

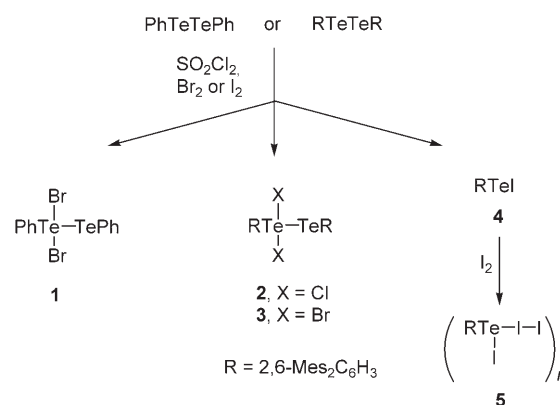
Formation of Mixed-Valent Aryltellurenyl Halides RX_2TeTeR^{**}

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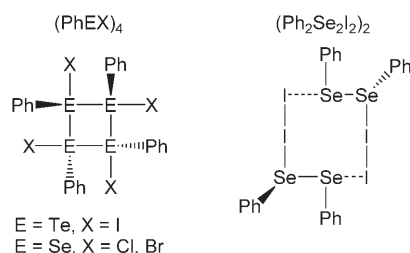
Extremely unstable sulfur difluoride, SF_2 , dimerizes to give the slightly more stable mixed-valent dinuclear F_3SSF .^[1] To date the only reported organic derivative of F_3SSF is $F_3CF_2SSCF_3$, which was identified during the disproportionation reaction of F_3CSF , affording F_3CSSCF_3 and F_3CSF_3 .^[2] While organoselenenyl(II) halides, such as $PhSeX$ ($X = Cl, Br$), are more stable and often even commercially available, aryltellurenyl(II) halides, $RTeX$, are also intrinsically unstable and usually observed only as short-lived intermediates in halogenation reactions of diorganoditellurides, $RTeTeR$, or in synproportionation reactions of $RTeTeR$ with $RTeX_3$.^[3] A common decomposition pathway of aryltellurenyl(II) halides, $RTeX$, proceeds by migration of the organic substituents and produces diaryltellurium(IV) dihalides, R_2TeX_2 , and elemental tellurium. From the decomposition of the sterically more encumbered $MesTeI$ ($Mes = 2,4,6-Me_3C_6H_2$), the dinuclear complex $Mes_2TeTeMesI$, $MesTeI_3$, and elemental tellurium were isolated.^[4] Very few kinetically stabilized aryltellurenyl(II) halides, such as $2,4,6-tBu_3C_6H_2TeX$ ($X = Cl, Br, I$), are known,^[5] and they are monomers in the solid state. In 1974, Schulz and Klar described a number of metastable organotellurenyl halides, $RTeX$ (e.g. $R = Ph, 4-MeOC_6H_4, 4-PhC_6H_4$; $X = Br, I$) containing smaller organic substituents, which on the basis of their rather poor solubility in most organic solvents and their rather dark color were presumed to have oligomeric and/or polymeric structures.^[6] Recently, one of these compounds has been identified as the cyclic tetramer $(PhTeI)_4$.^[7] The halogenation of $PhSeSePh$ with Cl_2 and Br_2

produced analogous selenium tetramers $(PhSeX)_4$ ($X = Cl, Br$),^[8] whereas the reaction with I_2 afforded a related charge transfer complex $Ph_2Se_2I_2$ having a dinuclear structure with secondary $Se \cdots I$ contacts.^[9]

The prospect of finding novel oligomeric organotellurenyl(II) halides and related charge-transfer complexes prompted us to study the halogenation of two diarylditellurides in greater detail. The reaction of $PhTeTePh$ with one equivalent of bromine provided the mixed-valent phenyltellurenyl bromide $PhBr_2TeTePh$ (**1**) in nearly quantitative yield, and the melting point and red-brown color are reminiscent of the compound described by Schulz and Klar (Scheme 1).^[6]



Scheme 1. Synthesis of compounds **1–5**.



