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TRIHYDROCARBYLTIN COMPLEXES OF TRANSITION METAL CYANIDES

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Summary

Cationic single-bridged trinuclear cyano complexes are obtained by treating trihydrocarbyltin perchlorate $(SnR_3O_2ClO_2)$ (R = alkyl or aryl) with neutral transition metal monocyanides, whereas treatment with the corresponding neutral or anionic transition metal polycyanides gives neutral single-bridged polymeric cyano compounds. In every case the tin atoms are pentacoordinated with the three R groups in the equatorial and two CN groups (or, respectively one CN and one $OClO_3$ group) in the axial position of a trigonal bipyramid.

Introduction

The synthesis of covalent perchlorato derivatives from the corresponding halo complexes, according to eq. 1 and the subsequent substitution of the perchlorato

$$L_nMX + AgClO_4 \rightarrow AgX + L_nMOClO_3$$
 (1)

group, (which has a low coordinating capacity) by neutral (eq. 2) or anionic ligands (eq. 3) has frequently been used by us [1] and by others [2] for the

$$L_n MOClO_3 + L' \rightarrow [L_n ML']ClO_4$$
 (2)

$$L_nMOClO_3 + M'X \rightarrow M'ClO_4 + L_nMX$$
 (3)

preparation of neutral and anionic organometallic complexes of gold [1a-1c], nickel [2a-2c], palladium [1d-1g,2d], platinum [1h-1j] and rhodium [1k-1l].

If reaction 1 is carried out in a non-donor solvent under anhydrous conditions the intermediate perchlorato complex can sometimes be isolated, but is frequently unstable. Even so, after removal of the precipitated silver halide, the filtrate can in every case be used for processes 2 and 3.

Mainly because of the solubility of L_nMX it is, however, sometimes necessary to work in another type of solvent. In these cases it seems appropriate to employ

acetone, which, though it displaces the perchlorato group giving a (sometimes isolable) cationic perchlorate (eq. 4), can afterwards readily be displaced by

$$L_n MX + AgClO_4 \xrightarrow{Me_2CO} AgX + [L_n M(Me_2CO)_x]ClO_4$$
 (4)

reactions of type 2 or 3.

We consider below the reaction of solutions of perchloratotrialkyltin or perchloratotriaryltin compounds with several cyano complexes of transition metals. These behave as a ligand L' (eq. 2) since their cyano group still has a potentially N-donor atom which causes the displacement of the perchlorato group or the acetone. The process allows the preparation of polymeric (finite or infinite) complexes with the cyano group bridging the tin and the transition metal atoms.

Some of these results were the subject of a preliminary communication [3].

Results and discussion

The precursor solutions used in the present study were prepared by treating the organochlorocomplexes SnR₃Cl (R being Me, Bu or Ph) with benzene or acetone solutions of AgClO₄, and removal of the AgCl. As reported [4] for the methyl complex [SnMe₃O₂ClO₂]_x in the solid state the pentacoordinated tin atoms are bridged by the perchlorato ligands.

This bridging system is cleaved in methanol to give the pentacoordinated cationic perchlorate $[SnR_3S_2]ClO_4$ (S = solvent). Thus, whilst our benzene solutions are likely to contain $[Sn(O_2ClO_2)R_3]$, the acetone solutions are assumed to contain $[SnR_3(Me_2CO)_2]ClO_4$, but the reaction products with cyano-transition metal complexes are in each case the same.

(a) Reactions with monocyanides

2
$$Pd(CN)(C_6F_5)(PPh_3)_2 + [Sn(O_2ClO_2)R_3]$$

$$\begin{bmatrix}
PPh_3 & R & PPh_3 \\
C_6F_5 & Pd & CN & Sn & NC & Pd & C_6F_5
\end{bmatrix} ClO_4 (5)$$

$$C_6F_5 & Pd & CN & Sn & PPh_3 \\
PPh_3 & R & PPh_3
\end{bmatrix} ClO_4 (5)$$

$$(I,R = Me;$$

$$II,R = Bu;$$

$$II,R = Ph)$$

The reaction (1/2) of benzene or acetone solutions of perchlorato trihydro-carbyltin $Sn(O_2ClO_2)R_3$ with cyanopen afluorophenylbis (triphenylphosphine)-palladium(II) $[Pd(CN)(C_6F_5)(PPh_3)_2]$ gives cationic trinuclear complexes (see eq. 5). The complexes I and III are precipitated instantaneously whatever the used molar ratio of the reagents used but the butyl complex (II) can only be obtained by use of a 2/1 mole ratio as in eq. 5. A 1/1 mole ratio leads to the

precipitation of the neutral complex IV, (see eq. 6), which contains a monodentate $(C_{3\nu})$ perchlorato group (see below).

