

This is a repository copy of Syntheses, structures, and infrared spectra of the hexa(cyanido) complexes of silicon, germanium, and tin.

White Rose Research Online URL for this paper: https://eprints.whiterose.ac.uk/143581/

Version: Accepted Version

Article:

Smallwood, Z.M., Davis, M.F., Hill, J.G. orcid.org/0000-0002-6457-5837 et al. (2 more authors) (2019) Syntheses, structures, and infrared spectra of the hexa(cyanido) complexes of silicon, germanium, and tin. Inorganic Chemistry, 58 (7). pp. 4583-4591. ISSN 0020-1669

https://doi.org/10.1021/acs.inorgchem.9b00150

This document is the Accepted Manuscript version of a Published Work that appeared in final form in Inorganic Chemistry, copyright © American Chemical Society after peer review and technical editing by the publisher. To access the final edited and published work see https://doi.org/10.1021/acs.inorgchem.9b00150

Reuse

Items deposited in White Rose Research Online are protected by copyright, with all rights reserved unless indicated otherwise. They may be downloaded and/or printed for private study, or other acts as permitted by national copyright laws. The publisher or other rights holders may allow further reproduction and re-use of the full text version. This is indicated by the licence information on the White Rose Research Online record for the item.

Takedown

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.



Syntheses, structures and infrared spectra of the hexa(cyanido) complexes of silicon, germanium and tin

Zoe M. Smallwood, Martin F. Davis[†], J. Grant Hill, Lara J. R. James, Peter Portius*

Department of Chemistry, University of Sheffield, Brook Hill, S3 7HF, U.K.

KEYWORDS. cyanido complexes, DFT, silicon, germanium, tin.

ABSTRACT: The rare octahedral EC₆ coordination skeleton type is unknown for complexes with coordination centers consisting of Group 14 elements. Here, the first examples of such EC₆ species, the hexacoordinate homoleptic cyanido complexes $E(CN)_6^{2^-}$, E = Si, Ge, Sn, have been synthesized from element halides SiCl₄, GeCl₄ and SnF₄ and isolated as salts with PPN counter ions (PPN⁺ = (Ph₃P)₂N⁺) on a scale 0.2-1 g. Characterization by spectroscopic techniques and by structure determination through single crystal crystallographic methods show that these pseudohalogen complexes have effective octahedral symmetry in solution and in the solid state. Infrared spectra obtained in solution reveal that the T_{1u} symmetric IR-active vibrations in all three complexes have unusually small oscillator strengths. The observed reluctance of Si(CN)₆²⁻, Ge(CN)₆²⁻ and Sn(CN)₆²⁻ to form from chloro-precursors was rationalized in terms of Gibbs free energies, which were found by ab initio calculations at the CCSD(T)-F12b/aug-cc-pVTZ(-PP)-F12 level of theory to be small or even positive. The work demonstrates that $E(CN)_6^{2^-}$ complexes of silicon, germanium and tin are in fact stable at r.t. and exist as well-defined units in the presence of non-coordinating counter-ions. The results add to our understanding of the chemistry of pseudohalogens and structure and bonding.

1. INTRODUCTION

Negatively charged cyanido, nitrato, cyanato and azido molecules (Y-) amongst others, possess reactivity similar to that of halides. Birckenbach has rationalized this phenomenon by the concept of the pseudohalogen,^{1, 2} which requires primarily, that the Y⁻ species form Y–Y dimers and Y-X inter-pseudohalogens upon oxidation, have a stable protonated pro-ligand form H-Y, and produce MY_n salts as well as EY_n^{q-} complexes. As the non-existence of hexaazadiene that was proposed to be formed by the oxidation of N_3^- demonstrates,^{3,4} not all pseudohalides fulfil the set of criteria completely.² Pseudohalogen complexes of the type EY_n^{q-} are isolable for most p-block elements in groups 13 to 15: Y = N₃ (B-Tl, Si-Sn, P-Bi), NCS (P, Si-Sn), NCO (Si, Ge, Sn), NO₃ (B, Al, Si-Sn). In group 14, they take the form of EL₄ and EL₆^{2–} (E = Si-Pb, L = N₃, NO₃, N₃, NCO, NCS, NCSe).⁵⁻¹² However, there are only very few homoleptic cyanido complexes. This group of complexes is restricted to coordination centres of the less electronegative elements in groups 13 and 15 as $P(CN)_3$, $Ga(CN)_4^-$, $E(CN)_5^{2-}$ (E = In, Tl, Sb, Bi) and Bi(CN) $_{6}^{3-}$.⁵⁻¹⁶ In group 14, the set of known complexes consist of the E(CN)₄ type (E = Ge, Sn), for which, apart from in situ ¹¹⁹Sn NMR spectral data for the Cl / CN ligand exchange to form $Sn(CN)_6^{2-}$, little analytical data is available.¹⁷⁻²⁰ CN ligands possess a comparably low oxidation potential and this effects, for instance, the low thermal stability of P(CN)₅ which readily eliminates cyanogen at ambient temperature to form $P(CN)_3$.¹³ The related $P(CN)_6^$ complex, on the other hand, is thought to be stabilized by hypercoordination. However, the

synthesis of $P(CN)_6^-$ by means of conventional methods from (fluoro)phosphorus precursors is hampered by an affinity of $P(CN)_5$ toward F^- (590 kJ mol⁻¹) that is vastly greater than that toward CN⁻ (198 kJ mol⁻¹) and has, thus far, not been realised.¹⁴ Intriguingly, tetracyanosilane remains unknown despite numerous investigations into reactions involving silicon tetrahalides and cyanide salts.^{16, 21, 22} $E(CN)_6^{2-}$ structures (E = Si-Pb) are interesting in general as they contain the octahedral EC₆ coordination skeletons which have hitherto, as opposed to the related covalent EC_x networks in silicon carbide containing tetrahedral SiC₄ units,²³ not been reported (see ref. 23 for related coordination geometries). Owing to the strength of the Si-C bonds $[\Delta H_d]/$ $(kJ mol^{-1}) = 318.0$ to 338.9²⁴ and the advantageous match of crystal and coordination geometries, SiC possesses great hardness, and has found many applications due to this property.²³ Recently, germanium and tin complexes of the type $E(Bu)R_2^+$ and $Sn(R-2-py)_6^{2+}$ containing EC₆ coordination skeletons have been prepared with sterically hindered organic and other ligands. These are, however, either not octahedral, or part of larger coordination networks.^{25, 26} Using strategies involving exchange and transfer reactions employed previously for the synthesis of azido complexes,^{6, 10, 11} ionic and covalent cyanides were investigated as reagents for the formation of hexacyanido complexes of the type $E(CN)_6^{2-}$ (E = Si, Ge, Sn), and the results reported here.

2. EXPERIMENTAL SECTION

The moisture sensitivity of starting materials for complexes **1**, **2** and **3** (ECl₄ or Me₃SiCN, see Scheme 1) necessitates the complete exclusion of air, using standard Schlenk-tube and inert gas box techniques. Extended experimental details are contained in the electronic supporting information, alongside all crystallographic information, spectra and other analytical details. The raw-starting material bis(triphenylphosphine)iminium cyanide, (PPN)CN, was prepared from (PPN)Cl and KCN according to Songstad *et al.*²⁷ However, the preparation of chlorine-free 1 relies on the complete absence of chlorine from the PPN starting material which was ensured by the removal of the KCl by-product at an intermediate stage and the application of two batches of KCN in excess each time. Caution! PPN(CN) and the cyanido complexes **1-4** are toxic. The cyanido complexes hydrolyze readily; compounds **4a** and **4b** evolve hydrogen cyanide gas upon exposure to air.

