Synthesis and structural characterisation of two $\eta^{\text{1}}\text{-bonded}$ N-phenylthioformamidate complexes of rhodium

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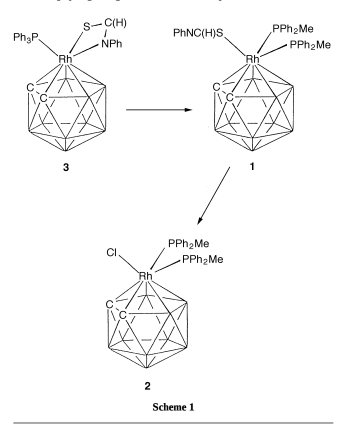
The complexes $[3-\{\eta^1-SC(H)NPh\}-3,3-(PMe_2Ph)_2-3,1,2-{\it closo-RhC}_2B_9H_{11}]$ and $[2-\{\eta^1-SC(H)NPh\}-2,2-(PMe_2Ph)_2-2,1-{\it closo-RhTeB}_{10}H_{10}]$ have been structurally characterised using X-ray crystallography and are the first η^1 -bonded thioformamidate complexes to be isolated.

Numerous metal complexes of ligands containing the S-C-N bond sequence have been characterised in the solid state. Reactions between isothiocyanates, RNCS, and metal hydrides usually lead to metal complexes with $[\eta^2-SC(H)NR]^-$ ligands which are bonded through both M-S and M-N bonds. A typical example is $[ZrCl{\eta^2-SC(H)NPh}(cp)_2]$ (cp = $\eta^5-C_5H_5$). Alternatively, more complex ligands derived from several SCNR molecules may be formed such as, $[\eta^2-S_2C(H)NR]^-$ in $[2-\{\eta^2-S_2C(H)NR]^-]$ S₂CN(H)Ph}-2-(PPh₃)-2,1-closo-RhTeB₁₀H₁₀].² Although there have been several reports of complexes containing monodentate ligands derived from pyridine-2-thiol and related compounds (see ref. 3 for a recent review), there has been no report to our knowledge of the structural characterisation of any complex with an η^1 -thioformamidato-to-metal functionality. Recently we proposed an $[\eta^1-SC(H)NPh]^-$ complex, $[3-\{\eta^1-\xi^2\}]$ SC(H)NPh}-3,3- $(PPh_2Me)_2$ -3,1,2-closo- $RhC_2B_9H_{11}$] 1, as an intermediate in the synthesis of the rhodacarborane [3,3- $(PPh_{2}Me)_{2}\text{--}3\text{--}Cl\text{--}3,1,2\text{--}\textit{closo}\text{--}RhC_{2}B_{9}H_{11}]~\textbf{2},~from~[3\text{--}\{\eta^{2}\text{--}SC(H)\text{--}H_{11}\}]$ NPh}-3-(PPh₃)-3,1,2-closo-RhC₂B₉H₁₁] **3**, Scheme 1.⁴ A study of the parallel reaction with PMe2Ph as the phosphine has afforded $[3-{\eta^1-SC(H)NPh}-3,3-(PMe_2Ph)_2-3,1,2-closo-RhC_2 B_9H_{11}$] 4, Fig. 1. This compound was isolated from the reaction between a ten-fold excess of PMe₂Ph (0.109 g, 0.790 mmol) and a solution of 3 (0.050 g, 0.079 mmol) in \tilde{CH}_2Cl_2 (20 cm³) at room temperature (r.t.) for 30 min. After evaporating the solvent under reduced pressure, the residue was washed with hexane $(3 \times 5 \text{ cm}^3)$ to remove the excess phosphine. The single product was recrystallised from CH₂Cl₂-hexane solution affording orange crystals of $[3-\{\eta^1-SC(H)NPh\}-3,3-(PMe_2Ph)_2-3,1,2$ closo-RhC₂B₉H₁₁] $4\cdot 0.93$ CH₂Cl₂, in 83% yield (0.047 g).† An analogous reaction (r.t., 15 min) with the rhodatelluraborane SC(H)NPh}-2-(PPh₃)-closo-2,1-RhTeB₁₀H₁₀] **5**, produced [2- $\{\eta^1\text{-SC(H)NPh}\}$ -2,2- $(PMe_2Ph)_2$ -2,1-closo-RhTeB₁₀H₁₀] **6**, Fig. 2. The rhodatelluraborane 6 was recrystallised from a CH₂Cl₂hexane solution in a yield of 86%.‡ Satisfactory microanalytic data (C, H, N) were obtained for both 4 and 6.