The tendency of tin to adopt a coordination number of 5 means that the reaction of $Sn(O_2ClO_2)R_3$ with di- or poly-cyano-transition metal complexes leads to polymeric end products. Thus the reaction with neutral polycyano complexes yields cationic polymers, whereas anionic polycyanides give neutral polymers.

(b) Reactions with di- and poly-cyanides

The reaction with trans-dicyanobis(triphenylphosphine)palladium(II), according to eq. 7 gives rise to the immediate precipitation of the cationic

$$\left[Pd(CN)_{2}(PPh_{3})_{2}\right] + \left[Sn(O_{2}ClO_{2})Ph_{3}\right] - \left[Sn(O_{2}ClO$$

polynuclear complex V. The reactions with other cationic dicyano complexes, such as $(Bu_4N)[Ag(CN)_2]$ or $K[Au(CN)_2]$ (which are very little soluble in benzene require the use of acetone $(Me_2CO = S)$ as reaction medium) also give neutral complexes (eq. 8).

$$Q[M(CN)_{2}] + [SnR_{3}S_{2}]ClO_{4} \rightarrow QClO_{4} + 2 S + \begin{bmatrix} R \\ Sn-NC-M-CN \end{bmatrix}_{x}$$

$$(VI, M = Ag, Q = Bu_{4}N, R = Me; VII, M = Ag, Q = Bu_{4}N, R = Ph; VIII, M = Au, Q = K, R = Me; IX, M = Au, Q = K, R = Bu; X, M = Au, Q = K, R = Ph)$$

$$(8)$$

The complexes separate out upon mixing the two solutions. The simultaneously

precipitated QClO₄ can be removed by washing with water. No reaction could be observed for M = Ag and R = Bu.

Complex V is decomposed by an acetone solution of NaBPh₄ finally giving trans-Pd(CN)₂(PPh₃)₂. Treatment with tetra- and hexa-cyanometallates gives immediate precipitation of the corresponding neutral polymers.

$$K_{2}[M(CN)_{4}] + 2[SnR_{3}S_{2}]ClO_{4} \rightarrow 2 \ KClO_{4} + 4 \ S + \{(R_{3}Sn)_{2}[(\mu\text{-}CN)_{4}]\}_{x}$$
 (9)
(XI, M = Pd, R = Me;
XII, M = Pd, R = Bu;
XIII, M = Pd, R = Ph;
XIV, M = Pt, R = Me;
XV, M = Pt, R = Bu;
XVI, M = Pt, R = Ph)

In the light of the square planar coordination of Pd or Pt, these polymers can be expected to have an infinite twodimensional structure (Fig. 1).

When $K_3[Fe(CN)_6]$ is used no reaction occurs for R = Me, whereas when R = Bu or Ph it gives neutral polymeric complexes (eq. 10), which are assumed to

$$K_3[Fe(CN)_6] + 3[SnR_3S_2]ClO_4 \rightarrow 3 KClO_4 + 6 S + \{[R_3Sn]_3[(\mu-CN)_6Fe]\}_x$$
 (10)

have a threedimensional structure.

(c) Solubility and conductivity

As expected for macromolecular substances, all the polymeric complexes (V—XVIII) are insoluble in water and organic solvents. The bi- and tri-nuclear complexes (I—IV) are soluble in chloroform, dichloromethane and nitromethane, and complexes II—IV are also soluble in acetone. All are insoluble in n-hexane

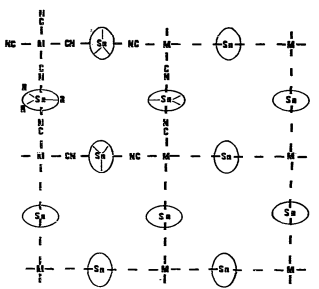


Fig. 1. Proposed structure of the $\{[R_3Sn]_2[(\mu-NC)_4M]\}_r$ complexes. (M = Pd or Pt; R = Me, Bu, Ph).

and benzene. The conductivities of the soluble complexes show [5] that they behave as 1/1 electrolytes, in accordance with the proposed formulae.