2.1. Synthesis of (PPN)₂Si(CN)₆ (1). In a Schlenk tube, a solution of (PPN)CN (5.012 g, 8.88 mmol, 6.8 equiv.) was prepared in the minimum amount of MeCN. SiCl₄ (0.15 cm³, 1.3 mmol, 1 equiv.) was added and the solution stirred at 60 °C for 1 h, resulting in the formation of a small amount of white precipitate. After removal of the precipitate by filtration, the volume of the colourless filtrate was diminished *in vacuo* until the point of saturation was reached and the solution then cooled to -28 °C for 2 h, forming a white crystalline solid. The solid was collected by filtration, re-dissolved in MeCN and then transferred into a new solution of (PPN)CN (1.000 g, 1.77 mmol, 1.35 equiv.) in the same solvent. The reaction solution was heated as described above which, again, resulted in the formation of a small amount of white solid, attributed to some decomposition of a silicon-containing compound which was removed by filtration. As before, crystallisation was achieved by evaporation of the filtrate *in vacuo* until a saturated solution was obtained at r.t. followed by lowering the temperature to -28 °C. This resulted in a white crop of large, colourless crystals consisting of pure PPN₂Si(CN)₆ (1, 0.696 g, 42%), m.p. 230-235 °C ($T_{dec} = 235$ °C), anal. calcd. for C₇₈H₆₀N₈P₄Si (1261.35 g mol⁻¹): C 74.27, H 4.79, N 8.88 %. Found: C, 74.22; H, 4.58; N, 8.66 %. IR (nujol) \overline{v} / cm⁻¹ = 3470(br), 3056, 2696, 2612, 2592, 2584, 2221(br), 2172, 2163, 2119, 2101, 1985, 1917, 1837, 1687, 1588, 1573, 1481, 1441, 1435, 1325, 1184, 1165, 1161, 1154, 1116, 1073, 1025, 998, 986, 932, 860,

850, 795, 763, 750, 746, 694, 663, 616, 583. ¹³C NMR (CD₃CN) δ / ppm = 128.2 (d, PPN), 130.3 (m, PPN), 133.2 (m, PPN), 134.5 (s, PPN), 139.9 (*C*N). IR (solution, MeCN) $v\overline{\prime}$ cm⁻¹ = 2182, 2175, 2169, 2135. TOF MS ES(–) m/z = 722 ([(PPN)Si(CN)₆]⁻, 100), 158 ([Si(CN)₅]⁻, 47), no signals were observed at 167 ([Si(CN)₅Cl⁻]) and 731 ([(PPN)Si(CN)₅Cl]⁻ or any other Cl-containing species in the m/z range 75 to 900); accurate masses (a.m.u.) found for [Si(CN)₆]⁻, [(PPN)Si(CN)₆]⁻, 157.9921, 722.1806; calcd.: 157.9928, 722.1813. TOF MS ES(+), m/z = 538 ([PPN]⁺, 100).

2.2. Synthesis of (PPN)₂Ge(CN)₆ (2). MeCN (20 ml) was added to (PPN)CN (3.142 g, 5.57 mmol, 6.32 equiv.), resulting in almost complete dissolution of all solid. To this stirred suspension, GeCl₄ (0.10 cm³, 0.88 mmol, 1 equiv.) was added. The suspension was immersed into an oil bath and heated to 65 °C for approximately 41 h; after ~30 mins of heating, only a small amount of solid (<100 mg) remained undissolved. The reaction mixture was allowed to cool to r.t. and settle and then the colorless supernatant solution was filtered off from a white filter residue (112 mg) that was discarded. The volume of the filtrate was the diminished in *vacuo* until the onset of precipitation occurred at which stage all solids were re-dissolved by warming the vessel in hot air. Crystallization was achieved at -25 °C. An initial crop of colorless crystals were obtained by filtration. The volume of the filtrate was further diminished and crystallization induced as described giving a second crop of colorless crystals which were shown to be spectroscopically identical to the first. Both crops were combined to give 2.210 g of mixture of $(PPN)_2Ge(CN)_xCl_y$ (x + y = 6), (PPN)Cl and (PPN)CN. In order to complete the Cl / CN exchange, a mixture of a part of the combined crop (0.459 g, 0.35 mmol if x = 5), and NaCN (0.186 g, 3.80 mmol, 10.86 equiv.), were suspended in MeCN (15 ml). After ~5 mins stirring at r.t., the suspension had changed consistency owing to the dissolution of the germanium complex.

The resultant reaction suspension was immersed in an oil bath set to a temperature of 65 °C. After a total of 196 h heating, the white suspension was filtered and the filter residue discarded. The colorless supernatant solution, containing the hexa(cyanide)complex was concentrated in vacuo until the onset of precipitation, warmed in hot air to re-dissolve all solids, and then placed in a freezer to induce crystallization. This gave colorless crystals which appear white in bulk under a pale yellow solution, which were collected by filtration to give pure $(PPN)_2Ge(CN)_6$ (2, 0.196 g, 82% with respect to GeCl₄) as colorless cuboidal crystals, mp. 188 °C (dec.). Anal. calcd. for C₇₈H₆₀N₈P₄Ge (1305.92 g mol⁻¹) C, 71.74; H, 4.63; N, 8.58; Cl, 0%. Found C, 71.37; H, 4.66; N, 8.63; Cl, <0.3 %. IR (nujol) $\bar{\nu}$ / cm⁻¹ = 3173, 3145, 3079, 3063, 2251(br), 2217(br), 2165, 2159, 1985, 1918, 1906, 1824, 1780, 1684, 1613, 1588, 1575, 1482, 1439, 1436, 1316, 1301, 1284, 1265, 1185, 1162, 1116, 1109, 1073, 1027, 997, 795, 757, 750, 695, 689, 663, 617, 550, 534. ¹H NMR (400 MHz, CD₃CN, r.t.) δ / ppm = 7.45-7.71 (m, PPN). ¹³C NMR (100 MHz, CD₃CN, rt, ppm) δ / ppm = 128.2 (d, PPN), 130.3 (m, PPN), 133.2 (m, PPN), 134.6 (s, PPN), 140.0 (s, CN). IR (solution, MeCN) \overline{v} / cm⁻¹ = 2161, 2135, 2089, 2080(sh), 2051. TOF MS $ES(+) m/z = 538 ([PPN]^+, 100), TOF MS ES(-) m/z = 245(100), 141(81), 123(17), spectra$ dominated by oxo(hydroxo) germanium species; no signals were attributable to $Ge_x(CN)_v^{z-}$.

2.3. Synthesis of Sn(CN)₄(MeCN)₂ (4a). SnF₄ (0.261 g, 1.34 mmol, 1 equiv.) was suspended in MeCN (15 ml). Me₃SiCN (0.67 ml, 5.36 mmol, 4.0 equiv.) was added. After stirring at r.t. for ca. 17 h, a noticeably thicker white suspension had formed. The suspension was filtered, resulting in Sn(CN)₄(MeCN)₂ (0.234 g, 78%) as a white/cream solid filter residue. Anal. calcd for C₈H₆N₆Sn (304.78 g mol⁻¹): C, 31.50; H, 1.98; N, 27.57 %. Found C, 5.46; H, 0; N, 5.41. Note: Sn(CN)₄(MeCN)₂ as well as Sn(CN)₄(py)₂ are extremely air sensitive and produce unreliable

data). IR (nujol) \overline{v} / cm⁻¹ = 3320(br), 2321, 2289, 2241(br), 2182, 1593(br), 1260, 1207, 1169, 844, 804.

