Synthesis, characterization, and ligand behaviour of a new ditelluroether \((C_{10}H_7)Te(CH_2)_4Te(C_{10}H_7)\) and the concurrently formed ionic \([(C_{10}H_7)Te(CH_2)_4]Br\)†

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The reaction of 1-naphthyl bromide with \(n\)-butyl lithium, elemental tellurium, and 1,4-dibromobutane in THF affords both \((C_{10}H_7)Te(CH_2)_4Te(C_{10}H_7)\) (1) and \([(C_{10}H_7)Te(CH_2)_4]Br\) (2) in good yields. 1 is preferentially formed at low temperatures and is a rare example of a structurally characterized ditelluroether in which the tellurium atoms are bridged by a hydrocarbon chain. In the solid state, 1 shows secondary bonding Te⋯Te interactions, which connect the molecules into layers which are further linked to 3-dimensional frameworks by Te⋯H hydrogen bonds. \([(C_{10}H_7)Te(CH_2)_4]Br\) (2) is formed concurrently during the synthesis of 1 and is the main product, when the reaction is carried out at room temperature. The revPBE/def2-TZVPP calculations of the reaction profiles indicate that the formation of 2 is somewhat more favourable than that of 1. Furthermore, at room temperature the activation energy for the formation of 2 is lower than that of 1. At low temperatures the activation energy of the reaction leading to 1 is lower than that to 2, which is consistent with the synthetic observations. When 1 was treated with CuBr, \([Cu_2(\mu-Br)_2(\mu-(C_{10}H_7)Te(CH_2)_4Te(C_{10}H_7))_2\) (3) was formed. It crystallizes as two polymorphs (3a) and (3b) in which both the packing and the conformation of the ditelluroether ligands are different. The reaction of 1 with HgCl_2 produces \([(C_{10}H_7)Te(CH_2)_4]_2[\text{HgCl}_4]\cdot\text{CH}_2\text{Cl}_2\) and that of 1 with CuCl_2 affords \([(C_{10}H_7)Te(CH_2)_4]Cl\) (5). 2 and 5 are isomorphous.

Introduction

Although the first preparation of ditelluroethers \(RTe(CH_2)_nTeR\) (\(R = 4\text{-MeO-C}_6\text{H}_4, 4\text{-EtO-C}_6\text{H}_4, C_6\text{H}_5, C_6\text{H}_5CH_2, 4\text{-Me-C}_6\text{H}_4\)) from diazomethane and ditellurides dates back to the 1970s, the chemistry of polytelluroethers has seen much slower progress than that of the related polythio- and polyselenoethers (for reviews, see ref. 2). Levason and Reid inferred that the difficulty in the preparation of di- and polytelluroether ligands is due to the inherent instability of Te–H bonds, which hinders the use of tellurols (RTeX) as synthons, the weakness of the Te–C bond leading to its facile cleavage, and the stability of the +IV oxidation state of tellurium.

In addition to the reaction between diazomethane and ditellurides, \(RTe(CH_2)_nTeR\) (\(R = \text{Ph}, 4\text{-EtOC}_6\text{H}_4, 4\text{-MeOC}_6\text{H}_4\) or \(\text{Me}_2\)) \(n = 1, 3\) have been prepared by the reaction of \(RTeX\) with organic dihalides \(X(CH_2)_nX\) (\(X = \text{Cl, Br}\)) at low temperatures. The reaction, however, is dependent on the starting materials and reaction conditions. At room temperature, the reaction of \(RTeX\) and \(X(CH_2)_nX\) (\(n = 2\) or 3) afforded \(RTeX\) and alkenes. Interestingly, it has also been reported that the reaction of \((4\text{-EtOC}_6\text{H}_4)TeNa\) and \(X(CH_2)_nX\) (\(X = \text{Br or I}; n = 3\) or 4) in aqueous ethanol does not yield the ditelluroether \((4\text{-EtOC}_6\text{H}_4)Te(CH_2)_nTe(4\text{-EtOC}_6\text{H}_4)\) because of the faster formation of the telluronium salt \([(4\text{-EtOC}_6\text{H}_4)Te(CH_2)_nX]X\). It only is with the chain lengths of \(n = 6\)–10 that the related reaction affords \((4\text{-EtOC}_6\text{H}_4)Te(CH_2)_nTe(4\text{-EtOC}_6\text{H}_4)\). In the case of \(X(CH_2)_nX\), both the ditelluroether and the telluronium halogenide are obtained. Furthermore, the treatment of \(RTeX\) with \(X(CH_2)_nX\) (\(R = \text{Me, Ph}; X = \text{Cl, Br}\)) resulted in a mixture of \(RTeX\) and cyclic tetrahydrotellurophene, \(Te(CH_2)_nTe\), while that of \(Cl(CH_2)_nCl\) resulted in a mixture of \(RTeX\), \(RTeX\), and \(Te(CH_2)_nTe\).