Both compounds 4 and 6 contain rhodium-sulfur bonded η¹-

† Crystal data for **4**. C₂₅H₃₉B₉NP₂RhS·0.93CH₂Cl₂, orange platelet, 0.39 × 0.26 × 0.12 mm, M= 726.80, monoclinic, $P2_1/c$, a = 10.0955(12), b = 21.897(2), c = 15.960(2) Å, β = 96.599(9)°, U= 3504.6(6) ų, Z= 4, D_c = 1.39 g cm⁻³, λ (Mo-K α) = 0.7107 Å, μ (Mo-K α) = 0.81 mm⁻¹, F(000) = 1484, T= 294 K. Data for 8046 reflections were measured, of which 7625 reflections were unique ($R_{\rm int}$ = 0.009) and of these the 5575 with I > 2 σ (I) were labelled 'observed'. $R(F_o)$ = 0.0348, R'(F^z) = 0.0885 for all measured data where $R(F_o)$ = $\Sigma ||F_o| - |F_c|| / \Sigma |F_o|$, R'(F^z) = $\{\Sigma [w(F_o^2 - F_c^2)^2] / \Sigma (wF_o^2)\}^{\frac{1}{2}}$, and w = 1/[σ^2 (F_o^2)].

SC(H)NPh ligands. However, the orientation of these ligands with respect to the RhC_2B_3 or $RhTeB_4$ moieties in each of the compounds is clearly different. The phenylthioformamidate group in compound 4 interacts solely with the rhodium atom of the RhC_2B_9 cage, Fig. 1, whereas in compound 6 there is also a



‡ Crystal data for **6**. $C_{23}H_{38}B_{10}NP_2RhSTe$, orange needle, $0.40\times0.20\times0.20$ mm, M=761.15, triclinic, $P\bar{1}$, a=9.7040(13), b=11.895(2), c=14.4469(15) Å, $\alpha=76.211(11)$, $\beta=80.557(11)$, $\gamma=83.680(12)^\circ$, U=1593.2(3) ų, Z=2, $D_c=1.587$ g cm³, $\lambda(Mo-K\alpha)=0.7107$ Å, $\mu(Mo-K\alpha)=1.616$ mm¹, F(000)=752, T=294(1) K. Data for 5450 reflections were collected and of these the 4402 with $I>2\sigma(I)$ were labelled 'observed'. $R(F_o)=0.0292$, $R'(F^2)=0.0800$ for all measured data, $R(F_o)$ and $R'(F^2)$ as for **4**.

Structure solution of **4** and **6**. Data were collected using an Enraf-Nonius CAD4 diffractometer to a maximum θ of 27° using graphite-monochromated Mo-Ka radiation. Data were corrected for Lorentz, polarisation and absorption effects (from ψ scans). The structures were solved by Patterson and Fourier methods and refined by full-matrix least-squares calculations initially using the NRCVAX system of programs ⁵ and finally with SHELXL 93^{6} using all F^{2} data. The H atoms were allowed for as riding atoms using the appropriate AFIX commands in SHELX 93. Diagrams were prepared with the aid of ORTEP ⁷ and PLATON. ⁸ Atomic coordinates, thermal parameters, and bond lengths and angles have been deposited at the Cambridge Crystallographic Data Centre (CCDC). See Instructions for Authors, *J. Chem. Soc., Dalton Trans.*, 1997, Issue 1. Any request to the CCDC for this material should quote the full literature citation and the reference number 186/334.