(d) IR spectra

The discussion above requires that the reaction products contain only bridging CN groups while the starting materials contain only terminal CN groups. In accordance with this, the $\nu(\text{CN})$ stretching vibration of the resulting complexes is always shifted towards higher energy, and only a single band is observed [6]. Table 1 shows the values of $\Delta\nu$ ($\Delta\nu = \nu(\text{CN})_{\text{bridge}} - \nu(\text{CN})_{\text{terminal}}$) for all the new complexes.

All the pentafluorophenyl derivatives exhibit absorption at \sim 1500vs, 1050s, 950vs and 800s cm⁻¹ due to the C₆F₅ ligand [7] along with those of the two mutually *trans*-PPh₃ groups [8] at 525s, 515s and 502s cm⁻¹ (complexes I—IV).

All the methyl derivatives show a strong absorption at 560-555 cm⁻¹ assignable to the $\nu_{\rm as}(\rm Sn-C)$ stretching mode, and only complex XIV also shows a very weak band at 505 cm⁻¹ assignable to $\nu_{\rm asym}(\rm Sn-C)$. This is in good agreement with the proposed trigonal bipyramidal structure, with the three methyl groups in equatorial position and the two cyano groups at axial sites. This agrees with the structure proposed for other trimethyl derivatives of $\rm SnR_3X$, with X being a bridging ligand (X = F [9]; Cl, Br [10]; ClO₄ [4,11]; BF₄ [12], AsF₆, SbF₆ [13]; RCOO [14] or alkyl- and aryl-imidazole [15]).

The corresponding absorptions for the butyl derivatives located in the 650–600 cm⁻¹ region, which are always weak [16] are in our complexes masked by bands due to the PPh₃, ClO₄ or CN group. Our phenyl derivatives show two medium bands at 280–270 and 240–230 cm⁻¹, assignable to symmetric and assymetric $\nu(\text{Sn-C})$ [17], showing that the Sn-C bonds are not coplanar.

Finally, all the ionic complexes (I, II, III and V) show absorptions at 1100s and 630m cm⁻¹ assignable, respectively, to v_3 or v_4 vibrations of the perchlorato group (T_d). In complex IV the bands expected at 1100 and 630 cm⁻¹ are split, and are located, respectively, at 1140 and 1040 cm⁻¹ and at 615 and 625 cm⁻¹, corresponding to the v_1 , v_4 and v_3 , v_5 vibrations of the OClO₃ group (C_{3v}). This complex shows another band (A_1) at 915 cm⁻¹ assignable to v(Cl-O), which would be IR active for C_{3v} symmetry, and this confirms the formulation proposed for this compound [18].

Experimental

C, H and N analyses were carried out with a Perkin—Elmer 240 microanalyzer. Melting points were measured on a Buchi apparatus (model Dr. Tottoli) and are uncorrected. Conductivities were determined in approximately $5 \times 10^{-4} M$ solutions with a Philips PW 9501/01 conductimeter. IR spectra were recorded (over the range $4000-200 \text{ cm}^{-1}$) on a Perkin—Elmer 599 spectrophotometer using Nujol mulls between polyethylene plates.

 $Pd(CN)(C_6F_5)(PPh_3)_2$ was prepared as described elsewhere [19]. $Pd(CN)_2$ -(PPh_3)₂ was obtained by treating a solution of $PdI_2(PPh_3)_2$ (9.01 g, 10 mmol) in 2C0 ml of acetone with freshly prepared AgCN (2.60 g, 20 mmol) for 60 min at room temperature with exclusion of light. The AgI was filtered off and