All the above-mentioned products have been identified and characterized by NMR and mass spectroscopy and in some cases by \(^{125}\text{Te}\) Mössbauer spectroscopy. The only known crystal structures of ditelluroethers \(RTeX\) are those of \([4\text{-MeO(C}_6\text{H}_4Te)]_2CH_2^{6a,b}\) and bis[2-fluoro-3-pyridyltelluro]methane. The crystal structures of some transition
metal complexes with chelating6,7 or bridging6,8 RTe
(CH3)nTeR (n = 1 or 3) ligands have also been determined. Monotelluroethers R2Te (R = C6H5, C4H3S, Me3SiCH2) have also been shown to form versatile metal complexes with silver(I) and copper(I).9

In this contribution, we report the low-temperature synthesis and structural characterization of naphthyl ditelluroether (C10H7)2Te(CH2)4Te(C10H7) (1). Upon prolonged reaction time and at a higher temperature, [(C10H7)Te(CH2)4Br (2) is obtained as a main product. The factors affecting the concurrent formation of 1 and 2 have been discussed on the basis of DFT calculations. We have also explored the ligand behavior of 1. When (C10H7)2Te(CH2)4Te(C10H7) (1) reacts with CuBr, an unprecedented dinuclear Cu(1) complex [Cu2(µ-Br)2(µ-(C10H7)Te(CH2)4Te(C10H7))2] (3) is formed. The reaction of 1 with HgCl2 produces [(C10H7)Te(CH2)4Te(C10H7)2][HgCl4] (4) containing a similar Te-cation to that in 2. All compounds have been characterized by single-crystal X-ray diffraction and NMR spectroscopy. The structural formulae of 1–4 have been depicted in Chart 1.

### Experimental

#### General

All reactions and manipulations of air- and moisture-sensitive reagents were carried out under an argon atmosphere. Tellurium (Aldrich), n-ButLi (Aldrich), HgCl2 (Merck), and CuCl2 (Aldrich) were used as purchased. 1,4-Dibromobutane (Aldrich) and 1-bromonaphthalene (Merck) were dried with molecular sieves and bubbled with argon prior to use. CuBr was prepared according to a literature method.10 All solvents were dried and distilled under an argon atmosphere prior to use. Tetrahydrofuran (Lab-Scan) and diethyl ether (Lab-Scan) were dried over Na/benzophenone and dichloromethane (Lab-Scan) on P2O5.

#### NMR spectroscopy

13C{1H} and 125Te NMR spectra were recorded unlocked in CH2Cl2 on a Bruker Avance III spectrometer operating at 100.62 MHz and 126.29 MHz, respectively. The typical respective spectral widths were 24.04 kHz and 125.00 kHz. The pulse widths were 4.8 μs and 8.17 μs, respectively. The 13C{1H} pulse delay was 6.50 s and that for 125Te was 1.60 s. 13C{1H} accumulations contained ca. 1000 transients and those for 125Te 30,000 transients. The 13C spectra were referenced to the solvent resonance and are reported relative to Me4Si. For 125Te, a saturated solution of H4TeO6 (aq) was used as an external reference. The 125Te chemical shifts are reported relative to neat Me4Te [δ(Me4Te) = δ(H4TeO6) + 712].11

#### X-ray crystallography

Diffraction data of 1–5 were collected on a Bruker Nonius Kappa-CCD diffractometer using graphite monochromated Mo Kα radiation (λ = 0.71073 Å; 55 kV, 25 mA). Crystal data and the details of the structure determinations are given in Table 1. Structures were solved by direct methods using SHELXS-2013 and refined using SHELXL-2013.12 After the full-matrix least-squares refinement of the non-hydrogen atoms with anisotropic thermal parameters, the hydrogen atoms were placed in calculated positions in the aromatic rings (C–H = 0.95 Å), and in the CH2 groups (C–H = 0.99 Å). The scattering factors for the neutral atoms were those incorporated with the programs.

One naphthyl group in 1 is disordered with two alternative orientations. This disorder was resolved in terms of two alternative orientations for the fused aromatic ring. The site occupation factors were refined by constraining the anisotropic displacement parameters of the disordered pairs of atoms to be equal.