The halogenation of the sterically more encumbered compound $RTeTeR$ ($R = 2,6-Mes_2C_6H_3$) with one equivalent of bromine or sulfuryl chloride produced the mixed-valent aryltellurenyl halides RX_2TeTeR (**2**, $X = Cl$; **3**, $X = Br$) in almost quantitative yield as blue and green crystalline materials, respectively (Scheme 1). Compounds **1–3** can be regarded as heavier congeners of $F_3CF_2SSCF_3$.^[2] The molecular structures of **1–3** are shown in Figure 1 and Figure 2.^[10] The $Te-Te$ bond lengths of 2.7966(5) (**1**), 2.759(6) (**2**), and 2.7835(11) Å (**3**) are somewhat longer than those of the parent compounds $PhTeTePh$ (2.712(2) Å)^[11] and $RTeTeR$ (2.711(1) Å, $R = 2,6-Mes_2C_6H_3$),^[12] whereas the average $Te-Cl$ and $Te-Br$ bond lengths 2.517(5) and 2.695(2) Å compare well with those of Ph_2TeCl_2 (2.505(3) Å) and Ph_2TeBr_2 (2.6818(6) Å), respectively.^[13] In the crystal lattice, individual molecules of $PhBr_2TeTePh$ (**1**) are associated by intermolecular $Te \cdots Br$ interactions of 3.328(4) Å that may explain the darker color and poorer solubility when compared to **2** and **3**, which lack similar contacts.

The reaction of $RTeTeR$ ($R = 2,6-Mes_2C_6H_3$) with one and with three equivalents of iodine afforded $2,6-Mes_2C_6H_3TeI$ (**4**)

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[**] $X = Cl, Br$; $R = Ph, 2,6-Mes_2C_6H_3$; $Mes = 2,4,6-Me_3C_6H_2$. We thank Mrs. Irene Brüdgam (Freie Universität Berlin) for the X-ray data collection. The Deutsche Forschungsgemeinschaft (DFG) is gratefully acknowledged for financial support.

Supporting information for this article is available on the WWW under <http://www.angewandte.org> or from the author.

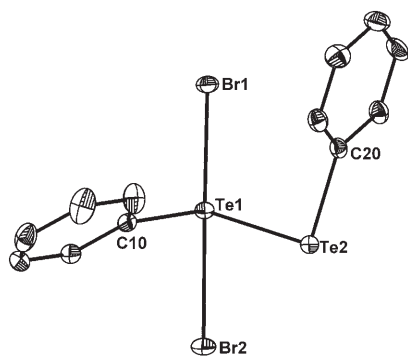


Figure 1. Molecular structure of $\text{PhBr}_2\text{TeTePh}$ (**1**) with thermal ellipsoids set at 30% probability. Hydrogen atoms have been omitted for clarity. Selected bond lengths [Å] and angles [°]: Te1–C10 2.132(4), Te1–Br1 2.5995(6), Te1–Br2 2.7842(6), Te1–Te2 2.7966(5), Te2–C20 2.108(3); C10–Te1–Te2 98.52(10), C20–Te2–Te1 97.19(9), Br1–Te1–C10 90.73(9), Br1–Te1–Te2 96.98(2), Br2–Te1–C10 87.69(9), Br2–Te1–Te2 80.299(15), Br1–Te1–Br2 176.61(2), C10–Te1–Te2–C20 98.19(14).

