The Phenyl Group as a Ligand in Divalent Tellurium Complexes

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Complexes of benzenetellurenyl chloride and bromide with one molecule of thiourea, $C_6H_5Te(tu)Cl$ (I) and $C_5H_5Te(tu)Br$ (II), and of benzenetellurenyl chloride with two molecules of thiourea, $C_6H_5Te(tu)_2Cl$ (III), were reported some time ago ¹. The mono-thiourea compounds form isomorphous crystals, space group $C_{2b}^5-P2_1/n$ with Z=4, and $^1a=6.32$ Å, b=10.62 Å, c=15.16 Å, $\beta=90\frac{1}{2}^\circ$ for I and a=6.46 Å, b=10.73 Å, c=15.36 Å, $\beta=91\frac{1}{2}^\circ$ for II. The di-thiourea chloride crystallizes in space group $D_2^4-P2_12_12_1$ with Z=4, and a=11.98 Å, b=20.80 Å, c=5.79 Å. The crystal and molecular structures of the three compounds have been determined by X-ray methods, by use of CuKa radi-

ation, through projections along the shortest and next shortest crystal axes. An account of the main molecular features is given here.

In all three compounds the tellurium atom is primarily three-coordinated, being bonded to one phenyl group, one thiourea sulphur atom and a halogen atom in I and II and to a phenyl group and two thiourea sulphur atoms in III. In view of the tendency of divalent tellurium to form four-coordinated, square-planar complexes ², we regard the phenyl compounds as based on square-planar coordination with one position, opposite to the phenyl group, vacant.

The arrangement around tellurium, as seen along the b crystal axis of I and II and the c axis of III, is pictured in Fig. 1. In III the Te-S bond lengths are 2.63 and 2.70 Å \pm 0.02 Å (uncertainties quoted are estimated standard deviations). The S-Te-S angle is 173° and the S-Te-C angles 87° and 86°, all \pm 1°. In I, Te-S = 2.50 Å and Te-Cl = 3.00 Å, both \pm 0.015 Å; \angle S-Te-Cl = 172°, \angle S-Te-C = 90°, \angle Cl-Te-C = 84°, all \pm 0.5°. In II, Te-S = 2.50 \pm 0.015 Å, Te-Br = 3.11 \pm 0.01 Å, \angle S-Te-Br = 174°, \angle S-Te-C = 90°, \angle Br-Te-C = 86°,

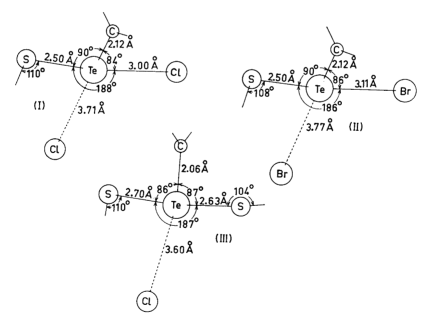


Fig. 1. The bonding to tellurium in (I) $C_6H_5Te(tu)Cl$, (II) $C_6H_5Te(tu)Br$ and (III) $C_6H_5Te(tu)_2Cl$.

all \pm 0.5°. The phenyl groups have been assumed regular hexagons with Te-C₁···C₄ linear. The Te-C₁ bonds are, within the accuracy, of normal length, about 2.10 Å.

accuracy, of normal length, about 2.10 Å. The mean length, 2.67 Å, of the two, approximately equal Te-S bonds in III is within the accuracy the same, 2.68 Å, as found for Te-S in centrosymmetric square-planar complexes 2. In I and II the Te-S bonds are shorter, 2.50 Å, and the Te-Cl and Te-Br bonds, in directions approximately linear to the Te-S bonds, are correspondingly longer. This is the same bond lengthening effect on Te-halogen bonds, of thiourea ligands in transpositions to halogen, as observed 2 in cis-Te(tu)₂Cl₂ and cis-Te(tu)₂Br₂.

The phenyl group has, however, a far more pronounced bond lengthening effect in a direction trans to itself than has thiourea. This effect of the phenyl group (strictly: the phenate ion when regarded as a coordinating ligand) is so large so as to virtually expel the ligand trans to it, and to make the complexes three-coordinated. In the crystals of III, there is a chloride ion in a direction 165° to the C-Te bond at a distance of 3.60 \pm 0.02 Å from tellurium; correspondingly, in I and II there are chlorine and bromine atoms, respectively, in directions of 164° at distances Te···Cl = 3.71 ± 0.015 Å and $Te \cdot \cdot \cdot Br = 3.77 \pm 0.01$ Å. These may be regarded as the fourth, missing ligands of square-planar arrangements, or at any rate, as very loosely bound ligands. In terms of a bonding scheme for divalent tellurium complexes based on p-orbitals 2, the highly nucleophilic phenate engages a tellurium 5p-orbital in bonding so effectively, that little or no bonding power of this p-orbital is left for bonding in a direction trans to the C-Te bond.

The work has been aided by a grant from Norges Almenvitenskapelige Forskningsråd.

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Received July 18, 1963.

Structure of the Triselenocyanate Ion in the Potassium Salt

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The pseudohalogens, thiocyanogen 1,2 and selenocyanogen 3,4 add the respective pseudohalide ions to give trithiocyanate 5,6 and triselenocyanate 4,7 ions, like the halogens. Salts of the former are very unstable. In rounding off our work 8,9 on divalent tellurium complexes, we sought for a divalent selenium complex suitable for crystal structure analysis, and have succeeded in determining the structure of potassium triselenocyanate hemihydrate, K(SeCN)₃·½H₂O, first prepared by Verneuil 7 although by him described without crystal water.

We prepared the salt from concentrated aqueous potassium selenocyanate by oxidation at room temperature with bromine dissolved in benzene. The reddish brown crystals which separated out were recrystallized from the aqueous mother liquor, by dissolving them through gentle heating and allowing the solution to cool slowly. (Found: Se 65.07; 65.37. Calc. for $K(SeCN)_3 \cdot \frac{1}{2}H_2O$: Se 65.25).

The salt forms long monoclinic prisms elongated along the b axis, with a = 17.00 $A, b = 4.44 A, c = 13.86 A, \beta = 122^{\circ}.$ The space group is C_2^3-C2 and there are four formula units per unit cell; density, calc. 2.72, found 2.74 g/cm³. The intensities of 206 h0l and 43 hk0 reflections were read from zero-level Weissenberg photographs taken with CuKa radiation. The approximate positions of the selenium atoms in the b- and c-axis projections were found from the Patterson maps, and the projections were refined through Fourier and difference syntheses. The reliability index R is 0.087 for the h0l and 0.109 for the hk0 reflections. The selenium coordinates, with origin on twofold axis at $y(\text{Se}_1)$ are, Se_1 : $x=0.1553,\ y=0,\ z=0.1351;\ \text{Se}_2$: $x=0.3035,\ y=0.0019,$