#### Syntheses

(C10H7)2Te(CH2)4Te(C10H7) (1). Lithium naphthalenide was prepared according to a slightly modified literature procedure.13 1-Bromonaphthalene (1.75 mL, 12.5 mmol) was dissolved in 50 mL of diethyl ether. The solution was cooled to −40 °C and n-ButLi (5.00 mL of 2.5 M in hexanes, 12.5 mmol) was added. The solution was stirred for 1.5 hours at −40 °C during which a white precipitate of lithium naphthalenide was formed. The ether solution was decanted after which the precipitate was washed with 100 mL of dry hexane and dried under vacuum.

Lithium naphthalenide was dissolved in 50 mL of THF and the solution was cooled to −30 °C. Freshly ground tellurium (1.616 g, 12.67 mmol) was added. The solution was stirred for 1.5 hours during which the temperature of the solution rose to −25 °C, and 1,4-dibromobutane (0.75 mL, 6.28 mmol) was added. The orange red solution quickly turned yellow. After one hour the reaction solution was filtered and the solvent was evaporated. The residue was dissolved in dichloromethane and filtered. The filtrate was concentrated and cooled during which a light yellow precipitate of (C10H7)2Te(CH2)4Te(C10H7) (1) was formed. Crystallization from THF afforded light yellow crystals of 1 suitable for X-ray crystallography. Yield: 1.747 g (49%, based on n-ButLi). Anal. calc. for C24H22Te2: C 50.96, H 3.92; found: C 50.42, H 3.78%. 125Te-NMR(CH2Cl2): 352 ppm.
Table 1  Crystal data and details of the structure determinations of (C10H7)Te(CH2)4Te(C10H7) (1), [(C10H7)Te(CH2)4]Br (2), [Cu2(µ-Br)2µ-(C10H7)Te(CH2)4Te(C10H7)] (3a, 3b), [(C10H7)Te(CH2)3]Br[HgCl4] CH2Cl2 (4), and [(C10H7)Te(CH2)3]Cl (5)

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[a] I ≥ 2σ(Fo). b R2 = Σ||Fo|−|Fc||2/Σ|Fo|2, wR2 = [Σw(Fo2−Fc2)]2/ΣwFo2.  

13C-NMR(CH2Cl2): 138.1, 136.0, 133.2, 131.7, 128.6, 128.4, 126.4, 125.9, 125.8, 115.0, 33.4, 7.1 ppm.

[(C10H7)Te(CH2)4]Br (2). [(C10H7)Te(CH2)3]Br (2) was prepared as previously described for (C10H7)Te(CH2)3Te(C10H7) (1)

using 1-bromonaphthalene (1.75 mL, 12.5 mmol), n-BuLi (5.00 mL, 12.5 mmol), tellurium (1.621 g, 12.70 mmol), and 1,4-dibromobutane (0.75 mL, 6.28 mmol). In this case, the reaction mixture was stirred overnight instead of one hour. During this time the solution was allowed to warm slowly to room temperature.

The solvent in the yellow brownish reaction mixture was evaporated. The residue was dissolved in dichloromethane (60 mL), filtered, and the solvent was evaporated. A white precipitate was formed upon dissolving the residue in THF. After filtration and washing the precipitate with THF and hexane, followed by recrystallization from CH2Cl2, colourless crystals of [(C10H7)Te(CH2)3]Br (2), which were suitable for X-ray structure determination, were obtained. Yield: 0.973 g (40%, based on lithium naphthalenide).

A light yellow precipitate of (C10H7)2Te(CH2)3Te(C10H7) (1) was formed. Yield: 0.804 g (23% based on lithium naphthalenide).

[(Cu2(µ-Br)2µ-(C10H7)Te(CH2)3]Br[Cu2(µ-Br)2µ-(C10H7)Te(CH2)3] (3). (C10H7)2Te(CH2)3Te(C10H7) (1) (0.294 g, 0.520 mmol) was dissolved in dichloromethane (5 mL) and added into a flask containing CuBr (0.06 g, 0.418 mmol) and 2 mL of dichloromethane. The solution was stirred for 3 hours. A light yellow precipitate was isolated and washed with dichloromethane. Colourless crystals of [(Cu2(µ-Br)2µ-(C10H7)Te(CH2)3Te(C10H7)] (3a and 3b) suitable for single crystal X-ray diffraction were formed in a dilute CH2Cl2 solution. Yield: 0.297 g (94%, based on CuBr). Anal. calc. for C45H32BrCuTe2: C 40.65, H 3.13; found: C 40.22, H 3.14, 125Te-NMR(CH2Cl2): 660 ppm.