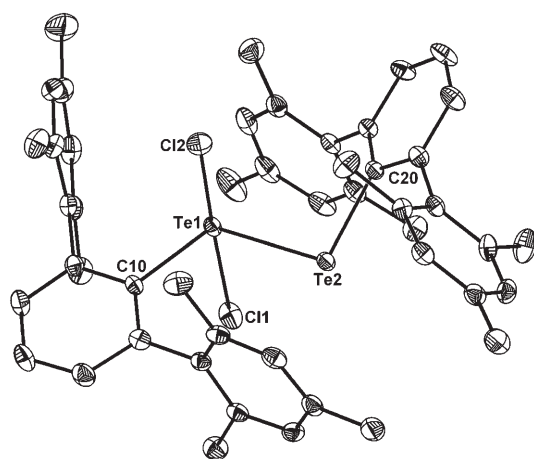


Figure 2. Molecular structure of RCl_2TeTeR (**2**, $\text{R} = 2,6\text{-Mes}_2\text{C}_6\text{H}_3$) with thermal ellipsoids set at 30% probability. Hydrogen atoms have been omitted for clarity. Selected bond lengths [Å] and angles [°]: Te1–C10 2.159(5), Te1–Cl1 2.498(5), Te1–Cl2 2.536(4), Te1–Te2 2.759(6), Te2–C20 2.149(5); C10–Te1–Te2 109.39(12), C20–Te2–Te1 96.35(12), Cl1–Te1–C10 86.68(11), Cl1–Te1–Te2 89.25(3), Cl2–Te1–C10 91.83(11), Cl2–Te1–Te2 86.08(3), Cl2–Te1–Cl1 174.33(4), C10–Te1–Te2–C20 154.71(16). The molecular structure of RBr_2TeTeR (**3**, $\text{R} = 2,6\text{-Mes}_2\text{C}_6\text{H}_3$) is almost identical. Selected bond lengths [Å] and angles [°]: Te1–C10 2.186(9), Te1–Br1 2.7011(15), Te1–Br2 2.6928(15), Te1–Te2 2.7835(11), Te2–C20 2.156(9); C10–Te1–Te2 109.2(2), C20–Te2–Te1 98.6(2), Br1–Te1–C10 87.8(2), Br1–Te1–Te2 87.53(3), Br2–Te1–C10 91.2(2), Br2–Te1–Te2 85.67(3), Br2–Te1–Br1 172.43(4), C10–Te1–Te2–C20 155.0(4).

and the charge transfer complex $2,6\text{-Mes}_2\text{C}_6\text{H}_3\text{TeI}\cdots\text{I}_2$ (**5**), respectively, as crystalline materials (Scheme 1). Complex **4** is a green solid similar to **3**, whereas the color of the charge-transfer complex **5** is red in the transmitted light of the microscope, but green with an intense metallic luster under reflective daylight (pleochroism). The molecular structures of **4** and **5** are shown in Figure 3 and Figure 4.^[10] In the solid state, two molecules of **4** form a centrosymmetric dimer, which are associated by secondary $\text{Te}\cdots\text{Te}$ contacts of 4.057(2) Å. In contrast, one of the few known aryltellurenyl

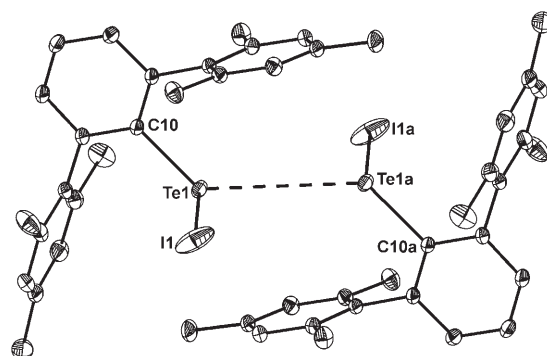


Figure 3. Molecular structure of $2,6\text{-Mes}_2\text{C}_6\text{H}_3\text{TeI}$ (**4**) with thermal ellipsoids set at 30% probability. Hydrogen atoms have been omitted for clarity. Symmetry code: $a = -x, -y, -z$. Selected bond lengths [Å] and angles [°]: C10–Te1 2.130(5), Te1–I1 2.676(1), Te1 $\cdots$ Te1a 4.057(2), C10–Te1–I1 98.58(7).

