂ SePh)] ^e	-249 (5200)	445	Inverting
[MnBr(CO) ₃ (PhSeCH ₂ CH ₂ SePh)] ^e	-346 (4590)	443	Inverting
[MnI(CO) ₃ (PhSeCH ₂ CH ₂ SePh)] ^e	-537 (3170)	438	Inverting
$[MnCl(CO)_{3}\{C_{6}H_{4}(SeMe)_{2}-o\}]$	$-247^{d}(2000), -311(2600), -394(4500)$	386, 400 (DL), 406 (meso-2)	1:5:4
$[MnBr(CO)_{3}\{C_{6}H_{4}(SeMe)_{2}-o\}]$	$-325^{d}(1070), -414(1860), -507(3000)$	375, 386 (DL), 400 (meso-2)	1:10:5
$[MnI(CO)_3\{C_6H_4(SeMe)_2-o\}]$	-465, ^{<i>d</i>} -581 (1500), -702 (3200)	374, 388 (DL), 396 (meso-2)	1:14:4

^{*a*} Spectra recorded in CHCl₃–CDCl₃ solution at 300 K. ^{*b*} At 89.27 MHz and referenced to external aqueous KMnO₄. ^{*c*} At 68.68 MHz and referenced to neat external Me₂Se. δ (⁷⁷Se) for 'free' ligands: MeSeCH₂CH₂SeMe, 121; MeSeCH₂CH₂CH₂SeMe, 74; PhSeCH₂CH₂SePh, 340; C₆H₄(SeMe)₂-*o*, 202. ^{*d*} Minor isomer. ^{*e*} Inverting at 300 K.



Fig. 1 The ⁵⁵Mn (89.27 MHz, CDCl₃) (a) and ${}^{13}C{}^{1}H$ (90.1 MHz, CDCl₃) NMR spectrum (b) of [MnBr(CO)₃{C₆H₄(SeMe)₂-o}] (methyl region only).

The ⁵⁵Mn NMR studies also confirm that at 300 K only the complexes involving PhSeCH₂CH₂SePh are undergoing rapid pyramidal inversion on the NMR timescale, showing only a single broad resonance. For the other compounds up to three distinct resonances attributable to the presence of up to three NMR distinguishable stereoisomers were observed (Fig. 1). The linewidths (<3000 Hz) are very similar to those for the thioether analogues reported previously, and again considerably smaller than for phosphine systems such as [MnX(CO)₃-(PPh₂H)₂] ($w_{1/2}$ ca. 10 000 Hz).¹¹ For the thioether species, ⁵⁵Mn NMR studies showed that for the compounds involving MeSCH₂CH₂SMe and C₆H₄(SMe)₂-o pyramidal inversion is slow on the ⁵⁵Mn NMR timescale at 300 K, whereas for MeSCH₂CH₂CH₂SMe or PhSCH₂CH₂SPh inversion is rapid at

this temperature. Thus, the energy barrier to inversion is higher for the manganese(I) selenoether complexes than for the corresponding thioethers. This is consistent with the trends observed in other transition metal complexes of these ligands.¹⁶ As for the thioether complexes we observe a shift in δ ⁽⁵⁵Mn) to low frequency upon changing X from Cl to Br to I. Of greatest significance, however, is the observation that $\delta(^{55}Mn)$ for the diselenoether compounds occurs to low frequency (more shielded) of the thioether compounds by ca. 100 ppm. This is consistent with the trend in v(CO) discussed above, *i.e.* v(CO)for the selenoether compounds is to low frequency of v(CO) for the thioethers. These results indicate that there is more electron density associated with the manganese(I) centre in the selenoether compounds (greater occupation of the CO π^* orbitals). This can arise in three ways: greater π donation from the selenoether donor, poorer π acceptance by the selenoether donor, or increased σ donation by the selenoether donor. Since there is little or no evidence for π bonding from thio- or seleno-ether donors to metal centres, it is highly likely that these results indicate an increase in $L \rightarrow Mn^{I} \sigma$ donation from thio- to selenoether. This would also support the work of Hoffmann and coworkers¹⁷ who predicted theoretically that $R_2 E \rightarrow M \sigma$ donation (E = S, Se or Te) would increase in the order $S < Se \ll Te$. Abel and co-workers¹⁸⁻²⁰ have described the preparations