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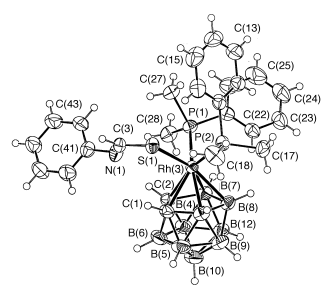


Fig. 1 An ORTEP view of compound 4 with the atom numbering scheme. Displacement ellipsoids are at the 50% level except for H atoms which are drawn as small spheres of an arbitrary size. Selected interatomic distances (Å) and angles (°): Rh(3)–S(1) 2.4010(8), Rh(3)–C(1) 2.246(3), Rh(3)–C(2) 2.212(3), Rh(3)–B(4) 2.280(4), Rh(3)–B(7) 2.216(3), Rh(3)–B(8) 2.274(3), Rh(3)–P(1) 2.3242(10), Rh(3)–P(2) 2.3346(8), C(1)–C(2) 1.613(4), S(1)–C(3) 1.724(3), C(3)–N(1) 1.254(4), N(1)–C(41) 1.424(4), C–B distances range from C(1)–B(6) 1.674(5) to C(2)–B(7) 1.753(4) and B–B distances from B(5)–B(6) 1.752(7) to B(4)–B(8) 1.818(5); C(3)–S(1)–Rh(3) 109.31(11), N(1)–C(3)–S(1) 126.7(2), C(41)–N(1)–C(3) 119.7(3), S(1)–Rh(3)–P(1) 82.11(3), S(1)–Rh(3)–P(2) 91.51(3), P(1)–Rh(3)–P(2) 95.83(3), S(1)–Rh(3)–C(1) 85.64(8), S(1)–Rh(3)–C(2) 103.55(8)

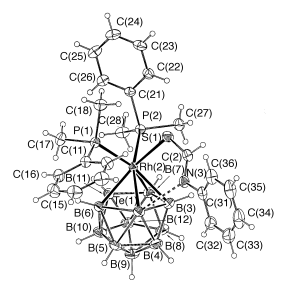


Fig. 2 An ORTEP view of compound **6** with the atom numbering scheme. Displacement ellipsoids are at the 30% level except for H atoms which are drawn as small spheres of an arbitrary size. Selected interatomic distances (Å) and angles (°): Rh(2)–S(1) 2.4147(10), Rh(2)–Te(1) 2.5788(4), Rh(2)–B(3) 2.336(4), Rh(2)–B(6) 2.362(4), Rh(2)–B(7) 2.241(4), Rh(2)–B(11) 2.253(4), Rh(2)–P(1) 2.3732(10), Rh(2)–P(2) 2.3601(10), S(1)–C(2) 1.712(4), C(2)–N(3) 1.256(5), N(3)–C(31) 1.423(5), Te(1) ··· N(3) 2.737(3), Te-B distances range from Te(1)–B(4) 2.288(5) to Te(1)–B(3) 2.390(5) and B–B distances from B(10)–B(12) 1.745(7) to B(5)–B(6) 1.896(6); C(2)–S(1)–Rh(2) 114.83(14), N(3)–C(2)–S(1) 127.7(3), C(31)–N(3)–C(2) 120.1(3), S(1)–Rh(2)–P(1) 87.41(4), S(1)–Rh(2)–P(2) 81.74(4), P(1)–Rh(2)–P(2) 96.58(3), S(1)–Rh(2)–Te(1) 93.27(3)

weak but significant intramolecular $Te\cdots N$ contact, Fig. 2. The S(1)-C(3)-N(1)-C(41) plane in $\bf 4$ is at an angle of $37.5(3)^\circ$ to the best-fit plane containing the C_2B_3 face, while the S(1)-C(2)-N(1)-C(31) plane in $\bf 6$ is at an angle of $60.8(3)^\circ$ to the TeB_4 face. The SC(H)NPh ligand in $\bf 6$ is clearly positioned to facilitate the $Te\cdots N$ interaction.