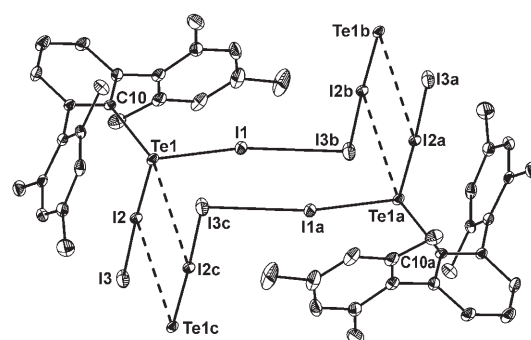


Figure 4. Molecular structure of $2,6\text{-Mes}_2\text{C}_6\text{H}_3\text{TeI}\cdots\text{I}_2$ (**5**) with thermal ellipsoids set at 30% probability. Hydrogen atoms have been omitted for clarity. Symmetry code: $b = x-1, y, z$; $c = 1-x, -y, -z$. Selected bond lengths [Å] and angles [°]: C10–Te1 2.148(5), Te1–I1 2.741(8), Te1–I2 2.839(9), I2–I3 3.003(10), I1–I3b 3.296(9); Te1 $\cdots$ I2c 3.684(1), I1–Te1–I2 91.72(2), Te1–I2–I3 177.16(2), Te1–I1–I3b 170.97(2), I1–I3b–I2b 85.43(2).

halides, $2,4,6\text{-}t\text{Bu}_3\text{C}_6\text{H}_2\text{TeI}$, has both secondary $\text{Te}\cdots\text{I}$ and $\text{I}\cdots\text{I}$ interactions,^[5b] and $2,4,6\text{-}t\text{Bu}_3\text{C}_6\text{H}_2\text{TeX}$ has only $\text{Te}\cdots\text{X}$ contacts ($\text{X} = \text{Cl}, \text{Br}$).^[5c,d] However, similar intermolecular homoatomic ($\text{Cl}\cdots\text{Cl}$) interactions were also observed in the crystal lattice of ClF .^[14] The synthesis and full characterization of the tritelluride anions $[(\text{RTe})_3]^-$ ($\text{R} = \text{Ph}, \text{CF}_3$)^[15] and the related diiodotelluride anion $[(\text{I}_2\text{TeCF}_3)]^-$ and their similarity to the triiodide ion, I_3^- have demonstrated that the RTe group has substantial pseudohalogen character, which is also evident in the charge-transfer complex **5**.

The structure of **5** is that of a 1D coordination polymer consisting of an alternating sequences of $-\text{Te}-\text{I}-\text{I}-$ chains. Adjacent 1D chains are associated by $\text{Te}\cdots\text{I}$ interactions (3.684(1) Å), whereas secondary $\text{I}\cdots\text{I}$ contacts are absent. The primary $\text{Te}-\text{I}$ bond lengths of 2.741(8) and 2.839(9) Å are slightly longer than that of $2,6\text{-Mes}_2\text{C}_6\text{H}_3\text{TeI}$ (**4**) being 2.676(1) Å. The $\text{I}-\text{I}$ bond lengths of 3.003(10) and 3.296(9) Å lie midway between intramolecular (2.715 Å) and intermolecular (3.496 Å) bond lengths of molecular iodine. The two $\text{Te}-\text{I}-\text{I}$ angles are almost linear, whereas the $\text{I}-\text{I}-\text{I}$ and the $\text{I}-\text{Te}-\text{I}$ angles are close to right angles.

Compounds **1–5** are readily soluble in most common organic solvents. At -40°C , the ^{125}Te NMR spectrum ($[\text{D}_8]\text{toluene}$) of $\text{PhBr}_2\text{TeTePh}$ (**1**) shows two equally intense, broad signals at $\delta = 1291.0$ and 823.7 ppm, which suggests that the molecular structure is retained in solution.^[16] However, no spectrum of **1** was obtained at room temperature, which points to a dynamic exchange processes taking place under these conditions. The ^{125}Te NMR spectrum (CDCl_3) of RBr_2TeTeR (**3**, $\text{R} = 2,6\text{-Mes}_2\text{C}_6\text{H}_3$) exhibits only one sharp resonance at $\delta = 1683.8$ ppm, which is consistent with the idea that **3** undergoes a reversible, presumably entropically favored rearrangement reaction to $2,6\text{-Mes}_2\text{C}_6\text{H}_3\text{TeBr}$ (**3a**) upon dissolution.^[16] The Te–Br and Te–Te bonds appear to be kinetically labile.