Abel and co-workers¹⁸⁻²⁰ have described the preparations of the rhenium(I) complexes $[ReX(CO)_3(L-L)]$ [X = Cl, Br or I; L-L = MeE(CH₂)_nEMe or MeECH=CHEMe (E = S or Se, n = 2 or 3)], including the crystal structure of *fac*-[ReI(CO)₃(MeSeCH₂CH₂SeMe)]. Detailed ¹H NMR studies revealed that the energy barrier associated with inversion at selenium is higher than at sulfur, and that changing the halogen has a negligible effect upon the overall energy barrier. The results from our work on manganese(I) compounds are consistent with these observations. Also, the relative populations of the invertomers in solution vary with the particular system although the *meso*-2 and DL forms are always dominant, and population of the *meso*-1 form decreases with increasing halide size (due to unfavourable Me···X or Ph···X interactions).

X-Ray crystallography

A search of the Cambridge Crystallographic Database revealed that there are no structurally characterised examples of selenoether complexes of manganese(I) carbonyl halides, although the structure of the dinuclear species $[{Mn(C_5H_4Me)-(CO)_2}_2(SeMe_2)]$ is known.¹⁴ In order to identify the isomeric arrangement in the solid state and to compare the Mn–Se bond lengths with those in the Me₂Se-bridged dimer [${Mn(C_5H_4Me)-(CO)_2}_2(SeMe_2)$] and with d(Mn–S) in the thioether analogues, we have conducted single crystal X-ray analyses on [MnCl-(CO)₃(MeSeCH₂CH₂SeMe)], [MnCl(CO)₃(MeSeCH₂CH₂CH₂-



Fig. 2 View of the structure of $[MnCl(CO)_3(MeSeCH_2CH_2SeMe)]$ with numbering scheme adopted. Ellipsoids are drawn at 40% probability.



SeMe)] and $[MnBr(CO)_3 \{C_6H_4(SeMe)_2 - o\}]$. Crystals of the complexes were obtained from vapour diffusion of light petroleum into a solution of the appropriate complex in CH₂Cl₂. The crystal structures of [MnCl(CO)₃(MeSeCH₂CH₂-SeMe)] (Fig. 2, Table 3) and [MnCl(CO)₃(MeSeCH₂CH₂CH₂-SeMe)] (Fig. 3, Table 4) both show a distorted octahedral geometry at Mn, with a fac arrangement for the three CO ligands and the chelating diselenoether adopting the DL arrangement, with one Me group on each side of the MnSe₂C₂ plane: [MnCl(CO)₃(MeSeCH₂CH₂SeMe)], Mn-Se(1) 2.481(3), Mn-Se(2) 2.467(3), Mn-Cl(1) 2.406(4), Mn-C(5) 1.79(2), Mn-C(6) 1.79(2), Mn-C(7) 1.80(2) Å; [MnCl(CO)₃(MeSeCH₂-CH₂CH₂SeMe)], Mn-Se(1) 2.474(2), Mn-Se(2) 2.482(2), Mn-Cl(1) 2.379(3), Mn-C(6) 1.792(9), Mn-C(7) 1.805(10), Mn-C(8) 1.810(9) Å. The Se-Mn-Se angles involved in the chelate rings are 88.79(8) and 89.29(5)° respectively. The DL isomer was also identified in the crystal structure of [MnCl(CO)₃-(MeSCH₂CH₂SMe)₂], suggesting that this might be the preferred solid state arrangement for [Mn(CO)₃X{MeE- $(CH_2)_n EMe$], E = S or Se, and indeed this is the dominant isomer in solution at room temperature for these species. The