2.4. Synthesis of (PPN)₂Sn(CN)₆ (3) from Sn(CN)₄(MeCN)₂ (4a). A Schlenk tube was charged with a mixture of 4a (0.209 g, 0.686 mmol, 1 equiv.) and (PPN)CN (1.262 g, 2.24 mmol, 3.27 equiv.) and then suspended in MeCN (15 ml). The tube was then immersed in an oil bath and the suspension stirred and heated to 60 °C for 21 h. After the reaction mixture had cooled to r.t., a small amount (< 10 mg) of solid material was removed by filtration and discarded. The volume of the pale-yellow filtrate was diminished in vacuo until the onset of precipitation, any solids visible in the solution were re-dissolved by gentle heating and the resultant clear solution cooled to -28 °C. This resulted in an initial crop of yellow block-shaped crystals under a yellow supernatant solution, which was collected by filtration. Further reduction of the volume of the filtrate under vacuum resulted in a second crop of crystals which were spectroscopically identical to the first crop. The combined crops gave analytically pure **3** (0.914 g, 98% with respect to **4a**); mp. 271-275 °C. Anal. calcd. for $C_{78}H_{60}N_8P_4Sn(1351.99 \text{ g mol}^{-1})$ C, 69.30; H, 4.47; N, 8.29; Cl, 0%; found C, 68.74; H, 4.54; N, 8.04; Cl, <0.3%. IR (nujol, cm⁻¹) \overline{v} / cm⁻¹ = 3173, 3146, 3081, 3064, 3039, 3025, 2794, 2787, 2696, 2252, 2218(br), 2157, 2153, 2049(br), 1985, 1915, 1833, 1616, 1588, 1575, 1482, 1439, 1436, 1260, 1252, 1186, 1179, 1114, 1069, 1028, 1020, 998, 941, 929, 924, 864, 797, 755, 695, 691, 664, 616, 550, 532. ¹H NMR (400 MHz, CD₃CN, r.t.) δ / ppm = 7.89-7.22 (m, PPN). ¹³C NMR (100 MHz, CD₃CN, r.t.) δ / ppm = 128.2 (dd, PPN), 130.3 (m, PPN), 133.2 (m, PPN), 134.6(s, PPN), 138.5 (s, CN). IR (solution, MeCN) $\bar{\nu}$ / $cm^{-1} = 2157, 2153(sh), 2135. TOF MS ES(-), MeCN, m/z = 283 ([((NC)_2C_3N_3)_2N_a]^-, 100), 250$ ([Sn(CN)₅]⁻, 33), 198 ([Sn(CN)₃]⁻, 20), 194 ([((NC)₂C₃N₃)Na(MeCN)]⁻, 22), 142

 $([(C_3N_3)Na(MeCN)]^-, 23);$ TOF MS ES(+), MeCN, m/z = 538 ([PPN]⁺, 100); NB no signals were observed at 259 ([Sn(CN)₄Cl⁻]) and 207 ([Sn(CN)₂Cl]⁻) or any other Cl-containing species.

3. RESULTS AND DISCUSSION

3.1. Syntheses. At 60 °C, solutions containing the group 14 chloride and the bis(triphenylphosphine)iminium cyanide (PPN)CN form mixed-ligand complexes of the type $E(CN)_x Cl_y^{2-} (x + y = 6)$ in which the distribution of chlorido(cyanido)complexes depends on the stoichiometric ratio of starting material (Scheme 1).



Scheme 1. Reaction sequences to form $PPN_2Si(CN)_6$ (1), $PPN_2Ge(CN)_6$ (2), $PPN_2Sn(CN)_6$ (3) from element halides *via* $[E(CN)_xCl_y]^{2-}$ intermediates (x + y = 6) and solvate $(Sn(CN)_4(MeCN)_2,$ 4a; $Sn(CN)_4(py)_2$, py = pyridine, 4b) complexes and (PPN)CN, $PPN^+ = (PPh_3)_2N^+$.

When crystallization is induced in the reaction mixture, the close chemical similarity of these $E(CN)_x Cl_y^{2-}$ complexes causes the formation of solid solutions. In the case of silicon, this finding is not only supported by crystallographic data (*vide infra*) but also by microanalytically determined CHN content in these crystals. At CN^- / SiCl₄ starting material ratios of 2 and 10 mol mol⁻¹, the C, H, N contents (% *w/w*) of 69.70, 4.22, 5.43 and 72.98, 4.41, 8.12, respectively,

were found to be consistent to within $\pm 0.5\%$ with the empirical formulae Si(CN)_{2.9}Cl_{3.1}²⁻ and Si(CN)_{5.2}Cl_{0.8}²⁻. This suggests that complete Cl / CN exchange requires molar ratios above 50 mol mol⁻¹, rendering this direct one-step method expensive. Since the $(PPN)_2Si(CN)_xCl_y$ salts are separable from the reaction mixture by crystallization, de-coordinated Cl⁻ can be removed as (PPN)Cl and a supposed reaction equilibrium shifted toward $Si(CN)_6^{2-}$ by exposing $(PPN)_2Si(CN)_xCl_v$ to further amounts of (PPN)CN in a second step under otherwise identical conditions. After this treatment, the Cl content in the newly crystallized material had decreased to below 0.3% which translates to coefficients of x > 5.9 and y < 0.1. GeCl₄ and SnCl₄ also react with (PPN)CN to form chlorido(cyanido) complexes. However, in these cases the preparation of the Cl-free complexes is unfeasible due to an even less favorable position of the equilibrium. Instead, the intermediary $(PPN)_2Ge(CN)_xCl_y$ was converted in the second step to $(PPN)_2Ge(CN)_6$ (2) by applying a large excess of NaCN instead of (PPN)CN and a longer reaction time (196 h). It was found that AgCN is not suitable as a CN-ligand transfer reagent since the crystallization from the solution containing $(PPN)_2Ge(CN)_xCl_v$ and AgCN removes the PPN cation as part of the silver salt (PPN)Ag(CN)₂,²⁸ which was identified by single crystal XRD (see ESI p. 29). This reaction is also likely to occur if AgCN is used in syntheses of the Si and Sn analogs 1 and 3. SnCl₄ showed insufficient reactivity in the presence of either NaCN or (PPN)CN, hence, $Sn(CN)_{6}^{2-}$ cannot be obtained by direct ligand exchange using ionic reagents. It was found that SnF₄ reacts readily with Me₃Si-CN in pyridine or acetonitrile under formation of the adducts $Sn(CN)_4(L)_2$ (L = MeCN, 4a; pyridine, 4b), which react further with (PPN)CN to afford analytically pure (PPN)₂Sn(CN)₆ (3). The degree of progress of the halide / cyanide exchange reactions to all three E(CN)₆²⁻ complexes was established by C, H, N, Cl elemental analyses and electrospray ionization mass spectrometry (see Figs. S23-S25). The mass spectrum of 1 shows

fragments corresponding to $[(PPN)Si(CN)_6]^-$ at m/z = 722 and of $[Si(CN)_6]^-$ at m/z = 158, but none at 731 and 167 that would correspond to a putative $[Si(CN)_5Cl]^-$ species. Similarly, the signals detected of $[Sn(CN)_5]^-$ and $[Sn(CN)_3]^-$ in the spectra of **3** were free of accompanying masses that are likely to occur if Cl were still present. While the solid hexacyanido complexes are moderately air-sensitive, the solutions, especially those of **2** and **3**, hydrolyze readily. Upon exposure to air, the decomposition of the solid white powders **4a** and **4b** also occurs rapidly and is accompanied by color change and the release of HCN. Even in hot polar solvent, the latter compounds are only sparingly soluble thus preventing further structural analysis.

3.2. Crystallographic studies. Colorless (1, 2) or pale yellow (3), cuboidal single crystals suitable for single-crystal X-ray diffraction were obtained from hot, saturated MeCN solutions upon cooling to temperatures below -25 °C. X-ray diffraction studies performed on specimen of each of these types of crystals confirmed that in the three investigated reactions, hexa(cyanido) complexes had formed as part of 2 : 1 salts with the PPN counterions: (PPN)₂E(CN)₆. This result allows for the structural analysis of the E(CN)₆²⁻ complexes of Si (1), Ge (2) and Sn (3) given below (see also ESI p. S9ff). The EC₆ coordination skeletons of the investigated complexes adhere closely to the octahedral symmetry (Figs. 1 to 3) with only small deviations in the interligand angles of no more than $3.80(6)^{\circ}$ and $2.32(6)^{\circ}$ from the ideal geometry of 90° and 180° , respectively, which is ascribed to the differences in the packing imposed by the crystal systems (1, orthorhombic; 2 and 3, monoclinic, Fig. 4). The cyanido ligands adopt the essentially linear *end-on*-C coordination centers of Groups 13 and $15.^{15,29}$ Structure models tested during crystallographic structure refinement that feature isocyanido ligands E(CN)_x(NC)_y²⁻ all were

significantly inferior in predicting the diffraction data, leading to substantially increased disagreement factors (R).