To estimate the relative stabilities of monomeric species REX, mixed-valent dinuclear species RX_2EER , and the disproportionation products REX_3 and REER ($\text{E} = \text{S, Se, Te}$; $\text{X} = \text{F, Cl, Br, I}$), ab initio calculations^[17,18] of suitable model compounds with $\text{R} = \text{CH}_3$ were performed; the results are summarized in Table 1. Within the fluoride series, there is

Table 1: Zero-point-energy-corrected reaction energies per chalcogen atom of the dimerization ΔE_1 and disproportionation ΔE_2 of H_3CEX .^[a]

	F	Cl	Br	I
S	–11.93 –13.40	–2.19 –0.87	–1.13 –0.06	0.05 0.27
Se	–14.42 –14.22	–6.12 –3.22	–4.50 –1.82	–8.01 –4.23
Te	–19.44 –18.74	–12.58 –8.72	–11.11 –6.93	–8.29 –4.44

[a] Upper value: ΔE_1 for dimerization, $2\text{H}_3\text{CEX} \rightarrow \text{H}_3\text{CX}_2\text{EECH}_3 + 2\Delta E_1$. Lower value: ΔE_2 for disproportionation, $3\text{H}_3\text{CEX} \rightarrow \text{H}_3\text{CEX}_3 + \text{H}_3\text{CEECH}_3 + 3\Delta E_2$. $\text{E} = \text{S, Se, Te}$; $\text{X} = \text{F, Cl, Br, I}$. Values in kcal mol^{-1} .

a strong thermodynamic driving force of the monomers H_3CEF ($\text{E} = \text{S, Se, Te}$) to undergo dimerization and disproportionation reactions to form either $\text{H}_3\text{CF}_2\text{EECH}_3$ or H_3CEF_3 and H_3CEECH_3 , respectively. For both processes, the associated reaction energies per chalcogen atom ($\Delta E_1/\Delta E_2$) are almost identical and range between -11.93 and -19.44 kcal mol^{-1} , which is consistent with the observation that an equilibrium exists between F_3CSF and $\text{F}_3\text{CF}_2\text{SSCF}_3$ prior to disproportionation to F_3CSF_3 and F_3CSSCF_3 .^[2] Amongst all other halides, the tellurenyl species H_3CTeCl and H_3CTeBr reveal the highest propensity to undergo dimerization to form $\text{H}_3\text{CCl}_2\text{TeTeCH}_3$ ($\Delta E_1 = -12.58$ kcal mol^{-1}) and $\text{H}_3\text{CBr}_2\text{TeTeCH}_3$ ($\Delta E_1 = -11.11$ kcal mol^{-1}), respectively, while the alternative disproportion reactions are less favored ($\Delta E_2 = -8.72$ and -6.93 kcal mol^{-1}). In all remaining cases, the driving force to undergo rearrangement reactions appears to be too small (ΔE_1 , $\Delta E_2 > -10$ kcal mol^{-1}), which explains the stability of most monomeric tellurenyl iodides and selenenyl halides.^[5,17]

Experimental Section

1–5: A solution of the appropriate diarylditelluride (PhTeTePh : 0.410 g, 1.00 mmol for **1**, RTeTeR ($\text{R} = 2,6\text{-Mes}_2\text{C}_6\text{H}_3$):^[12] 0.882 g, 1.00 mmol for **2–5**) in Et_2O or CH_2Cl_2 (30 mL) was cooled to 0°C and slowly treated with the appropriate halogen or synthetic equivalent (Br_2 : 160 mg, 1.00 mmol for **1**, SO_2Cl_2 : 135 mg, 1.00 mmol for **2**, Br_2 : 160 mg, 1.00 mmol for **3**, I_2 : 253 mg, 1.00 mmol for **4**, I_2 : 761 mg, 3.00 mmol for **5**). Analytically pure samples were obtained by crystallization from $\text{CH}_2\text{Cl}_2/\text{pentane}$ (**1**), diethyl ether (**2–4**) and THF (**5**). Yields 550 mg, 0.96 mmol of **1** (96% , m.p. 40°C); 858 mg, 0.90 mmol of **2** (90%); 990 mg, 0.95 mmol of **3** (95%); 1.08 g, 1.90 mmol of **4** (95%), 1.50 g, 1.82 mmol of **5** (91%). Compounds **2–5** decompose without melting.