$\overline{\text{Se}(1)}$ – $Mn(1)$	2.481(3)	Se(1) - C(1)	1.96(1)
Se(1)-C(2)	1.98(2)	Se(2)-Mn(1)	2.467(3)
Se(2)-C(3)	1.95(1)	Se(2)-C(4)	1.96(1)
Mn(1)-Cl(1)	2.406(4)	Mn(1) - C(5)	1.79(2)
Mn(1)-C(6)	1.79(2)	Mn(1)-C(7)	1.80(2)
O(1)–C(5)	1.16(2)	O(2)–C(6)	1.18(2)
O(3)–C(7)	1.15(2)	C(2) - C(3)	1.47(2)
Mn(1)-Se(1)-C(1)	109.0(5)	Mn(1)-Se(1)-C(2)	101.6(4)
C(1)-Se(1)-C(2)	96.6(7)	Mn(1)-Se(2)-C(3)	101.2(4)
Mn(1)-Se(2)-C(4)	107.9(5)	C(3)-Se(2)-C(4)	96.3(7)
Se(1)-Mn(1)-Se(2)	88.79(8)	Se(1)-Mn(1)-Cl(1)	92.2(1)
Se(1)-Mn(1)-C(5)	86.4(5)	Se(1)-Mn(1)-C(6)	176.6(5)
Se(1)-Mn(1)-C(7)	90.2(5)	Se(2)-Mn(1)-Cl(1)	82.7(1)
Se(2) - Mn(1) - C(5)	95.2(4)	Se(2)-Mn(1)-C(6)	90.1(4)
Se(2)-Mn(1)-C(7)	175.1(5)	Cl(1)-Mn(1)-C(5)	177.5(5)
Cl(1)-Mn(1)-C(6)	90.8(5)	Cl(1)-Mn(1)-C(7)	92.6(5)
C(5)-Mn(1)-C(6)	90.4(6)	C(5)-Mn(1)-C(7)	89.5(6)
C(6)-Mn(1)-C(7)	91.1(7)		

Table 4 Selected bond lengths (Å) and angles (°) for $[MnCl(CO)_3(MeSeCH_2CH_2SeMe)]$

Se(1)–Mn(2)	2.474(2)	Se(1)–C(1)	1.939(10)
Se(1)-C(2)	1.96(1)	Se(2)-Mn(2)	2.482(2)
Se(2) - C(4)	1.961(10)	Se(2) - C(5)	1.94(1)
Mn(2)-Cl(1)	2.379(3)	Mn(2) - C(6)	1.792(9)
Mn(2)-C(7)	1.805(10)	Mn(2)-C(8)	1.810(9)
O(1)–C(6)	1.14(1)	O(2) - C(7)	1.16(1)
O(3)–C(8)	1.14(1)	C(3) - C(4)	1.53(2)
C(2) - C(3)	1.50(2)		
Mn(2)-Se(1)-C(1)	105.4(3)	Mn(2)-Se(1)-C(2)	107.4(3)
C(1)-Se(1)-C(2)	95.5(5)	Mn(2)-Se(2)-C(4)	110.1(3)
Mn(2)-Se(2)-C(5)	107.3(3)	C(4)-Se(2)-C(5)	97.9(5)
Se(1)-Mn(2)-Se(2)	89.29(5)	Se(1) - Mn(2) - Cl(1)	90.65(8)
Se(1)-Mn(2)-C(6)	88.0(3)	Se(1)-Mn(2)-C(7)	179.2(3)
Se(1)-Mn(2)-C(8)	90.5(3)	Se(2) - Mn(2) - Cl(1)	86.56(8)
Se(2)-Mn(2)-C(6)	89.9(3)	Se(2)-Mn(2)-C(7)	90.2(3)
Se(2)-Mn(2)-C(8)	178.8(3)	Cl(1)-Mn(2)-C(6)	176.2(3)
Cl(1)-Mn(2)-C(7)	89.8(4)	Cl(1)-Mn(2)-C(8)	92.3(3)
C(6)-Mn(2)-C(7)	91.4(5)	C(6) - Mn(2) - C(8)	91.3(4)
C(7)-Mn(2)-C(8)	90.0(4)		

crystals of $[MnBr(CO)_3 \{C_6H_4(SeMe)_2-o\}]^{\dagger}$ were of poorer quality and hence the data less reliable, therefore detailed comparisons of the geometric parameters requires caution. However, the molecular arrangement (Fig. 4) is very similar to the structures described above, with three mutually *fac* CO ligands, a chelating diselenoether and a Br ligand completing the distorted octahedron. In this case the co-ordinated diselenoether adopts the *meso*-2 form with both Me groups directed to the opposite side of the MnSe₂C₂ plane from the Br ligand, Mn– Se(1) 2.454(6), Mn–Se(2) 2.468(6), Mn–Br(1) 2.562(6), Mn– C(9) 1.79(4), Mn–C(10) 1.82(3), Mn–C(11) 1.82(3) Å. This is also the isomer which dominates in solution at room temperature. The Se–Mn–Se angle in the chelate ring is 88.3(2)°. As expected on the basis of the radii of Se *vs.* S, the Mn–Se bond lengths in these compounds are approximately 0.10–0.15 Å