[(C10H7)Te(CH2)3]Br[HgCl4]-CH2Cl2 (4-CH2Cl2). (C10H7)2Te(CH2)3Te(C10H7) (1) (0.160 g, 0.283 mmol) was dissolved in dichloromethane (5 mL) and added into a flask containing HgCl2 (0.077 g, 0.284 mmol) and 2 mL of dichloromethane. The reaction mixture was stirred overnight. The cloudy bright yellow solution was filtered. Upon concentration and cooling, a light yellow precipitate was formed. The precipitate was washed with cold dichloromethane. X-ray-quality crystals of [(C10H7)2Te(CH2)3]Br[HgCl4]-CH2Cl2 (4-CH2Cl2) were obtained by recrystallization from CH2Cl2 at 0 °C. Yield: 0.081 g.
(54%, based on 1). Anal. calc. for C_{29}H_{32}Cl_{6}HgTe_{2}: C 33.20, H 3.07; found: C 33.48, H 2.98%. \(^{125}\)Te-NMR(CH_{2}Cl_{2}): 680 ppm. \([(C_{10}H_{7})Te(CH_{2})_{4}]Cl\) \(5\) (C_{10}H_{7})Te(CH_{2})_{4}Te(C_{10}H_{7}) \(1\) \((0.220 \text{ g}, 0.389 \text{ mmol})\) was added into a flask containing CuCl\(_2\) \((0.035 \text{ g}, 0.260 \text{ mmol})\) and 8 mL of dichloromethane. The reaction mixture was stirred for three hours and the cloudy bright yellow solution was filtered. The white precipitate was washed with cold dichloromethane. X-ray-quality crystals of \([(C_{10}H_{7})Te(CH_{2})_{4}]Cl\) \(5\) were obtained by recrystallization from acetonitrile at 0 °C.

**Computational details**

All structures considered in this work were optimized with the ORCA program using gradient techniques\(^{14}\) and employing a revPBE GGA functional\(^{15}\) and def2-TZVPP\(^{16}\) basis sets together with the RI approximation\(^{17}\) and Grimme’s dispersion corrections.\(^{18}\) Energies for species in THF were calculated using the COSMO polarizable continuum model\(^{19}\) implemented in ORCA. The fundamental frequencies were calculated to assess the nature of stationary points and to estimate the zero-point energy (ZPE) corrections and Gibbs reaction energies. The reported transition states correspond to structures with a single imaginary vibrational mode along the reaction coordinate.

To account for the formation of solid \([(C_{10}H_{7})Te(CH_{2})_{4}]Br\) in the reaction, the formation of the ion pair \([(C_{10}H_{7})Te(CH_{2})_{4}]^{+}\) and \(Br^{-}\) was computed at the revPBE/def2-TZVPP level and corrected for lattice effects by estimating the sublimation energy using solid-state DFT calculations. The Crystal14 program package,\(^{20}\) which utilizes periodic boundary conditions, and the PBE0 functional\(^{15a,b,21}\) were employed for solid-state DFT calculations. Triple-zeta valence basis sets that are designed for solid-state calculations and include polarization functions pob-TZVP\(^{22}\) were used for all other elements except for tellurium, for which a locally modified basis set was used.\(^{23}\) Grimme’s corrections as implemented in Crystal14 were employed to account for dispersion interactions.\(^{24}\) The Hamiltonian matrix was diagonalized in a set of \(k\)-points in a reciprocal space generated according to the Pack–Monkhorst method for sampling the first Brillouin zone with a shrinking factor \((4, 4)\). For the evaluation of Coulomb and exchange integrals (TOLINTEG), tolerance factors of 8, 8, 8, 8, and 16 were used.\(^{25}\) The default SCF convergence threshold on total energy \((10^{-7} \text{ hartree})\) was used for optimizations while for frequency calculations the threshold was increased to \(10^{-11} \text{ hartree}\). Vibrational corrections to energies for the crystal structure were determined using harmonic phonon frequencies calculated at the \(\Gamma\)-point of the Brillouin zone without considering phonon dispersion.\(^{25}\)

**Results and discussion**

**General**

At low temperatures, \((C_{10}H_{7})Te(CH_{2})_{4}Te(C_{10}H_{7})\) \(1\) was obtained in moderate yields from 1-bromonaphthalene, as shown in Scheme 1. After the addition of 1,4-dibromobutane, the workup and isolation of \(1\) was carried out in 1 h. Pure \(1\) is relatively stable under argon. In solution, the compound shows two-fold symmetry. Consequently, the NMR spectra in CH\(_2\)Cl\(_2\) exhibit one \(^{125}\)Te resonance at 352 ppm and twelve \(^{13}\)C resonances at 138.1–115.0 ppm are attributable to the carbon atoms in the naphthyl group and those at 33.4 and 7.1 ppm are due to the two inequivalent CH\(_2\) groups in the \((CH_{2})_{4}\) chain.