Analytical data: **1:** ^{125}Te NMR (126 MHz, $[\text{D}_8]\text{-toluene}$, -40°C): $\delta = 1291.0$ (integral 50%), 823.7 ppm (integral 50%). UV (Et_2O , 0.1 mmol): $\lambda_{\text{max}} = 422$ nm. Elemental analysis (%) calcd for $\text{C}_{12}\text{H}_{10}\text{Br}_2\text{Te}_2$ (569.22 g mol^{-1}): C 25.32 , H 1.77 ; found: C 25.19 , H 1.71 .

2: ^{125}Te NMR (126 MHz, CDCl_3): $\delta = 1374.9$ (integral 37%), 1090.4 (integral 26%), 1027.3 ppm (integral 37%). UV (Et_2O , 0.1 mmol): $\lambda_{\text{max}} = 534$ nm. Elemental analysis (%) calcd for $\text{C}_{48}\text{H}_{50}\text{Cl}_2\text{Te}_2$ (953.09 g mol^{-1}): C 60.49 , H 5.29 ; found: C 60.20 , H 5.02 .

3: ^1H NMR (400 MHz, CDCl_3): $\delta = 7.37$ (t, $J = 7$ Hz, 1H , Ar), 7.07 (d, $J = 7$ Hz, 2H , Ph), 6.84 (s, 4H , Mes), 2.27 (s, 6H , CH_3), 1.99 ppm (s, 12H , CH_3). ^{13}C NMR (100 MHz, CDCl_3): $\delta = 148.2$, 139.0 , 137.8 , 136.4 , 130.9 , 128.4 , 128.3 , 128.0 (Ar), 21.2 , 20.8 ppm (CH_3). ^{125}Te NMR (126 MHz, CDCl_3): $\delta = 1683.8$ ppm. UV (Et_2O , 0.1 mmol): $\lambda_{\text{max}} = 559$ nm. Elemental Analysis (%) calcd for $\text{C}_{48}\text{H}_{50}\text{Br}_2\text{Te}_2$ (1042.00 g mol^{-1}): C 55.33 , H 4.84 ; found: C 54.95 , H 4.82 .

4: ^1H NMR (400 MHz, CDCl_3): $\delta = 7.49$ (t, $J = 8$ Hz, 1H , Ar), 7.14 (d, $J = 8$ Hz, 2H , Ph), 6.96 (s, 4H , Mes), 2.37 (s, 6H , CH_3), 2.07 ppm (s, 12H , CH_3). ^{13}C NMR (100 MHz, CDCl_3): $\delta = 149.6$, 140.5 , 137.6 , 136.1 , 130.9 , 128.3 , 128.0 , 112.8 (Ar), 21.2 , 21.0 ppm (CH_3). ^{125}Te NMR (126 MHz, CDCl_3): $\delta = 1018.0$ ppm. UV (Et_2O , 0.1 mmol): $\lambda_{\text{max}} = 622$ nm. Elemental analysis (%) calcd for $\text{C}_{24}\text{H}_{25}\text{I}\text{Te}$ (568.00 g mol^{-1}): C 50.75 , H 4.44 ; found: C 50.35 , H 4.41 .

5: ^1H NMR (400 MHz, CDCl_3): $\delta = 7.15$ (t, $J = 8$ Hz, 1H , Ar), 6.98 (d, $J = 8$ Hz, 2H , Ph), 6.95 (s, 4H , Mes), 2.29 (s, 6H , CH_3), 2.19 ppm (s, 12H , CH_3). ^{13}C NMR (100 MHz, CDCl_3): $\delta = 138.2$, 136.4 , 135.7 , 131.0 , 129.0 , 128.7 , 127.8 , 127.7 , (Ar), 21.2 , 21.0 ppm (CH_3). ^{125}Te NMR (126 MHz, CDCl_3): $\delta = 905.1$; (C_6D_6): $\delta = 945.6$ ppm. UV (toluene, 0.1 mmol): $\lambda_{\text{max}} = 496$ nm. Elemental analysis (%) calcd for $\text{C}_{24}\text{H}_{25}\text{I}_3\text{Te}$ (821.81 g mol^{-1}): C 35.08 , H 3.07 ; found: C 35.24 , H 3.32 .