[†] Crystal data: *M* 482.96, orthorhombic, space group *Pna2*₁, *a* = 21.34(2), *b* = 9.50(2), *c* = 7.08(1) Å, *V* = 1448(6) Å³, *Z* = 4, *D_c* = 2.214 g cm³, μ (Mo-K α), 86.97 cm⁻¹. 1513 unique observed reflections, 1148 with $I_o > 2\sigma(I_o)$, *R* = 0.084, *R'* = 0.105. The structure was solved in *Pna2*₁ (since the data indicated an acentric space group) by direct methods²¹ and developed by iterative cycles of full-matrix least-squares refinement and difference Fourier syntheses (an attempt to solve the structure in *Pnma* yielded no solution).²² The Mn, Br, Se and O atoms were refined anisotropically while H atoms were placed in fixed, calculated positions with *d*(C–H) = 0.96 Å. The weighting scheme $w^{-1} = \sigma^2(F)$ gave satisfactory agreement analyses and the Flack parameter indicated that the hand chosen was the correct enantiomorph.



Fig. 4 View of the structure of $[MnBr(CO)_3 \{C_6H_4(SeMe)_2-o\}]$ with the numbering scheme adopted. Ellipsoids are drawn at 40% probability. Mn-Se(1) 2.454(6), Mn-Se(2) 2.468(6), Mn-Br(1) 2.562(6), Mn-C(9) 1.79(4), Mn-C(10) 1.82(3), Mn-C(11) 1.82(3) Å.

longer than d(Mn-S) in the corresponding thioether compounds. However, the increased σ bonding in the selenoethers compared to the thioethers deduced from the spectroscopic studies would be expected to have a very small effect on the M-S and M-Se distances, and is therefore not expected to be detected by X-ray crystallography. The Mn-Se bond lengths in the selenoether compounds reported here are longer than those observed in the SeMe₂ bridged species $[{Mn(C_5H_4Me)(CO)_2}_2]$ (SeMe₂)] [2.375(1), 2.378(1) Å].¹⁴ This may be attributed to the repulsive effect of the second Se-based lone pair in the mononuclear species, whereas in the bridged dimer there is no such repulsion.

Experimental

Infrared spectra were measured as CsI discs using a Perkin-Elmer 983 spectrometer over the range 200-4000 cm⁻¹, or in solution using NaCl plates on a Perkin-Elmer 1600 FTIR spectrometer, mass spectra by fast-atom bombardment (FAB) using 3-nitrobenzyl alcohol as matrix on a VG Analytical 70-250-SE normal geometry double focusing mass spectrometer, ¹H NMR spectra in CDCl₃ at 300 MHz unless otherwise stated, using a Bruker AM300 spectrometer, ¹³C-{¹H}, ⁵⁵Mn and ⁷⁷Se-{¹H} NMR spectra using a Bruker AM360 spectrometer operating at 90.1, 89.27 or 68.68 MHz respectively and referenced to Me₄Si, external saturated, aqueous K[MnO₄] and external neat Me₂Se respectively (δ 0); [Cr(acac)₃] was added to the NMR solutions prior to recording $^{13}\text{C-}\{^1\text{H}\}$ and $^{77}\text{Se-}\{^1\text{H}\}$ spectra and a pulse delay of 2 s was employed for the ¹³C-{¹H} spectra to take account of the long relaxation times. The compounds $[MnX(CO)_5]$ (X = Cl, Br or I), MeSeCH₂CH₂-SeMe, MeSeCH₂CH₂CH₂SeMe, PhSeCH₂CH₂SePh and C₆H₄- $(SeMe)_2$ -o were prepared according to literature procedures.^{23,24}

Preparations

All of the compounds were synthesized by the same general procedure, hence only one is described in detail below.

