## Cyclopentadienylmanganese Dicarbonyl Thiocarbonyl

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Summary The thiocarbonyl complex, C<sub>5</sub>H<sub>5</sub>Mn(CO)<sub>2</sub>(CS), has been isolated from the reaction of C<sub>5</sub>H<sub>5</sub>Mn(CO)<sub>2</sub>-(cis-cyclo-octene) with carbon disulphide.

FOLLOWING the discovery in 1966 of the first thiocarbonyl complex of a transition metal,1 a molecular orbital calculation indicated2 that transition-metal thiocarbonyl complexes should be more stable than their carbonyl analogues. Despite this, surprisingly few thiocarbonyl complexes have been reported and then only for the Group VIII metals, rhodium,1-4 irridium,4 cobalt,5 iron,6 and ruthenium.7

We have been attempting to extend the range of the known thiocarbonyl complexes and have succeeded in isolating in low yield the first thiocarbonyl complex of manganese, C<sub>5</sub>H<sub>5</sub>Mn(CO)<sub>2</sub>(CS) (I), from the reaction of  $C_5H_5Mn(CO)_2(cis$ -cyclo-octene) with carbon disulphide. The reaction is complex owing to the concomitant formation of C5H5Mn(CO)3 and an, as yet, unidentified compound. The new compound, (I), forms fairly air stable, pale-green crystals, m.p. 53°. It analyses correctly and from its mass spectrum it has the correct molecular weight and the expected fragmentation pattern. The i.r. spectrum exhibits two strong terminal C-O stretching absorptions and one strong terminal C-S stretching absorption (Table). These frequencies may be compared with those also shown in the Table for the isoelectronic and presumably isostructural cationic iron thiocarbonyl complex, [C<sub>5</sub>H<sub>5</sub>Fe(CO)<sub>2</sub>(CS)]<sup>+</sup>. The absence of the positive charge on (I) would be expected to result in an increase in the extent of back-bonding between the metal and the CO and CS groups when compared to that for the iron complex. Consequently, the C-O and C-S stretching frequencies of (I) would be expected to be lower than those of the iron complex, as is observed.

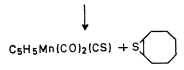
I.r. spectra (cm<sup>-1</sup>) of cyclopentadienylmetal dicarbonyl thiocarbonyl complexes in Nujol mull

Compound		νco		VC8
$C_5H_5Mn(CO)_2(CS)^3$		2010s	1959s	1271s
$[C_5H_5Fe(CO)_2(CS)]PF_6$ b		2093s	2064s	1348s

- a In CS<sub>2</sub> solution: ν<sub>CO</sub> 2006s 1954s; ν<sub>CS</sub> 1266s.
- b From ref. 6a.

The formation of (I) from C<sub>5</sub>H<sub>5</sub>Mn(CO)<sub>2</sub>(cis-cyclooctene) and CS2 is a novel reaction as it provides the first example of the substitution of a co-ordinated olefin by the two-electron ligand CS. Most probably the olefin acts as a sulphur acceptor in the reaction and in doing so it is converted into an episulphide

$$C_5H_5Mn(CO)_2(C_8H_{14}) + CS_2$$



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