Figure 1. Thermal ellipsoid plots of the Si(CN)₆^{2–} complexes of **1** drawn at the 50% probability level; black, C; blue, N; grey, "i" denotes symmetry-equivalent sites. Bond lengths [Å] and selected angles [°]: Si(1)-C(1) 1.9501(15), Si(1)-C(2) 1.9547(15), Si(1)-C(3) 1.9588(15), C(1)-N(1) 1.152(2), C(2)-N(2) 1.125(2), C(3)-N(3) 1.115(2); C(1)-Si(1)-C(2) 89.55(6), C(1)-Si(1)-C(2) 90.45(6), C(1)-Si(1)-C(3) 88.61(6), C(1)-Si(1)-C(3) 91.39(6), C(2)-Si(1)-C(3) 89.08(6), C(2)-Si(1)-C(3) 90.92(6), Si(1)-C(1)-N(1) 178.01(13), Si(1)-C(2)-N(2) 177.59(14), Si(1)-C(3)-N(3) 175.27(14).



Figure 2. Thermal ellipsoid plots of the $Ge(CN)_6^{2^-}$ complex of 2 drawn at the 50% probability level; black, C; blue, N; grey. Bond lengths [Å] and selected angles [°]: Ge(1)-C(1) 2.0584(15), Ge(1)-C(2) 2.0441(15), Ge(1)-C(3) 2.0491(17), Ge(1)-C(4) 2.0480(15), Ge(1)-C(5) 2.0573(15), Ge(1)-C(6) 2.0407(16), C(1)-N(1) 1.1455(19), C(2)-N(2) 1.145(2), C(3)-N(3) 1.146(2), C(4)-N(4) 1.145(2), C(5)-N(5) 1.146(2), C(6)-N(6) 1.144(2); C(1)-Ge(1)-C(6) 90.99(6), C(2)-Ge(1)-C(5) 179.54(6), C(6)-Ge(1)-C(5) 91.5251(6), C(2)-Ge(1)-C(4) 89.97(6), C(6)-Ge(1)-C(4) 88.52(6), C(5)-Ge(1)-C(4) 89.8685(6), C(2)-Ge(1)-C(3) 91.85(6), C(6)-Ge(1)-C(3) 177.68(6), C(5)-Ge(1)-C(3) 88.58(6), C(4)-Ge(1)-C(3) 93.80(6), C(2)-Ge(1)-C(1) 89.92(6), C(6)-Ge(1)-C(1) 90.99(6), C(5)-Ge(1)-C(1) 90.25(6), C(4)-Ge(1)-C(1) 179.50(5), C(3)-Ge(1)-C(1) 86.70(6), N(1)-C(1)-Ge(1) 176.56(14), N(2)-C(2)-Ge(1) 174.12(14), N(3)-C(3)-Ge(1) 174.90(13), N(4)-C(4)-Ge(1) 174.92(13), N(5)-C(5)-Ge(1) 177.24(13), N(6)-C(6)-Ge(1) 176.14(13).



Figure 3. Thermal ellipsoid plots of the $Sn(CN)_6^{2-}$ complex of 3 drawn at the 50% probability level; black, C; blue, N; grey, "i" denotes symmetry-equivalent sites. Bond lengths [Å] and selected angles: Sn(1)-C(3) 2.203(2), Sn(1)-C(1) 2.225(2), Sn(1)-C(2) 2.229(2), N(1)-C(1) 1.134(3), N(2)-C(2) 1.146(3), N(3)-C(3) 1.159(3); C(3)-Sn(1)-C(1) 91.65(8), C(3)-Sn(1)-C(1) 88.35(8), C(3)-Sn(1)-C(2) 88.43(8), C(3)-Sn(1)-C(2) 88.92(8), C(1)-Sn(1)-C(2) 91.08(8), C(3)-Sn(1)-C(2) 91.58(8), C(1)-Sn(1)-C(2) 88.92(8), N(1)-C(1)-Sn(1) 174.89(19), N(2)-C(2)-Sn(1) 175.73(19), N(3)-C(3)-Sn(1) 179.1(2).

Within each $E(CN)_6^{2-}$ complex, the lengths of C-N and E-C bonds vary only slightly at a level similar to that observed previously in crystals of K₃Fe(CN)₆,²⁷ for instance (see Table 1). This variation is attributable to the crystal lattice-imposed, non-cubic packing around the complex anions resulting in various sets of cation - complex anion contacts (Fig. 4), rather than to electronic structure effects arising from the complexes themselves. It is a well-known fact that

impurities in crystals may lead to deficiencies in crystal structure models (vide supra, discussion on intermediary chlorido(cyanide) species). Chlorine as the most likely candidate for forming solid solutions of the type $(PPN)_2E(CN)_xX_v$ (x + y = 6, X = Cl, and by implication similarly-sized ligands), can be ruled out as sufficiently abundant contaminant. Support for this conclusion is found in the Cl contents of the substances used to grow the crystals for X-ray data collection, which were below the microanalytical detection limit (0.3%). This implies that the x / y ratio, which accounts for any Cl ligands potentially present in the crystals, is close to or above 5.9 / 0.1. Therefore, no more than one additional electron per formula unit is unaccounted for by a structure model that assumes x / y = 6 / 0 (further discussion of residual electron densities relating to Cl can be found in the ESI). The system $(PPN)_2Si(CN)_xCl_y$ (x + y = 6) is particularly prone to the formation of mixed crystals from solutions of incompletely Cl / CN -exchanged complexes, which results in residual electron density at distances from the coordination center typical for Si-Cl bonds in other hexacoordinate Si(IV) complexes (2.25-2.32 Å), but also in a reduction in the unit cell volume in line with typical crystallographic volumes of ligands ($y \sim 2.3$, $V = 6490.5(4) \text{ Å}^3$, y < 0.1, $V = 6662.7(2) \text{ Å}^3$, details see ESI, p. S9).

Table 1. Key structural parameters of the $E(CN)_6^{2-}$ complexes of 1, 2, 3 in comparison with $K_3M(CN)_6$, M = Cr, Fe, derived from X-ray diffraction studies.

Compound	d _{C-N} / Å ^e	d _{E-C} / Å ^e	< E-C-N $\Delta_{max} / ^{\circ}$	< C-E-C Δ _{max} / °	< C-E-C Δ _{max} / °	Ref.
$\mathbf{E} = \mathrm{Si}(1)^{a}$	1.115(2)- 1.152(2)	1.950(2)- 1.959(2)	4.7(1)	1.39(6)	±0	f
$\mathbf{E} = \mathbf{Ge} \left(2 \right)^{b}$	1.144(2)- 1.146(2)	2.041(2)- 2.059(2)	5.9(1)	3.80(6)	2.32(6)	f
$\mathbf{E} = \mathbf{Sn} \left(3 \right)^{b}$	1.134(3)-	2.203(2)-	5.1(1)	1.65(8)	±0	f

	1.159(3)	2.229(2)				
K ₃ Fe(CN) ₆	1.131(22)- 1.167(26)	1.927(14)- 1.971(29)	8.6(1.5)	1.6(7)	±0	С
K ₃ Cr(CN) ₆	1.099(22)- 1.167(17)	2.057(12)- 2.100(10)	2.7(1.0)	1.1(5)	1.0(5)	d

^{*a*} data for 130 K, ^{*b*} data for 100 K, ^{*c*} see refs. 30, 31, ^{*d*} see ref. 32, ^{*e*} minimum-maximum values given, ^{*f*} this work.



Figure 4. Projection of the unit cells of **1** (top) and **3** (bottom) down the crystallographic *a* axes; C, dark grey; N, blue; P, orange; Si, yellow; Sn, olive; H, light grey; ellipsoids set at the 50% level.

A comparison with the formula unit volumes found in the crystal structures of the related PPN₂E(N₃)₆ salts (all *P*-1 symmetry)^{6,10,11} reveals a trend, in which the E(CN)₆²⁻ complexes occupy consistently less space than E(N₃)₆²⁻ complexes: $\Delta V / \text{Å}^3 = -28$ (Si), -66 (Ge), -50 (Sn).