[MnCl(CO)₃(MeSeCH₂CH₂SeMe)]. To a degassed solution of [MnCl(CO)₅] (0.070 g, 0.30 mmol) in CHCl₃ (30 ml), MeSeCH₂CH₂SeMe (0.066 g, 0.305 mmol) was added. The resulting mixture was gently refluxed for 24 h, and the solution IR spectrum monitored until the reaction had gone to completion. The solvent volume was reduced in vacuo to ca. 2 ml to give a yellow solution. Addition of light petroleum (bp 40-60 °C) at 0 °C afforded a yellow solid which was filtered off, washed with light petroleum and dried in vacuo. The compound was recrystallised from CHCl3-light petroleum (yield 0.087 g, 73%) (Found: C, 21.3; H, 2.7. Calc. for C₇H₁₀ClMnO₃Se₂: C, 21.5; H, 2.6%). ¹H NMR spectrum: δ 3.28, 3.04, 2.75 (CH₂), 2.30, 2.12, 2.02 (sh) (Me). $^{13}C-{^1H}$ NMR spectrum: δ 222.5–

218.4 (CO), 29.6, 29.4, 29.0 (CH₂), 12.6 (Me, meso-1), 11.9 and 8.9 (Me, DL).

[MnCl(CO)₃(MeSeCH₂CH₂CH₂SeMe)]. [MnCl(CO)₅] (0.100 g, 0.43 mmol) and MeSeCH₂CH₂CH₂SeMe (0.100 g, 0.43 mmol), giving a yellow precipitate (yield 0.111 g, 63%) (Found: C, 23.4; H, 3.3. Calc. for C₈H₁₂ClMnO₃Se₂: C, 23.7; H, 3.0%). ¹H NMR spectrum: δ 3.25, 3.10, 2.78, 2.65 (SeCH₂), 2.40, 2.18 (Me) and 1.98 (CH₂CH₂CH₂). $^{13}C-\{^{1}H\}$ NMR spectrum: δ 223.1–216.7 (CO), 26.9, 26.8, 25.3, 25.0, 24.5 (CH₂), 13.4, 12.6, 11.2, 10.7 (Me).

[MnCl(CO)₃(PhSeCH₂CH₂SePh)]. [MnCl(CO)₅] (0.047 g, 0.21 mmol) and PhSeCH₂CH₂SePh (0.070 g, 0.21 mmol), giving a yellow precipitate (yield 0.078 g, 75%) (Found: C, 39.4; H, 2.8. Calc. for $C_{17}H_{14}ClMnO_3Se_2$: C, 39.7; H, 2.7%). ¹H NMR spectrum: δ 7.25–7.70 (Ph), 3.25–3.78 (CH₂). ¹³C-{¹H} NMR spectrum: δ 223.2–219.6 (CO), 130.8 (sh), 130.4, 129.9 (Ph) and 30.7 (CH₂).

[MnCl(CO)₃{C₆H₄(SeMe)₂-o}]. [MnCl(CO)₅] (0.024 g, 0.100 mmol) and C₆H₄(SeMe)₂-o (0.027 g, 0.100 mmol), giving a yellow precipitate (yield 0.031 g, 68%) (Found: C, 27.1; H, 2.3. Calc. for C₁₁H₁₀ClMnO₃Se₂·CH₂Cl₂: C, 27.5; H, 2.7%). ¹H NMR spectrum: δ 7.80, 7.45 (*o*-C₆H₄), 2.70, 2.55, 2.40 (Me). ¹³C-{¹H} NMR spectrum: δ 221.7–218.8 (CO), 134.6, 133.5, 131.1, 130.5 (o-C₆H₄), 20.0 (Me, meso-2), 19.5, 15.1 (Me, DL) and 14.2 (Me, meso-1).

[MnBr(CO)₃(MeSeCH₂CH₂SeMe)]. [MnBr(CO)₅] (0.085 g, 0.31 mmol) and MeSeCH₂CH₂SeMe (0.067 g, 0.31 mmol) giving an orange precipitate (yield 0.117 g, 87%) (Found: C, 19.8; H, 2.5. Calc. for C₇H₁₀BrMnO₃Se₂: C, 19.3; H, 2.3%). ¹H NMR spectrum: δ 3.36, 3.23 (sh), 3.10, 2.82 (CH₂), 2.37 (sh), 2.35, 2.23 (Me). ${}^{13}C-{}^{1}H$ NMR spectrum: δ 124.3–116.5 (CO), 29.6, 29.3 (CH₂), 15.5, 13.4, 12.9 and 10.3 (Me).