Within each SC(H)NPh ligand, the bond lengths and most of the bond angles are essentially the same. In both cases the S-C-N-C atoms are virtually coplanar with torsion angles of $177.5(3)^{\circ}$ in compound **4** and $178.3(3)^{\circ}$ in **6**. The S(1)-C(3)-N(1) and C(3)-N(1)-C(41) angles in 4 are 126.7(2) and 119.7(3)°, while the corresponding angles in 6 are 127.7(3) and 120.1(3)°. The S-C distances of 1.724(3) and 1.712(4) Å respectively in 4 and 6 are typical of delocalised sp² hybridised carbon-sulfur bonds (1.720 Å), i.e. longer than the typical $S=C_{sp^2}$ distance of 1.681 Å in thioureas and shorter than the typical S-C_{sp3} distance of 1.808 Å in thiols.⁵ The phenyl carbonto-nitrogen and the methine carbon-to-nitrogen distances are respectively 1.424(4) and 1.254(4) Å in 4 and 1.423(5) and 1.256(5) Å in 6, and are essentially identical. These bond lengths are respectively longer than typical N-C_{ar} bonds and shorter than typical N=C_{sp²} bonds.9

The Te \cdots N interaction in compound **6** has implications for cluster electron counting. Although the Te···N distance in 6 is long, 2.737(3) A compared with a typical Te-N distance (in covalent bonds 2.15 Å), it is considerably shorter than the sum of the van der Waals' radii of Te and N, 3.61 Å. Similar Te · · · N distances of 2.702 and 2.752 Å respectively have previously been reported in the compounds bis[2-(4'methoxyphenyl)iminomethinylphenyl]telluride 10 and 1,6-bis[(2butyltelluro)phenyl]-2,5-diazahexa-1,5-diene.11 If the donation of electron density from the nitrogen lone pair to the tellurium in the RhTeB₁₀ cage was strong it would imply an electronic character for 6 which is nido-type, but because the Te···N interaction is weak, a nido structure for 6 is not observed and the closo structure of the system is maintained. It is noteworthy, however, that the Rh-S bond length in 6, 2.4147(10) Å, is significantly longer than that in 4, 2.4010(8) Å. In overall electron density terms, the relative weakening of the Rh-S bond in 6 may be considered to balance the Te···N interaction. The rhodium-tellurium distance in 6, 2.5788(4) Å, is well within the known range of 2.529(4) 12 to 2.6172(4) Å 13 for rhodatelluraboranes and is close to the distance found in [2-{n²-S₂CN-(H)Ph}-2-(PPh₃)-closo-2,1-RhTeB₁₀H₁₀] [2.5812(3) Å].² The dimensions of the RhC₂B₉H₁₁ and RhTeB₁₀H₁₀ cages, Figs. 1 and 2 respectively, are typical of such structures and require no further comment.¹⁴

Finally, with respect to the previous suggestion that $[3-\{\eta^1-SC(H)NPh\}-3,3-(PPh_2Me)_2-3,1,2-closo-RhC_2B_9H_{11}]$ **1** is an intermediate in the formation of $[3,3-(PPh_2Me)_2-3-Cl-3,1,2-closo-RhC_2B_9H_{11}]$ **2** from $[3-\{\eta^2-SC(H)NPh\}-3-(PPh_3)-3,1,2-closo-RhC_2B_9H_{11}]$ **3**,⁴ Scheme 1, we wish to report that the rhodacarborane **1** has now been isolated from this reaction in 70% yield and characterised spectroscopically. The formation of the rhodium–chloride containing compounds $[3-Cl-3,3-(PPh_2Me)_2-3,1,2-closo-RhC_2B_9H_{11}]$ **2** (quantitative yield) or $[3-Cl-3,3-(PMe_2Ph)_2-3,1,2-closo-RhC_2B_9H_{11}]$ **7** (16% yield) is observed when the complexes **1** and **4** respectively are refluxed in CH_2Cl_2 solution for **48** h.

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