Upon prolonged stirring of the reaction mixture during which time the temperature was allowed to rise to ambient temperature, a white precipitate of \([(C_{10}H_{7})Te(CH_{2})_{4}]Br\) \(2\) was observed. It is stable in air. The precipitate can be dissolved in CH\(_2\)Cl\(_2\) and the NMR spectra show a \(^{125}\)Te resonance at 666 ppm, ten \(^{13}\)C resonances at 134.0–123.3, and two resonances at 39.6 and 32.5 ppm.

Our findings support the inferences of De Silva et al.\(^{42}\) that the room temperature synthesis of RTe(CH\(_2\))\(_n\)TeR \((R = p\text{-EtOC}_{6}H_{4}, \text{Ph}; n = 3, 4)\) failed because of the formation of tellurium salts due to the internal quaternarization, which was always faster than the nucleophilic attack by RTe\(^{-}\) on the second C–Br bond of the precursor Br(CH\(_2\))\(_n\)Br. We also concur that at room temperature the formation of the tellurium bromide \(2\) is faster than that of the ditelluroether \(1\) (see Scheme 1). The latter, however, seems to form more rapidly at low temperatures. In principle it is also possible that at higher temperatures the decomposition of \(1\) also takes place as shown in Scheme 2 producing additional tellurium bromide \(2\).

The reaction of \(1\) with CuBr in a molar ratio 1 : 1 produced a dinuclear complex \([Cu_{2}μ-Br]_{3}[μ-(C_{10}H_{7})Te(CH_{2})_{4}Te(C_{10}H_{7})]_{2}\) \(3\) in which two ditelluroether ligands and bromide ligands bridge two copper(i) centers (see Chart 1). 3 crystallized as two polymorphs 3a and 3b. There are a few bidental phosphane complexes, which contain similar structures where both the phosphane and halogenido ligands bridge two Cu(i) centers.\(^{26}\)
The reaction of 1 with CuCl₂ afforded colourless crystals of [[(C₁₀H₇)₂Te(CH₂)₄]Cl (5) and red crystals of [(C₁₀H₇)₂Te₂]. The crystal structure of the latter is known.²⁷

The reaction of (C₁₀H₇)₂Te(CH₂)₄Te(C₁₀H₇) (1) and HgCl₂ in CH₂Cl₂ afforded [[(C₁₀H₂)₄Te(CH₂)₄][HgCl₄]CH₂Cl₂ (4·CH₂Cl₂), which also contains the [[(C₁₀H₇)₂Te(CH₂)₄]⁺ cation like 2 and 5. The preparation of (Ph₃Te)₂[HgCl₄]²⁸ (Ph₃Te)₂[Hg₂Cl₆]²⁹ and (RTe)₂[Hg₂Cl₆] {R = 2,6-[O(CH₂CH₂)₂NCH₂]₂C₆H₃}²⁹ has also been reported, though only the [Hg₂Cl₆]²⁻ salts have been structurally characterized.

Crystal structures

(C₁₀H₂)₄Te(CH₂)₄Te(C₁₀H₂) (1) and [C₁₀H₇Te(CH₂)₄]X {X = Br (2), Cl (5); n = 1, X = Br (2), Cl (5); n = 2, X = [HgCl₄] (4)}. The molecular structure of (C₁₀H₂)₄Te(CH₂)₄Te(C₁₀H₂) (1) indicating the numbering of atoms and selected bond parameters is shown in Fig. 1.

The asymmetric unit in 1 contains two molecules. Te–C bond distances span a range of 2.118(9) Å–2.186(16) Å, which are comparable to those reported for bis(4-methoxyphenyltelluro)methane [2.13(2) Å]. The Te–C bond angles are 89.5(8)–95.0(3)°. The naphthyl group, which is bonded to Te₄, is disordered with the ring assuming two different orientations [s.o.f. 0.547(11) and 0.453(11); see Fig. 1].

Each tellurium atom is involved in two intermolecular Te⋯Te interactions and a number of H⋯Np (Np = naphthyl) π close contacts. These interactions link the discrete molecules into a three-dimensional supramolecular network (see Fig. 2).

Compounds [[(C₁₀H₂)₄Te(CH₂)₄]X [X = Br (2), Cl (5)] are isomorphous. The molecular structure indicating the labeling of atoms is shown in Fig. 3. The crystal structure of [(C₁₀H₂)₄Te(CH₂)₄][HgCl₄]CH₂Cl₂ (4·CH₂Cl₂) is shown in Fig. 4.