[MnBr(CO)₃(MeSeCH₂CH₂CH₂SeMe)]. [MnBr(CO)₅] (0.011 g, 0.40 mmol) and MeSeCH₂CH₂CH₂SeMe (0.092 g, 0.40 mmol), giving an orange precipitate (yield 0.079 g, 44%) (Found: C, 21.6; H, 2.9. Calc. for C₈H₁₂BrMnO₃Se₂: C, 21.4; H, 2.7%). ¹H NMR spectrum: δ 3.42, 3.20 (sh), 2.75, 2.62 (sh) (SeCH₂), 2.28 (Me) and 1.98 (CH₂CH₂CH₂). ¹³C-{¹H} NMR spectrum: δ 223.0–216.0 (CO), 27.3, 27.2 (sh), 25.6, 24.5 (CH₂), 14.2, 13.2, 12.4, 11.5 (Me).

[MnBr(CO)₃(PhSeCH₂CH₂SePh)]. [MnBr(CO)₅] (0.072 g, 0.26 mmol) and PhSeCH₂CH₂SePh (0.089 g, 0.26 mmol), giving a yellow precipitate (yield 0.109 g, 74%) (Found: C, 35.9; H, 2.3. Calc. for C₁₇H₁₄BrMnO₃Se₂: C, 36.5; H, 2.5%). ¹H NMR spectrum: δ 7.75–7.18 (Ph), 3.75–3.30 (CH₂). ¹³C-{¹H} NMR spectrum: δ 224.5–220.1 (CO), 130.0, 129.5 (Ph) and 30.9 (CH₂).

 $[MnBr(CO)_{3}\{C_{6}H_{4}(SeMe)_{2}-o\}].$ $[MnBr(CO)_{5}]$ (0.085 g, 0.31 mmol) and C₆H₄(SeMe)₂-o (0.082 g, 0.31 mmol), giving an orange precipitate (yield 0.095 g, 64%) (Found: C, 27.2; H, 2.0. Calc. for C₁₁H₁₀BrMnO₃Se₂: C, 27.3; H, 2.1%). ¹H NMR spectrum: δ 7.85, 7.54 (*o*-C₆H₄), 2.81 (sh), 2.62 (Me). ¹³C-{¹H} NMR spectrum: δ 222.9–218.6 (CO), 134.9, 133.7, 131.1, 130.4 (o-C₆H₄), 31.0 (Me, meso-1), 21.0 (Me, meso-2), 20.4 and 17.1 (Me, DL).

[MnI(CO)₃(MeSeCH₂CH₂SeMe)]. [MnI(CO)₅] (0.119 g, 0.37 mmol) and MeSeCH₂CH₂SeMe (0.080 g, 0.37 mmol) giving an orange precipitate (yield 0.108 g, 61%) (Found: C, 17.7; H, 1.9. Calc. for C₇H₁₀IMnO₃Se₂: C, 17.4; H, 2.1%). ¹H NMR spectrum: δ 3.45, 3.12, 2.88 (CH₂), 2.50, 2.31, 2.22, 2.05 (Me). ¹³C-{¹H} NMR spectrum: δ 225.0–220.5 (CO), 30.5, 30.1 (CH₂), 15.4 (Me, meso-2), 15.3, 1.3 (Me, DL).

 Table 5
 Crystallographic data collection and refinement parameters

	[MnCl(CO) ₃ (MeSe- CH ₂ CH ₂ SeMe)]	[MnCl(CO) ₃ (MeSe- CH ₂ CH ₂ CH ₂ SeMe)]
Formula	C7H10ClMnO3Se2	C ₈ H ₁₂ ClMnO ₃ Se ₂
М	390.47	404.49
Space group	Pbca	$P2_{1}2_{1}2_{1}$
Crystal symmetry	Orthorhombic	Orthorhombic
a/Å	12.283(4)	11.342(5)
b/Å	19.293(5)	12.223(7)
c/Å	10.154(4)	9.464(6)
V/Å ³	2406(1)	1311(1)
Ζ	8	4
$D_{\rm c}/{\rm g~cm^{-3}}$	2.155	2.048
μ (Mo-K α)/cm ⁻¹	72.51	66.53
Unique observed		
reflections	2430	1360
Observed reflections		
with $[I_o > 2\sigma(I_o)]$	1206	1157
No. parameters	127	136
R	0.058	0.036
R'	0.064	0.046