3.3. Spectral Properties. Group theory requires isotopically pure octahedral $E(CN)_6^{q^-}$ complexes to exhibit only a single band in the region of the C=N stretches (v_{CN}) of the IR spectrum due to the t_{1u} symmetric fundamental stretching vibrations. In MeCN solution, the hexacvanido complexes show this band at 2169 cm⁻¹ (1), 2161 cm⁻¹ (2) and 2157 cm⁻¹ (3), respectively (see Fig. 5 bottom and Table 2). Compared to their azido analogues (see refs. given above), the molar extinction coefficients of these bands are very small but still considerably larger than that of the CN⁻ anion itself at 2052 cm⁻¹ in the same medium (see Figs. S1-S13). The shift to higher wavenumbers in comparison to the latter is caused by electron donation from the σ^* orbital of the CN ligand and the ineffective back donation into the π^* acceptor levels due to the lack of *d*-electrons at the coordination center. Compared to the related transition metal complexes Fe(CN)₆⁴⁻, Ti(CN)₆³⁻³³ and Fe(CN)₆³⁻, which display CN stretches at 2098 cm⁻¹, 2071 cm⁻¹ and 2135 cm⁻¹, respectively, those of the title complexes are, intriguingly, all at much higher frequencies and closer to those of the covalently bound tetracoordinate Me₃E-CN cyanides at 2190 cm⁻¹ (E = Si, +21 cm⁻¹), ³⁴ 2181 cm⁻¹ (E = Ge, +20 cm⁻¹), ³⁵ 2175 cm⁻¹ (E = Sn, +18 cm⁻¹).^{36, 37} This finding suggests that the E-C bonds are unusually strong and covalent despite the hypercoordinate character of the $E(CN)_6^{2-}$ complexes. This conclusion is also reflected in the IR spectra of the complexes in the microcrystalline form where, however, additional v_{CN} bands are present (Fig. 5 top). Again, the analytically confirmed purity of each sample (vide supra) rules out any assignment to chlorido(cyanido) species as components of a solid solution. The additional bands are more likely to be caused by the crystal packing-imposed

symmetry at the sites of the E(CN)₆ complexes, which transforms with either the D_{2h} point group (1 and 3) or C_1 (2). Under D_{2h} , three fundamental v_{CN} stretches of E(CN)₆ are IR-active (B_{1u} , B_{2u} and B_{3u}). As these complexes have essentially octahedral geometry, their symmetry is closely related to D_{3d} and two of the three IR-active v_{CN} stretches (A_{2u} and E_u) become degenerate so that only two absorption bands result, which aligns with the observations made in the spectra in the mull of 1 (D_{3h} , 2172 cm⁻¹ intense, 2163 cm⁻¹ medium) and 2 (D_{3h} , 2164 cm⁻¹ medium, 2159 cm⁻¹ intense), whereas the greater distortion of 3 produces three bands (D_{2d} , 2157 cm⁻¹ intense, 2153 cm⁻¹ intense, 2146 cm⁻¹ weak shoulder).



Figure 5. Transmission IR spectra of (PPN)CN (–), PPN₂Si(CN)₆ (1 –), PPN₂Ge(CN)₆ (2 –), and PPN₂Sn(CN)₆ (3 –) between 2185 and 2135 cm⁻¹ (top, mulls) and of MeCN solutions of (PPN)CN (4.0×10^{-2}), 1 (2.0×10^{-2}), 2 (1.0×10^{-2}), and 3 (9.9×10^{-3}) between 2180 and 2145 cm⁻¹ (bottom, concentrations / mol dm⁻³ given in parentheses). The integrated intensities of those bands at 1186 cm⁻¹ and 739-782 cm⁻¹ attributed to the PPN cation were used for normalization.

Table 2. Frequencies of the v_{CN} band absorption maxima observed in the transmission IR spectra of compounds 1-4 and (PPN)CN and chlorido(cyanido) complexes.

Compound	Solid ^{<i>a, b</i>}	Solution <i>a</i> , <i>c</i>	
(PPN)CN	2074, 2058, 2047	е	
(PPN) ₂ Si(CN) ₆ (1)	2172, 2163 $(0.89)^d$	2168.8 $(0.53, 4.74(6))^d$	
$(PPN)_2Ge(CN)_6(2)$	2164, 2159 $(0.70)^d$	2160.8 (0.33, 5.64(6)) ^d	
(PPN) ₂ Ge(CN) _x Cl _y	2155, 2053	-	
(PPN) ₂ Sn(CN) ₆ (3)	2157, 2153, 2146 (1) ^d	2157.3 (1, 5.68(9)) ^{d}	
$(PPN)_2Sn(CN)_xCl_y$	2157, 2153	-	
$Sn(CN)_4(MeCN)_2$ (4a)	2182	f	
Sn(CN) ₄ (py) ₂ (4b)	2165	ſ	
$Pb(CN)_6^{2-}$	-	-	

^{*a*} frequency at band maximum / cm⁻¹, ^{*b*} nujol mulls, ^{*c*} in MeCN, ^{*d*} relative v_{CN} band area, ± 0.02 , and FWHM / cm⁻¹ in parentheses, areas based on compound **3** set to unity; ^{*e*} not observed, ^{*f*} insufficiently soluble.

The frequency of the v_{CN} stretch decreases from 1 to 3 - a trend which matches that found in other pseudohalogen complexes such as $E(N_3)_6^{2-}$ and in the series of Me₃E-CN compounds (E = Si, Ge, Sn, refs. see above). Due to their insolubility in inert solvents suitable for IR spectral investigation in the region of interest, spectra of 4a and 4b solutions were unattainable.

¹³C MAS NMR spectra were recorded of microcrystalline samples for all three hexacyanido complexes. The peaks arising from CN ligands could not be resolved due to overlap with the intense peaks of the PPN cations. ¹³C NMR spectroscopic studies of solutions of compound 1, however, display a well-resolved CN group resonance (at $\delta(^{13}C) = 138.5$ ppm) with a chemical shift in between those of cvanosilanes, e.g. Me₃Si-CN (126.9 ppm),³⁸ and cvanide salts, e.g. (PPN)CN (166.5 ppm in the same solvent). Similar claims can be made for 2 and 3 (Me₃Ge-CN, 127.0 ppm;³⁹ Me₃Sn-CN, ~134 ppm⁴⁰). ¹³C NMR spectra of dmso-*d*₆, solutions of the chargeneutral 4b, show a single resonance attributable to the CN ligands at $\delta = 113.8$ ppm, alongside the three signals of symmetrically coordinated pyridine ligands at 149.6, 136.1 and 123.9 ppm. To corroborate the findings in solution, ¹³C-¹¹⁹Sn cross-polarization experiments were performed at 30 s delay time. However, under these parameters, only signals that belong to the PPN cation were observed, similar to those detected earlier by direct observation of ¹³C. Only minor differences between (PPN)Cl and the complexes are noticeable and signals of CN ligand(s) could not be attributed due to overlap of bands. ¹H T_1 measurements on (PPN)₂Sn(CN)₆ showed long relaxivity times of ~25 s. Therefore, it can be extrapolated that ¹⁵N and ¹³C will require extremely long acquisition times in the order of weeks and these were deemed unfeasible. A ¹⁵N MAS NMR spectrum for compound 3, obtained at a relaxation time of ~ 125 s, displayed only one very weak signal at ~362 ppm, which is assign to the PPN cation.

While all attempts to observe the ²⁹Si and ⁷³Ge resonance of (PPN)₂Si(CN)₆ and (PPN)₂Ge(CN)₆ failed due the lack of sensitivity of the spectrometer probe, the ¹¹⁹Sn MAS NMR spectrum of PPN₂Sn(CN)₆ displays a peak in the range –600 to –1000 ppm, at δ (¹¹⁹Sn) = –871 ppm, which corresponds to the peak at *ca*. –879 ppm of the same compound in MeCN solution (¹*J*(¹¹⁹Sn,¹³C) = 1218 Hz). The variance of δ (¹¹⁹Sn) with that published previously at –916.2 ppm²⁰ is large. The latter is the result of an assignment for *in-situ*-generated $Sn(CN)_6^{2-}$ in CH₂Cl₂ solution; the authors state that $Sn(CN)_6^{2-}$ could not be isolated. Given the independent, multiply verified identity and purity of the substance obtained as compound **3**, and the similarity between MAS and solution ¹¹⁹Sn NMR spectra of this substance, we suggest that the previously observed chemical shift belongs to a Sn-containing species other than $Sn(CN)_6^{2-}$ (see also Figs. S14-22).