Received: May 29, 2007

Revised: July 11, 2007

Published online: September 18, 2007

Keywords: halides · hypervalent compounds · mixed-valent compounds · structure elucidation · tellurium

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- [10] a) Crystal data for **1**: (C₁₂H₁₀Br₂Te₂·CH₂Cl₂): M_r = 651.70, monoclinic space group $P2_1/c$, a = 11.863(3), b = 13.797(3), c = 11.379(2) Å, β = 103.615(4)°, V = 1810.0(6) Å³, Z = 4, ρ_{calcd} = 2.401 mg m⁻³, crystal dimensions 0.2 × 0.1 × 0.1 mm³. 19452 collected and 3386 unique reflections. After absorption correction, this and the following structures were solved by direct methods and refined anisotropically on F^2 . Final residuals R_1 = 0.0233, wR_2 = 0.0411 ($I > 2\sigma(I)$); R_1 = 0.0452, wR_2 = 0.0508 (all data). GooF = 1.050, 229 parameters; b) Crystal data for **2** (C₄₈H₅₀Cl₂Te₂): M_r = 953.00, triclinic space group $P\bar{1}$, a = 11.6979(16), b = 14.1242(19), c = 14.871(2) Å, α = 109.732(3), β = 97.715(3)°, γ = 109.289(3), V = 2097.7(5) Å³, Z = 2, ρ_{calcd} = 1.509 mg m⁻³, crystal dimensions 0.25 × 0.11 × 0.08 mm³. 26336 collected and 8788 unique reflections. Final residuals R_1 = 0.0402, wR_2 = 0.0798 ($I > 2\sigma(I)$); R_1 = 0.0925, wR_2 = 0.1068 (all data). GooF = 1.017, 469 parameters; c) Crystal data for **3** (C₄₈H₅₀Br₂Te₂): M_r = 1041.94, monoclinic space group $P2_1/n$, a = 12.036(5), b = 27.064(11), c = 13.914(5) Å, β = 108.288(8)°, V = 4304(3) Å³, Z = 4, ρ_{calcd} = 1.608 mg m⁻³, crystal dimensions 0.40 × 0.16 × 0.015 mm³. 34641 collected and 4882 unique reflections. Final residuals R_1 = 0.0543, wR_2 = 0.1239 ($I > 2\sigma(I)$); R_1 = 0.1078, wR_2 = 0.1568 (all data). GooF = 1.018, 469 parameters; d) Crystal data for **4** (C₂₄H₂₅ITe): M_r = 567.94, monoclinic space group $P2_1/n$, a = 11.6436(19), b = 12.6391(19), c = 15.191(2) Å, β = 92.911(4)°, V = 2232.7(6) Å³, Z = 4, ρ_{calcd} = 1.690 mg m⁻³, crystal dimensions 0.55 × 0.31 × 0.08 mm³. 27262 collected and 5772 unique reflections. Final residuals R_1 = 0.0350, wR_2 = 0.0925 ($I > 2\sigma(I)$); R_1 = 0.0437, wR_2 = 0.1013 (all data). GooF = 1.050, 235 parameters; e) Crystal data for **5** (C₂₄H₂₅ITe): M_r = 821.74, monoclinic space group $P2_1/n$, a = 8.263(2), b = 14.659(4), c = 21.186(5) Å, β = 94.571(6)°, V = 2558.0(11) Å³, Z = 4, ρ_{calcd} = 2.134 mg m⁻³, crystal dimensions 0.23 × 0.11 × 0.04 mm³. 30687 collected and 5051 unique reflections. Final residuals R_1 = 0.0460, wR_2 = 0.1066 ($I > 2\sigma(I)$); R_1 = 0.0862, wR_2 = 0.1251 (all data). GooF = 1.012, 253 parameters. f) CCDC-651201 (**1**), CCDC-651202 (**2**), CCDC-651203 (**3**), CCDC-651204 (**4**), and CCDC-651205 (**5**) contain the supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif.
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