[MnI(CO)₃(MeSeCH₂CH₂CH₂SeMe)]. [MnI(CO)₅] (0.124 g, 0.38 mmol) and MeSeCH₂CH₂CH₂SeMe (0.087 g, 0.38 mmol), giving an orange precipitate (yield 0.126 g, 66%) (Found: C, 19.5; H, 2.6. Calc. for $C_8H_{12}IMnO_3Se_2$: C, 19.4; H, 2.4%). ¹H NMR spectrum: δ 3.30, 2.85 (SeCH₂), 2.70, 2.40 (Me) and 2.00 (CH₂CH₂CH₂). ¹³C-{¹H} NMR spectrum: δ 226.7–217.5 (CO), 27.8, 27.5, 27.2, 26.3, 24.6, 24.3 (CH₂), 15.7, 14.5, 13.9, 12.2 (Me).

[MnI(CO)₃(PhSeCH₂CH₂SePh)]. [MnI(CO)₅] (0.063 g, 0.20 mmol) and PhSeCH₂CH₂SePh (0.067 g, 0.20 mmol), giving an orange precipitate (yield 0.096 g, 81%) (Found: C, 33.3; H, 2.4. Calc. for $C_{17}H_{14}$ IMnO₃Se₂: C, 33.7; H, 2.3%). ¹H NMR spectrum: δ 7.55–7.25 (Ph), 3.80–3.40 (CH₂). ¹³C-{¹H} NMR spectrum: δ 226.0–220.2 (CO), 131.1, 130.3, 129.8 (Ph) and 31.3 (CH₂).

[MnI(CO)₃{C₆H₄(SeMe)₂-\sigma}]. [MnI(CO)₅] (0.032 g, 0.098 mmol) and C₆H₄(SeMe)₂- σ (0.026 g, 0.098 mmol), giving an orange precipitate (yield 0.043 g, 82%) (Found: C, 21.6; H, 1.9. Calc. for C₁₁H₁₀IMnO₃Se₂·CHCl₃: C, 22.1; H, 1.7%). ¹H NMR spectrum: δ 7.90, 7.55 (σ -C₆H₄), 2.95, 2.60, 2.48 (sh), 2.35 (Me). ¹³C-{¹H} NMR spectrum: δ 223.8–219.6 (CO), 134.8, 134.1, 131.0 (σ -C₆H₄), 22.7 (Me, *meso*-2), 21.9, 20.7, 14.3 (Me, *meso*-1 and DL).

$\label{eq:crystal structures of [MnCl(CO)_3(MeSeCH_2CH_2SeMe)] and \\ [MnCl(CO)_3(MeSeCH_2CH_2CH_2SeMe)] \\$

Details of the crystallographic data collection and refinement parameters are given in Table 5. The crystals were grown by vapour diffusion of light petroleum into solutions of the complexes in CH₂Cl₂. Data collection used a Rigaku AFC7S four-circle diffractometer operating at 150 K, with graphitemonochromated Mo-K α X-radiation ($\lambda = 0.71073$ Å). No significant crystal decay or movement was observed. The structures were solved by heavy atom Patterson methods²⁵ and developed by iterative cycles of full-matrix least-squares refinement and Fourier-difference syntheses which located one complete molecule in the asymmetric unit.²² All non-H atoms were refined anisotropically while H atoms were placed in fixed, calculated positions with d(C-H) = 0.96 Å. The weighting scheme $w^{-1} = \sigma^2(F)$ gave satisfactory agreement analyses and the Flack parameter indicated that the correct enantiomorph of [MnCl(CO)₃(MeSeCH₂CH₂CH₂SeMe)] was refined.

CCDC reference number 186/1187.

Acknowledgements

We thank the EPSRC and the University of Southampton for support and the Director of the Cambridge Crystallographic Database for access.