ANALYTICAL AND OTHER DATA FOR POLYNUCLEAR COMPLEXES

TABLE 1

Complex		Foun (calcd.) %	led.) %		Λ _M (ohm ⁻¹	v(CN)	νΔ ا	M.p.
		Ć	н	z		(cm ⁻¹)	C may	3
1	{[Me3Sn](µ-NC)2[Pd(C6F5)(PPh3)2]2}C1O4	64.75	3.76	1,45	116 b	21.77	47	154(dec.)
Ħ	{ Bu3Sn](u-NC)2 Pd(C6F5)(PPh3)2 2 }ClO4	(55.93) 57.62	(3.64) 4,31	(1.47) 1.41	112 b	2155	26	195 (dec.)
,		(57.77)	(4.30)	(1.37)				
ш	{[Ph3Sn](µ-NC)2[Pd(C6F5)(PPh3)2]2}2}2C!O4	58.69	3,60	1.35	62.9	2155	26	179 (dec.)
ΙΛ	[Bu ₃ (O ₃ ClO)Sn](\(\mu\)-NC)[Pd(C ₆ F ₅)(PPh ₃) ₂]	(09,00) 74,46	(3.60) 4.85 3.85	(1.34) 1.16	130 p	21.55	25	203 (dec.)
		(64.44)	(4,73)	(1.16)				
>	$\{[Ph_3Sn](\mu\text{-NC})[Pd(PPh_3)_2](\mu\text{-CN})\}_{\mathcal{K}}(ClO_4)_{\mathcal{K}}$	59,11	4.04	2.59	೮	2150	30	192 (dec.)
ΙΛ	{[Me ₃ Sn](u-NC)[Ag](u-CN)}	(59.39) 17.91	(4.01) 2.59	(2.47) 8.58		2166	26	263
		(18.55)	(3,80)	(8.65)) 		
VII	$\{[Ph_3Sn](\mu \cdot NC)[Ag](\mu \cdot CN)\}_x$	47.02	2.92	5.13		2160	20	>300
		(47.11)	(2.96)	(6,49)				
VIII	$\{\text{IMe}_3\text{Sn}\}(\mu\cdot\text{NC})[\text{Au}](\mu\cdot\text{CN})\}_{x}$	14.55	2.14	6.74		2171	26	215 (dec.)
		(14.65)	(2.20)	((6.79)				
×	$\{[Bu_3Sn](\mu-NC)[Au](\mu-CN)\}_{x}$	31.38	5,03	60'9		2170	26	265 (dec.)
	[]	(31.19)	(5.05)	(6.20)		;	;	,
×	$\{[Pn_3sn](\mu-NC)[Au](\mu-CN)\}_{x}$	40.59	2,65	4.82		2170	52	220 (dec.)
		(40.10)	(2,62)	(4.68)			;	•
×	$\{[Me_3Sn]_2[(\mu-NC)_4Pd]\}_x$	22.36	3.25	10.24		2165	23	243 (dec.)
;		(22.32)	(3.27)	(10,41)		. !	,	
TIX	Lbu3snj21(4-NC)4PdJjx	42.30	6.69	86,9 96,1		2165	2	287 (dec.)
. !	((42.64)	(6.89)	(40.7)		1	;	:
XIII	$[[Ph_3Sh_2](\mu \cdot NC)_4 \cdot d_1]_x$	52.75	38.80	5.77		2165	2	210 (dec.)
V17.	[[Max Gm1_[[M], Dit]]	(02.77)	(3.32)	(0.10) 0.08		270	00	
	H(T, +1/), - L(1/17, -1/17)	(19.26)	(2,89)	(8.94)			3	
ΧV	{[Bu3Sn]2[(u-NC)4Pt]}.	38,78	6.27	6,05		2160	27	>300
		(38.25)	(6.19)	(6.37)				
XVI	$\{[Ph_3Sn]_2[(\mu \cdot NC)_4Pt]\}_x$	48.62	3.21	4.76		2160	27	>300
		(48.08)	(3.03)	(5,61)				
XVII	$\{\text{IBu}_3\text{Sn}\}_3[(\mu\text{-NC})_6\text{Fo}]\}_{x}$	45.61	7.07	7.78		2146	8 8	182 (dec.)
	•	(46.22)	(7.55)	(7.77)				
XVIII	{[Ph3Sn]3[(u·NC)6Fe]}x	56.46	3.59	7,16		2140	22	163 (dec.)
		(67.11)	(3.59)	(8.66)				
	فقيا فيسبون كالقراب فيفيد والباط ويترقف فيستوجون أسيدو ففيت ووهانا فيوورن أسيدون فالمنازي فقداني والمقارف							

a In nitromethane. b In acetone. c Insoluble.

the filtrate evaporated to dryness. The residue was recrystallized from dichloromethane/ethanol.