Fig. 1  The molecular structure of [(C₁₀H₂)₄Te(CH₂)₄Te(C₁₀H₂)] (1) indicating the numbering of atoms. Thermal ellipsoids have been drawn at 50% probability level. The atoms in the more abundant part of the naphthyl group [C₄₀A–C₄₉A; s.o.f. 0.547(11)] in 1 are shown in color, whereas those in the less abundant group [C₄₀B–C₄₉B; s.o.f. 0.453(11)] are shown in white. Hydrogen atoms have been omitted for clarity. Selected bond lengths (Å) and angles (°): Te₁–C₁ 2.161(11), Te₁–C₁₀ 2.141(12), Te₂–C₄ 2.153(15), Te₂–C₂₀ 2.186(16), Te₃–C₅ 2.164(12), Te₃–C₃₀ 2.150(14), Te₄–C₈ 2.156(13), Te₄–C₄₀A 2.118(9), Te₄–C₄₀B 2.130(9), C₁–Te₁–C₁₀ 94.4(5), C₄–Te₂–C₂₀ 92.6(6), C₅–Te₃–C₃₀ 95.0(3), C₈–Te₄–C₄₀A 94.8(7), C₈–Te₄–C₄₀B 89.5(8).

Fig. 2  The Te⋯Te and H⋯Np (Np = naphthyl) π close contacts connecting the individual [(C₁₀H₂)₄Te(CH₂)₄Te(C₁₀H₂)] (1) molecules into a three-dimensional supramolecular network.

Fig. 3  The molecular structure of [[(C₁₀H₂)₄Te(CH₂)₄]X [X = Br (2), Cl (5)] indicating the numbering of atoms. Thermal ellipsoids have been drawn at 50% probability level. Hydrogen atoms have been omitted for clarity. Selected bond lengths (Å) and angles (°): 2: Te₁–C₁ 2.130(11), Te₁–C₄ 2.122(10), Te₁–C₃₀ 2.161(10), C₁–Te₁–C₄ 85.4(4), C₁–Te₁–C₃₀ 92.2(4), C₄–Te₁–C₃₀ 96.1(4); 5: Te₁–C₁ 2.109(12), Te₁–C₄ 2.168(13), Te₁–C₃₀ 2.147(12), C₁–Te₁–C₄ 84.1(5), C₁–Te₁–C₃₀ 95.8(5), C₄–Te₁–C₃₀ 92.6(5).
The molecular structure and the crystal packing of 2 and 5 have similar features to those observed for other simple tellurium halogenide salts, as exemplified by Ph₃TeCl [31], Ph₃TeBr [32], and Ph₃TeI. [33] The three Te–C bonds in the cations of 2 and 5 are 2.122(10)–2.161(10) and 2.109(12) Å, respectively, and compare well with those in Ph₃TeX (X = Cl, Br, I). [31–33] The two Te···Br contacts in 2 or Te···Cl contacts in 5 [3.3419(5)–3.3651(5) and 3.219(3)–3.222(6) Å, respectively] expand the coordination polyhedron of the tellurium atom into a square pyramid (see Fig. 1S in the ESI†). Whereas in the solid state short Te···Cl contacts in Ph₃TeCl link the ion-pairs into dimers, [31,32] in Ph₃TeClC₂H₅ and Ph₃TeBr [31] the lattice is built up by ladder-like tetrameric units. By contrast, similar cation–anion contacts in 2 and 5 result in the formation of infinite chains. Further H···N p π interactions create three-dimensional supramolecular networks (see Fig. 1S in the ESI†).

The cation in 4–CH₂Cl₂ expectedly shows a similar structure to 2 and 5 (see Fig. 4). The cations and anions are linked into a three-dimensional network through Te···Cl contacts of 3.290(2)–3.336(3) Å. The shortest H···Cl hydrogen bonds between the cation and the anion are 2.783(2)–2.983(3) Å. Furthermore, the solvent CH₂Cl₂ molecules also show hydrogen bonds of 2.915(2)–3.303(6) Å to the tellurium cation. These solvent molecules also form dimers by two H···Cl contacts of 3.002(3) Å. The packing in the lattice is shown in Fig. 2S in the ESI.† [Cu₂(μ-Br₂)(μ-(C₉H₆)₂)Te(CH₂)₄Te(C₉H₆)₂] [3] and [Cu₂(μ-Br₂)(μ-(C₉H₆)₂)Te(CH₂)₄Te(C₉H₆)₂] [2] crystallize as two polymorphs in space groups P1 (3a) and P2₁/c (3b). Their molecular structures together with the numbering of the atoms are presented in Fig. 5(a) and (b).