References

- N. R. Champness, S. R. Jacob, G. Reid and C. S. Frampton, *Inorg. Chem.*, 1995, **34**, 396; N. R. Champness, S. J. A. Pope and G. Reid, *J. Chem. Soc., Dalton Trans.*, 1997, 1639; W. Levason, G. Reid and S. M. Smith, *Polyhedron*, 1997, **16**, 5253.
- 2 M. C. Durrant, S. Davies, D. L. Hughes, C. Le Floc'h, R. L. Richards, J. R. Sanders, N. R. Champness, S. J. A. Pope and G. Reid, *Inorg. Chim. Acta*, 1996, **251**, 13.
- 3 S. Davies, M. C. Durrant, D. L. Hughes, C. LeFloc'h, S. J. A. Pope, G. Reid, R. L. Richards and J. R. Sanders, J. Chem. Soc., Dalton Trans., 1998, 2191.
- 4 D. Rehder, *Multinuclear NMR*, ed. J. Mason, Plenum, New York, 1987, ch. 19.
- 5 A. Kekeci and D. Rehder, Z. Naturforsch., Teil B, 1981, 36, 20.
- 6 F. Calderazzo, E. A. C. Lucken and D. F. Williams, J. Chem. Soc. A, 1967, 154.
- 7 D. Rehder, H.-C. Bechtold, A. Kekeci, H. Schmidt and M. Z. Siewing, Z. Naturforsch., Teil B, 1982, 37, 631.
- 8 N. J. Holmes, W. Levason and M. Webster, J. Organomet. Chem., 1998, 568, 213.
- 9 A. M. Hill, W. Levason, M. Webster and I. Albers, *Organometallics*, 1997, 16, 5641.
- 10 J. Connolly, G. W. Goodban, G. Reid and A. M. Z. Slawin, J. Chem. Soc., Dalton Trans., 1998, 2125.
- 11 S. J. A. Pope and G. Reid, unpublished work.
- 12 E. W. Abel and G. V. Hutson, J. Inorg. Nucl. Chem., 1969, **31**, 3333.
- 13 D. Rehder, K. Ihmels, D. Wenke and P. Oltmanns, *Inorg. Chim. Acta*, 1985, **100**, L13.
- 14 A. Belforte, F. Calderazzo, D. Vitali and P. F. Zanazzi, Gazz. Chim. Ital., 1985, 115, 125.
- 15 E. W. Abel, S. K. Bhargava and K. G. Orrell, Prog. Inorg. Chem., 1984, 33, 1.
- 16 E. G. Hope and W. Levason, Coord. Chem. Rev., 1993, 122, 109.
- 17 H. Schumann, A. M. Arif, A. L. Rheingold, C. Janiak, R. Hoffmann and N. Kuhn, *Inorg. Chem.*, 1991, 30, 1618.
- 18 E. W. Abel, S. K. Bhargava, M. M. Bhatti, K. Kite, M. A. Mazid, K. G. Orrell, V. Sik, B. L. Williams, M. B. Hursthouse and K. M. A. Malik, J. Chem. Soc., Dalton Trans., 1982, 2065.
- 19 E. W. Abel, M. M. Bhatti, K. G. Orrell and V. Sik, J. Organomet. Chem., 1981, 208, 195.
- 20 E. W. Abel, S. K. Bhargava, M. M. Bhatti, M. A. Mazid, K. G. Orrell, V. Sik, M. B. Hursthouse and K. M. A. Malik, J. Organomet. Chem., 1983, 250, 373.
- 21 SHELXS 86, Program for crystal structure solution, G. M. Sheldrick, *Acta Crystallogr., Sect. A*, 1990, **46**, 467.
- 22 TEXSAN, Crystal Structure Analysis Package, Molecular Structure Corporation, Houston, TX, 1995.
- 23 K. J. Reimer and A. Shaver, Inorg. Synth., 1979, 19, 159.
- 24 D. J. Gulliver, E. G. Hope, W. Levason, S. G. Murray, D. M. Potter and G. L. Marshall, J. Chem. Soc., Perkin Trans. 2, 1984, 429.
- 25 PATTY, The DIRDIF Program System, P. T. Beurskens, G. Admiraal, G. Beurskens, W. P. Bosman, S. Garcia-Granda, R. O. Gould, J. M. M. Smits and C. Smykalla, Technical Report of the Crystallography Laboratory, University of Nijmegen, 1992.

Paper 8/06881J