3.4. Theoretical Studies. In order to rationalize the experimental findings, gas phase and solution energies of CN^- , Cl^- , $E(CN)_4$, $E(CN)^{5-}$, $E(CN)_6^{2-}$ and $E(CN)_5Cl^{2-}$ were calculated using density functional theory and coupled cluster methods (see ESI for full computational details). These energies are related to each other according to the reactions analogous to those evaluated by Haiges *et al.*¹⁵ (*i*)-(*iii*) shown below. In view of earlier reports on trimethylsilylcyanide, which exists as an equilibrium mixture with the N-bonded silylisocyanide through a rapid CN group exchange reaction,⁴⁶ the relative stability of the linkage isomers $E(CN)_5(NC)^{2-}$ was assessed relative to the homoleptic cyanido complexes, $E(CN)_6^{2-}$ (E = Si-Sn) (*iv*).

$$E(CN)_4 + CN^- \to E(CN)_5^- \qquad (i)$$

$$E(CN)_{5^{-}} + CN^{-} \rightarrow E(CN)_{6}^{2^{-}}$$
(*ii*)

$$E(CN)_5Cl^{2-} + CN^- \rightarrow E(CN)_6^{2-} + Cl^- \qquad (iii)$$

$$E(CN)_6^{2-} \rightarrow E(CN)_5(NC)^{2-} \qquad (iv)$$

Geometry optimizations and harmonic frequency calculations were carried out at the PBE0 level of theory⁴⁴ using the aug-cc-pV(T+d)Z basis set for C, N, Si and Cl,⁴² where the "+d" denotes additional tight *d* functions for second row atoms, and the aug-cc-pVTZ-PP basis for the heavier atoms.⁴³ The latter is matched to small-core relativistic pseudopotentials replacing 10, 28 and 60

electrons for Ge, Sn and Pb, respectively. The complexes $E(CN)_4$, $E(CN)_5^-$, $E(CN)_5Cl^{2-}$ and $E(CN)_6^{2-}$, E = Si-Pb, were optimized and were found to have T_d , D_{3h} , C_{4v} and O_h symmetry, respectively. Frequency calculations of the minimum geometries obtained after optimization resulted in no imaginary frequencies (see ESI p. S68). Bond lengths and angles derived from the calculated geometries for the $E(CN)_6^{2-}$ complexes **1-3** and those observed crystallographically (Table 3) are within, or close to, the experimental 1σ uncertainty. Therefore, it is suggested that the PBE0/aug-cc-pV(T+d)Z(-PP) model is sufficiently reliable for predicting geometries and spectral properties, and can be used in conjunction with coupled cluster methods for energetics.

Table 3. Observed (obs) and calculated (calc) bond lengths (d / Å), and frequencies (v / cm^{-1}) and intensities of the CN stretching vibrations of the $E(CN)_6^{2-}$ (a) and $E(CN)_5(NC)^{2-}$ (b) complexes.

		E-L _{obs} ^a	E-L _{calc}	C-N _{obs} ^a	C-N _{calc}	\mathcal{V} CN(obs) b	VCN(calc) ^c
E = Si	(a)	1.956(10)	1.961	1.144(16)	1.156	2168.8 (0.53)	2281.8(0.08)
	(b)	-	1.957^{d}	-	1.156 ^{<i>d</i>}	-	2284.4(0.34) ^g
			1.863 ^e		1.166 ^e		2202.6(207) ^{<i>h</i>}
E = Ge	(a)	2.052(7)	2.059	1.152(2)	1.156	2160.8 (0.33)	2277.2(0.02)
	(b)	-	2.048 ^d	-	1.156 ^{<i>d</i>}	-	2281.6(0.00) ^g
			1.999 <i>°</i>		1.166 ^e		2194.0(195) ^{<i>h</i>}
E = Sn	(a)	2.222(14)	2.238	1.156(14)	1.157	2157.3 (1)	2276.0(1.57)
	(b)	-	2.227 ^d	-	1.156 ^{<i>d</i>}	-	2279.5(1.11) ^g
			2.168 ^e		1.167 ^e		2191.5(193) ^{<i>h</i>}
E = Pb	(a)	-	2.329	-	1.157	-	2265.3(2.43)

^{*a*} Thermal motion in the crystal may produce an apparent shrinkage of molecular dimensions and is considered by calculating instantaneous bond lengths D_i , which are estimated by $D_i \approx D_0 + |U_A - U_B| / D_0$ using the distances D_0 and thermal displacement parameters U_A , U_B for atoms A, B of the refined crystallographic structure solution; E-C_{obs} and C-N_{obs} are listed as unweighted mean x_u of D_i with either standard deviation $\sigma = [\Sigma(D_i - x_u)^2/(n-1)]^{0.5}$ or distribution of means $\sigma_m = \sigma_i / n^{0.5}$ - whichever is larger - in parentheses; n = 3, Si, Sn; n = 6, Ge; ^{*b*} in MeCN solutions, relative band areas in parentheses; ^{*c*} oscillator strength (in km mol⁻¹) given in parentheses; ^{*d*} cyanido ligands, ^{*e*} isocyanido ligand; ^{*g*} *E*-symmetric C-N stretches; ^{*h*} isocyanido C-N stretch. Remarkably, the C=N bond lengths are influenced only slightly by a change of the coordination center, which indicates that the electronic situation of the C=N bond is only marginally dependent on the central atom in group 14. The calculated vibrational frequencies and absorption cross sections validate the IR spectroscopic findings reported above suggesting that the transition dipole moments⁴⁴ of the IR-active T_{1u} -symmetric CN stretches have a minimum for Ge(CN)₆²⁻ and a maximum for Pb(CN)₆²⁻. The calculated frequencies for the isocyanido CN stretches in the hypothetical complexes of the type E(CN)₅(NC)²⁻ (Si-Sn) are predicted to be about 80 cm⁻¹ below those observed, and their absorption coefficients to be three to four orders of magnitude more intense than those of the cyanido ligands. In the recorded spectra, this region is free of such bands and therefore we suggest that if isocyanido complexes are present in solution, then at a very low concentration.

Table 4 displays the net electronic energy changes for the reactions (*i*), (*ii*) and (*iii*), as well as the correction terms for changes in the Gibbs free energies caused by thermal and solvent effects. As can be expected, the Lewis acid-base reactions of the attachment of a cyanido anion to the tetracoordinate tetracyano complex are highly exothermic, whereas the attachment of another cyanido ligand to form the hexacoordinate complexes is endothermic. While both types of reactions are endergonic at 298 K, solvent effects cause a large reduction in the Gibbs free energies so that, according to the model used, reactions (*i*) and (*ii*) are spontaneous at r.t. for all evaluated coordination centers.

Е	Reaction	$E_{\text{CCSD(T)}}^{a}$	$\Delta G_{\text{therm}}{}^{b}$	$\Delta G_{ m solv}{}^c$	$\Delta G_{ m sol}{}^e$
Si	<i>(i)</i>	-295.4	+48.4	+147.6	-99.3
	(<i>ii</i>)	+26.5	+57.9	-176.9	-92.4
	(iii)	-50.8	+18.2	+20.6	-12.0
	<i>(iv)</i>	-	-	-	+4.4
Ge	<i>(i)</i>	-253.3	+45.5	+163.7	-44.1
	(<i>ii</i>)	+60.2	+54.7	-159.9	-44.9
	(iii)	-39.5	+17.6	+19.1	-2.8
	<i>(iv)</i>	-	-	-	+12.5
Sn	<i>(i)</i>	-302.5	+42.8	+185.4	-74.2
	(ii)	+20.6	+54.8	-133.0	-57.6
	(iii)	-27.2	+17.0	+18.1	+7.9
	(iv)	-	-	-	+10.2
Pb	<i>(i)</i>	-271.4	+40.3	+198.6	-32.5
	(ii)	+48.9	+52.1	-114.1	-13.1
	(iii)	-19.1	+16.5	+15.3	+12.7

Table 4. Reaction energies in units of kJ mol⁻¹ at 298 K for the formation of the E(CN)₆^{2–} complexes.