 $K[Au(CN)_2]$, $(BuN_4)[Ag(CN)_2]$, $K_2[Pd(CN)_4]$ and $K_2[Pt(CN)_4]$ were prepared by standard procedures [20,21]. $K_3[Fe(CN)_6]$ was acquired from commercial sources.

(i) Preparation of the organotin perchlorate complexes $[Sn(O_2ClO_2)R_3]$ (R = Me, Bu, Ph)

Depending on their subsequent use these complexes were prepared either in benzene or acetone by treating stoichiometric amounts of the organotin complex SnR₃Cl with AgClO₄ for 30 min at room temperature. After removal of the AgCl the filtrate was used for further reactions without additional treatment.

- (ii) $\{[R_3Sn](\mu\text{-}CN)_2[Pd(C_6F_5)(PPh_3)_2]_2\}CIO_4$ (I, R=Me; II, R=Bu, III, R=Ph) Addition of $Sn(O_2CIO_2)R_3$ (0.25 mmol) in 40 ml of benzene to a stirred solution of $Pd(CN)(C_6F_5)(PPh_3)_2$ (0.4120 g, 0.5 mmol) in 40 ml of benzene immediately gave a white microcrystalline precipitate which was filtered off and repeatedly washed with hexane (yields: Complex I 84%, complex II 75%, complex III (61%).
- (iii) $\{[Bu_3(O_3ClO)Sn](\mu-NC)[Pd(C_6F_5)(PPh_3)_2]\}$ (IV) Complex IV was similarly prepared but using a different molar ratio: Pd(CN)- $(C_6F_5)(PPh_3)_2$ (0.4120 g, 0.5 mmol) in 20 ml of benzene and $Sn(O_2ClO_2)Bu_3$ (0.5 mmol) in 40 ml of benzene (84% yield).
- (iv) {[Ph₃Sn](μ-CN)[Pd(PPh₃)₂](μ-CN)]_x(ClO₄)_x (V)
 To a solution of Sn(O₂ClO₂)Ph₃ (0.5 mmol) in 25 ml of benzene was added Pd(CN)₂(PPh₃)₂ (0.3415 g, 0.5 mmol) in 40 ml of dichloromethane. The microcrystalline precipitate, which was difficult to filter off, was separated by centrifugation and washed with hexane (70% yield).
- (v) $\{[R_3Sn](\mu\text{-NC})[M(\mu\text{-CN})]\}_x$ (VI, M = Ag, R = Me; VII, M = Ag, R = Ph; VIII, M = Au, R = Me; IX, M = Au, R = Bu; X, M = Au, R = Ph)

 To a solution of $Q[M(CN)_2]$ (M = Ag, $Q = Bu_4N$; M = Au, Q = K) (1 mmol)

To a solution of $Q[M(CN)_2]$ (M = Ag, Q = Bu₄N; M = Au, Q = K) (1 mmol) in 25 ml of acetone was added an equimolecular solution of $Sn(O_2ClO_2)E_3$ in 25 ml of acetone. The immediately precipitated white (or in the case of VIII yellowish) solid was filtered off and repeatedly washed with water/acetone (yields: complex VI, 63%; VII, 76%; VIII, 66%; IX, 64%; X, 81%).

(vi) $\{[R_3Sn]_2[(\mu-NC)_4]\}_x$ (XI, M = Pd, R = Me; XII, M = Pd, R = Bu; XIII, M = Pd, R = Ph; XIV, M = Pt, R = Me; XV, M = Pt, R = Bu; XVI, M = Pt, R = Ph)

To a solution of $K_2[M(CN)_4]$ (1 mmol) in 10 ml of H_2O and 15 ml of acetone was added a solution of $Sn(O_2ClO_2)R_3$ (2 mmol) in acetone (1/2 mole ratio). Stirring was continued for 30—60 min. The white precipitate was filtered off and washed with benzene or acetone and then with hexane (yields: complex XI, 81%; XII, 88%; XIII, 85%; XIV, 91%; XV, 81%; XVI, 93%).

(vii) $\{[R_3Sn]_3[(\mu-NC)_6Fe]\}_x$ (XVII, R = Bu; XVIII, R = Ph)

The reaction was carried out as before by treating 1 mmol of the iron complex with 3 mmol of the organotin perchlorato complex. The resulting compounds were washed with acetone (yields; XVII, 95%; XVIII, 96%).

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