The Cu–Br bonds are somewhat shorter for 3a [2.486(2)–2.5215(13) Å] and 3b [2.4433(18)–2.5495(18) Å] than for [Cu₂Br₂(dppb)₂] [dppb = bis(diphenylphosphino)butane] [2.5323(9) and 2.5726(7) Å]. [26c] The Te–Cu–Te bond angles of 3a [113.36(18) and 111.62(6)°, respectively] are significantly smaller than the P–Cu–P angles of 125.03(5)° in [Cu₂Br₂(dppb)₂].

It can be seen in Fig. 5 that the two polymorphs differ due to the conformations of the bridging ditelluroethyld ligands.

**Fig. 4** The molecular structure of [(C₁₀H₇)Te(CH₂)₄][HgCl₄]CH₂Cl₂ (4–CH₂Cl₂) indicating the numbering of the atoms. Thermal ellipsoids have been drawn at 50% probability level. Hydrogen atoms have been omitted for clarity. Selected bond lengths (Å) and angles (°): Te1–C1 2.143(8), Te1–C4 2.140(8), Te1–C10 2.148(7), Te2–C2 2.153(8), Te2–C8 2.147(8), Te2–C20 2.132(8), C1–Te1–C4 84.0(3), C1–Te1–C10 94.5(3), C4–Te1–C10 97.9(3), C5–Te1–C8 85.1(3), C5–Te2–C20 98.5(3), C8–Te2–C20 94.6(3).
The motifs in the torsional angles C10–Te1–C1–C2, Te1–C1–C2–C3, C1–C2–C3–C4, C2–C3–C4–Te2, and C3–C4–Te2–C20 are $\vartheta_1 = - - - +$ and $\vartheta_2 = + - - +$ for 3a, and 3b, respectively (+ indicates the clockwise rotation about the central bond defining the torsional angle and − the anticlockwise rotation; for actual metrical values, see Fig. 5). This leads to differences in packing, as shown in Fig. 3S in the ESI.† The shortest Br⋯H hydrogen bond in 3a is 3.0754(14) Å and those in 3b are 2.9072(14) and 3.0439(12) Å.

Formation of (C10H7)Te(CH2)4Te(C10H7) (1) and [C10H7Te(CH2)4]X (n = 1, X = Br (2), Cl (5); n = 2, X = [HgCl4] (4))

Although the syntheses of RTe(CH2)nTeR (R = Me or Ph; n = 1, 3) by the reaction of RTe with organic dihalides X(CH2)nX (X = Cl, Br) at low temperatures have been reported,4,5,6,7 the reaction involving X(CH2)nX at room temperature seems to result in the preferential formation of the ionic telluronium halogenide [RTe(CH2)n]X (R = p-ETOOC6H4 or Ph).4,5,6,7,8 We have also observed that (C10H7)Te(CH2)4Te(C10H7) (1) can be prepared with a moderate yield at −25 °C, but as the temperature of the reaction mixture rises to room temperature, an equally good yield of [(C10H7)Te(CH2)3]Br (2) together with some 1 can be obtained.

Since it was possible that the prolonged reaction time even at low temperatures could result in the formation of [(C10H7)Te(CH2)3]Br (2) in addition to that of (C10H7)Te(CH2)4Te(C10H7) (1), we carried out two small-scale experiments in THF at −40 °C and monitored the products by 125Te NMR spectroscopy. In one experiment, the reaction was discontinued after one hour. The reaction solution in THF was slightly cloudy, and the 125Te NMR spectrum showed the main resonance at 352 ppm, which is due to 1, with only a trace of the resonance at 666 ppm, which is due to the [(C10H7)Te(CH2)4]X cation of 2. The latter resonance is known to be slightly soluble in THF. Another reaction was discontinued after six hours. In this solution, there was a clear presence of the precipitate of 2. The 125Te NMR spectrum showed that the concentration of 2 had also increased somewhat during the prolonged stirring time at −40 °C, though 1 was still the main product. It can be concluded that both 1 and 2 are formed also at low temperatures, but under these conditions the formation of 1 is faster than that of 2.

Since it is conceivable that the formation of 1 and 2 can take place concurrently, as shown in Scheme 1, or sequentially, as shown in Scheme 2, we decided to carry out revPBE/def2-TZVPP calculations both at 298 K and at 200 K. The energy profiles of the three different reactions in question at room temperature (298 K) are shown in Fig. 6.