^{*a*} Gas phase data at the CCSD(T)-F12b/aug-cc-pVTZ(-PP)-F12 level of theory using PBE0optimised geometries; ^{*b*} thermodynamic Gibbs free energy (298 K) at the PBE0 level; ^{*c*} Gibbs free energy of solvation in acetonitrile at the PBE0 level using the SMD approach; ^{*e*} solution phase reaction energies.

Since the synthesis of complexes 1-3 was investigated with ECl₄ as starting material (*vide supra*), it is the $E(CN)_5Cl^{2-}$ complexes that are the ultimate intermediates to the hexacyano

complexes in the by Cl / CN exchange reactions. The solvent effect-corrected Gibbs free energies for 298 K, the temperature at which **1-3** were isolated from the reaction solutions, clearly corroborate the experimentally observed trend in the reduction of the driving force for this reaction. Based on free Gibbs energies, ΔG_{sol} , for solvated complexes, concentration ratios of E(CN)₆^{2–} / E(CN)₅Cl^{2–} at 298 K can be calculated as 1.3×10^2 , 3.1, 4.1×10^{-2} , 6.0×10^{-3} , respectively, which show that within this model, reaction (*iii*) to form Sn(CN)₆^{2–} from SnCl₄ with soluble ionic cyanides, cannot be driven to completion. The ΔG_{sol} energies furthermore indicate a clear preference for homoleptic hexacyanido complexes E(CN)₆^{2–} over their E(CN)₅(NC)^{2–} linkage isomers.

Conclusion

Hexa(cyanido) complexes of the type $E(CN)_6^{2^-}$, E = Si, Ge, Sn, have been synthesized, isolated and fully characterized as salt-like compounds with PPN counterions. The PPN₂E(CN)₆ compounds are moderately air sensitive and dissolve readily in polar aprotic solvents without noticeable dissociation or linkage isomerization of the $E(CN)_6^{2^-}$ complex anions. The synthetic methodologies applied to these hypercoordinate complexes are partially based on those used previously for the N₃, NCS and NCO analogues.^{6,8,45} However, more forcing conditions are required for the final reaction step $E(CN)_5Cl^{2^-} + CN^- \rightarrow E(CN)_6^{2^-} + Cl^-$, the reasons for which were identified with the help of systematic DFT and coupled cluster calculations. According to these calculations it is primarily the reduction in the net-loss of electronic energy in the reaction involving tin and, presumably, the similarity in Sn-CN and Sn-Cl bond strengths which cause the supposed reaction equilibrium to lie on the side of the starting material. Crystallographic studies have demonstrated that the CN ligands prefer the κ (C)-cyanido form in an only marginally distorted octahedral ligand sphere which demonstrates the viability of homoleptic octahedral complexes with the unusual EC_6 coordination skeletons in group 14.

ASSOCIATED CONTENT

Supporting Information.

The supporting information is available free of charge on the ACS Publications website at DOI: And contains details of the crystallographic structure determination for compounds **1**, **2**, **3** and ((PPN)Ag(CN)₂, FTIR, NMR and mass spectra, Cartesian coordinates of the computational models and further preparative and experimental details. The crystallographic information files for compounds **1** (CCDC 1820362), **2** (CCDC 1820360), **3** (CCDC 1820145), (PPN)Ag(CN)₂ (CCDC 1856451) are available free of charge from the Cambridge Crystallographic Data Centre (CCDC).

AUTHOR INFORMATION

Corresponding Author

*E-mail: p.portius@shefield. ac.uk

ORCID

Zoe Smallwood: 0000-0001-8962-6285

Martin Davis:

J. Grant Hill: 0000-0002-6457-5837

Lara James:

Peter Portius: 0000-0001-8133-8860

Notes

The authors declare no competing financial interests.

Present Address

† Martin F. Davis, Nanoco Technologies Ltd., Manchester M13 9NT, UK.

ACKNOWLEDGEMENT

This work was supported by the University of Sheffield; in particular, the authors thank the university for the award of a graduate teaching assistantship to ZS. The authors thank Prof. Wolfgang A. Herrmann (TU München) for his suggestions, Dr. Sandra van Meurs for recording NMR spectra and Harry Adams and Craig Robertson for their technical assistance in obtaining crystallographic data.

REFERENCES

(1) Birkenbach, L.; Kellermann, K. Über Pseudohalogene (I). Ber. Dtsch. Chem. Ges., 1925, 58B, 786-794.

(2) Golub, A. M.; Köhler, H.; Skopenko, V. V. "Chemistry of Pseudohalides", **1986**, Elsevier, New York.

(3) Greschner, M. J.; Zhang, M.; Majumdar, A.; Liu, H.; Peng, F.; Tse, J. S.; Yao, Y. A New Allotrope of Nitrogen as High-Energy Density Material. *J. Phys. Chem. A*, **2016**, *120*, 2920-2925.

(4) Najafpour, J.; Foroutan-Nejad, C.; Shafiee, G. H.; Peykani, M. K. How does electron delocalization affect the electronic energy? A survey of neutral poly-nitrogen clusters. *Comput. Theor. Chem.*, **2011**, *974*, 86-91.

(5) Wiberg, E.; Michaud, H. Silicon tetraazide, Si(N₃)₄. Z. Naturforsch., B: Chem. Sci., 1954, 9, 500.

(6) Filippou, A. C.; Portius, P.; Schnakenburg, G. The Hexaazidosilicate(IV) Ion: Synthesis, Properties and Molecular Structure. *J. Am. Chem. Soc.*, **2002**, *124*, 12396-12397.

(7) Seiler, O.; Burschka, C.; Goetz, K.; Kaupp, M.; Metz, S.; Tacke, R. The new λ⁶-Si-silicate dianion [Si(NCO)₆]²⁻: synthesis and structural characterization of [K(18-crown-6)]₂[Si(NCO)₆]. *Z. Anorg. Allg. Chem.*, **2007**, *633*, 2667-2670.

(8) Heininger, W.; Stucka, R.; Nagorsen, G. A new silicate anion: [Si(NCS)₆]²⁻. Z. *Naturforsch., B: Chem. Sci.*, **1986**, *41*, 702-707.

(9) Heininger, W.; Polborn, K.; Nagorsen, G. A further silicate anion: $[Si(NCSe)_6]^{2-}$. Z. Naturforsch., B: Chem. Sci., **1988**, 43, 857-861.

(10) Filippou, A. C.; Portius, P.; Neumann, D. U.; Wehrstedt, K.-D. The Hexaazidogermanate(IV) Ion: Syntheses, Structures, and Reactions. *Angew. Chem. Int. Ed.*, 2000, *39*, 4333-4336.

(11) Campbell, R.; Davis, M. F.; Fazakerley, M.; Portius, P. Taming Tin(IV) Polyazides. *Chem. Eur. J.*, **2015**, *51*, 18690-18698. (12) Portius, P.; Peerless, B.; Davis, M.; Campbell, R. Homoleptic Poly(nitrato) Complexes of Group 14 Stable at Ambient Conditions. *Inorg. Chem.*, **2016**, *55*, 8976-8984.

(13) Gall, H.; Schüppen, J. Über die Valenz-Grenze bei Phosphorcyaniden und Phosphorrhodaniden. *Chem. Ber.*, **1930**, *32*, 482-487.

(14) Bläsing, K.; Ellinger, S.; Harloff, J.; Schulz, A.; Sievert, K.; Täschler, C.; Villinger, A.;
Zur, C. Lewis Acid Catalyzed Synthesis of Cyanidophosphates. *Chem. Eur. J.*, 2016, 22, 4175-4188.

(15) Haiges, R.; Deokar, P.; Vasiliu, M.; Stein, T. H.; Dixon, D. A.; Christe, K. O. Preparation and Characterization of Group 13 Cyanides. *Chem. Eur. J.*, **2017**, *23*, 1-14.

(16) Arlt, S.; Harloff, J.; Schulz, A.; Stoffers, A.; Villinger, A. Cyanido Antimonate(III) and Bismuthate(III) Anions. *Inorg. Chem.*, **2016**, *55*, 12321–12328..

(17) Menzer, W. Zur Kenntnis des Germaniumtetracyanids. Angew. Chem., 1958, 70, 656.

(18) Jung, S.; Renz, F.; Klein, M.; Menzel, M.; Boča, R.; Stößer, R. Molecular switching in iron complexes bridged via tin-cyanides observed by Mössbauer and ESR spectroscopy. *J. Phys. Conf. Ser.*, **2010**, vol. *217*, p. 12027.

(19) Bither, T. A.; Knoth, W. H.; R. V. Lindsey, J.; Sharkey, W. H. Trialkyl- and Triaryl(iso)cyanosilanes. *J. Am. Chem. Soc.*, **1958**, *80*, 4151-4153.

(20) Dillon, K. B.; Marshall, A. Tin-119 Nuclear Magnetic Resonance Studies of some
Pseudohalogenoderivatives of [SnX₆]²⁻ (X = CI or Br). *J. Chem. Soc. Dalton Trans.*, **1987**, 315-317.

(21) Emerson, K. University of Minnesota, 1960.

(22) Howdeshell, J. Ph. D. Thesis. University of Wyoming, 1971.

(23) Fan, J.; Chu, P. K.-H. Silicon Carbide Nanostructures. Engineering Materials and Processes, Springer International Publishing, Switzerland, 2014, Chapter 2, Fabrication, Structure, Properties.

(24) Steele, W. C.; Nichols, L. D.; Stone, F. G. A. The Determination of Silicon-Carbon and Silicon-Hydrogen Bond Dissociation Energies by Electron Impact. *J. Am. Chem. Soc.*, **1962**, *84*, 4441-4445.

(25) Schrader, I.; Zeckert, K.; Zahn, S. Dilithium Hexaorganostannate(IV) Compounds. *Angew. Chem. Int. Ed.*, **2014**, *53*, 13698-13700.

(26) Yang, Y.; Panisch, R.; Bolte, M.; Müller, T. Bis-alkene Complexes of Stannylium and Germylium Ions. *Organometallics*, **2008**, *27*, 4847-4853.

(27) Martinsen, A.; Songstad, J. Preparation and properties of some bis(triphenylphosphine)iminium salts, [(Ph₃P)₂N]X. *Acta Chem. Scand. A*, **1977**, *31*, 645-650.

(28) Jaafar, M.; Liu, X.; Dielmann, F.; Hahn, F. E.; Al-Farhan, K.; Alsalme, A.; Reedijk, J. Synthesis, structure and spectroscopic properties of bis(triphenylphosphane)iminium(chlorido) (cyanido) argentates(I). *Inorg. Chim. Acta*, **2016**, *443*, 45-50.

(29) Deokar, P.; Leitz, D.; Stein, T. H.; Vasiliu, M.; Dixon, D. A.; Christe, K. O.; Haiges, R. Preparation and Characterization of Antimony and Arsenic Tricyanide and Their 2,2'-Bipyridine Adducts. *Chem. Eur. J.*, **2016**, *22*, 13251-13257.

(30) Vannerberg, N.-G. OD [order-disorder] structures of potassium hexacyanoferrate(III) and potassium hexacyanocobaltate(III). *Acta Chem. Scand.*, **1972**, *26*, 2863-2876.

(31) Shimanouchi, T.; Nakagawa, I. Infrared spectroscopic study on the co-ordination bond-II:I Infrared spectra of octahedral metal cyanide complexes. *Spectrochim. Acta*, **1962**, *18*, 101-103.

(32) Jagner, S.; Ljungstrom, E.; Vannerberg, N. G. Crystal structure of potassium hexacyanochromate(III). *Acta Chem. Scand, A*, **1974**, *28*, 623-630.

(33) Entley, W. R.; Treadway, C. R.; Wilson, S. R.; Girolami, G. S. The Hexacyanotitanate Ion: Synthesis and Crystal Structure of [NEt₄]₃[Ti^{III}(CN)₆].4MeCN. *J. Am. Chem. Soc.*, **1997**, *119*, 6251-6258.

(34) Booth, M. R.; Frankiss, S. G. The constitution, vibrational spectra and proton resonance of trimethylsilyl cyanide and isocyanide. *Spectrochim. Acta A*, **1970**, *26*, 859-869.

(35) Watari, F. Vibrational Spectra and normal coordinate calculations for (CH₃)₃GeCN and (CD₃)₃GeCN. *J. Mol. Struct.*, **1976**, *32*, 285-295.

(36) Seyferth, D.; Kahlen, N. Trimethyl(iso)cyanogermane and trimethyltin (iso)cyanide. *J. Org. Chem.*, **1960**, *25*, 809-812.

(37) Lorberth, J. Preparation of organotin cyanides of the types R₃SnCN and R₂Sn(CN)₂. *Chem. Ber.*, **1965**, *98*, 1201-1204.

(38) Arnold, D. E. J.; Cradock, S.; Ebsworth, E. a. V.; Murdoch, J. D.; Rankin, D. W. H.; Skea,D. C. J.; Harris, R. K.; Kimber, B. J. Nuclear Magnetic Resonance Spectra of Some Silyl andTrimethylsilyl Pseudo Halides. *Dalton Trans.*, **1981**, 1349-1351.

(39) Drake, J. E.; Glavincevski, B. M.; Humphries, R. E.; Majid, A. The carbon-13 chemical shifts of various methylgermanium derivatives. *Can. J. Chem.*, **1979**, *57*, 1426-1430.

(40) Brimah, A. K.; Siebel, E.; Fischer, R. D.; Davies, N. A.; Apperley, D. C.; Harris, R. K.
Towards organometallic zeolites: Spontaneous self-assembly of Et₃SnCN, CuCN and
("Bu₄N)CN to supramolecular [("Bu₄N)(Et₃Sn)₂Cu(CN)₄]. *J. Organomet. Chem.*, **1994**, *475* 85-94.

(41) Adamo, C.; Barone, V. Toward reliable density functional methods without adjustable parameters: The PBE0 model. *J. Chem. Phys.*, **1999**, *110*, 6158-6170.

(42) Dunning Jr, T. H.; Peterson, K.; Wilson, A. Gaussian basis sets for use in correlated molecular calculations. X. The atoms aluminum through argon revisited. *J. Chem. Phys.*, **2001**, *114*, 9244.

(43) Peterson, K. A. Systematically convergent basis sets with relativistic pseudopotentials. I. Correlation consistent basis sets for the post-d group 13-15 elements. *J. Chem. Phys.*, **2003**, *119*, 11099-11112.

(44) Gribov, L. A. The theory of intensities in the infrared spectra of polyatomic molecules. *Pure Appl. Chem.*, **1969**, *18*, 339-351.

(45) Portius, P.; Davis, M. A new hexakis(isocyanato)silicate(IV) and the first neutral Lewisbase adducts of silicon tetraisocyanate. *Dalton Trans.*, **2010**, *39*, 527-532.

(46) Booth, M. R.; Frankiss, S. G. Spectrochim. Acta, Part A, 1970, 26, 859-869.



The previously unknown hexacoordinate homoleptic cyanido species $Si(CN)_6^{2^-}$, $Ge(CN)_6^{2^-}$ and $Sn(CN)_6^{2^-}$ were synthesized and found to be stable at r.t. as well-defined octahedral complexes in solution and in solid state in the presence of non-coordinating PPN counter-ions. The Cl/CN ligand exchange required to access $E(CN)_6^{2^-}$ from Cl-containing precursors and ionic CN-transfer reagents is thermodynamically hindered. The complexes were characterized by spectroscopic, crystallographic and computational methods.