The first step in both reactions is the formation of acyclic (C10H7)Te(CH2)4Br from Li+[(C10H7)Te(CH2)4]− and Br(CH2)4Br. This reaction is relatively straight-forward and since it is common for the formation of both 1 and 2, we used it as a starting point for our calculations. It can be seen from the Gibbs energy values shown in Fig. 6 that in the formation of both 1 and 2, the driving force is the formation of solid salts, namely LiBr(s) in the case of 1, and that of 2 itself. The activation energies are reasonably low and relatively close to each other with that for the reaction leading to 1 (shown in green) slightly higher than that for the reaction leading to 2 (shown in red). The total energy of the formation of 2 is somewhat more favourable than that of 1. Furthermore, the conversion of 1 to

![Fig. 6 revPBE/def2-TZVPP reaction profiles of the formation of (C10H7)Te(CH2)4Te(C10H7) (1) (shown in green) and of [C10H7]Te(CH2)4+Br (2) (shown in red) from C10H7Te(CH2)4Br. The reaction profile of the decomposition of 1 to 2 (shown in blue). Each of the three transition states T51–T53 show only one imaginary frequency.](image-url)
2 goes via a higher activation barrier (shown in blue). All these findings support the earlier conclusion that at room temperature the quaternation at tellurium and the formation of the telluronium salt are faster than the reaction of RTe(CH2)4Br with another equivalent of RTe−.4a

We have also computed the activation barriers for reactions via the transition states TS1 and TS2 at 200 K. The activation energy for the formation of ditelluroether 1 (transition state TS1) was lowered to 32.1 kJ mol−1, while the activation barriers for the formation of the telluronium bromide 2 (transition state TS2) and the decomposition of the ditelluroether 1 to the telluronium bromide 2 (transition state TS3) remained approximately unchanged. This is consistent with our experimental observation that at low temperatures the reaction of (C10H7)Te− with Br(CH2)4Br affords (C10H7)Te(CH2)4Te(C10H7) (1) faster than [(C10H7)Te(CH2)4]Br (2). While in principle spontaneous, the decomposition of 1 to 2 does not seem to be a significant alternative route.

However, we tested the possibility of the conversion of 1 to 2 by adding an excess of LiBr to the THF solution of 1 and stirring the mixture for several days. Only trace amounts of 2 were formed. It further supports the conclusion that 2 is mainly formed concurrently with 1 and that the relative rates of the formation of the two products are dependent on the temperature.

In contrast, the treatment of 1 with HgCl2 resulted in the formation of [(C10H7)Te(CH2)4]2[HgCl4] (4) in a good yield. The likely reaction is presented in eqn (1).

\[
2(C_{10}H_7)Te(CH_2)_4Te(C_{10}H_7) + 2HgCl_2 \rightarrow [(C_{10}H_7)Te(CH_2)_4]_2[HgCl_4] + [Hg(Te(C_{10}H_7))]_2
\]

(1)

It is possible that the reaction is fast due to the catalytic effect of mercury resulting in the formation of the salt 4. We also tested a related reaction by treating 1 with CuCl2. The 125Te NMR spectrum of the reaction mixture in CH2Cl2 indicates that this reaction affords the chloride salt 5 and approximately a half equivalent of (C10H7)TeTe(C10H7). Both products could be isolated and structurally characterized. A possible path of the reaction involves the reduction of CuII with the consequent oxidation of (C10H7)Te− to (C10H7)TeTe(C10H7) (see eqn (2)):

\[
(C_{10}H_7)Te(CH_2)_4Te(C_{10}H_7) + CuCl_2 \rightarrow [(C_{10}H_7)Te(CH_2)_4]Cl + \frac{1}{2}(C_{10}H_7)TeTe(C_{10}H_7) + CuCl
\]

(2)

Conclusions

The low-temperature preparation of a rare new ditelluroether (C10H7)Te(CH2)4Te(C10H7) (1) from naphthyl bromide, n-butyl lithium, elemental tellurium, and 1,4-dibromobutane has been described. The crystal structure of 1 shows two independent molecules, which are linked by Te−−Te secondary bonding interactions and Te−−H hydrogen bonds.

Upon prolonged stirring of the reaction solution during which time the temperature rose to the room temperature, a telluronium salt [(C10H7)Te(CH2)4]Br (2) was observed and isolated. A similar cation is obtained in the reactions of 1 with M(II) metal salts HgCl2 and CuCl2, which form [(C10H7)Te(CH2)4][HgCl4] (4) and [(C10H7)Te(CH2)4]Cl (5), respectively. The two [C10H7Te(CH2)4X [X = Br (2), Cl (5)] salts are isomorphous. The cation–anion contacts in 2 and 5 result in the formation of infinite chains.

The dinuclear complex [Cu2(µ-Br)2(µ-(C10H7)Te(CH2)4Te(C10H7))] (3) was formed by the reaction of 1 with CuBr. It crystallizes as two polymorphs 3a and 3b, which both show a similar bridging arrangement of both the ditelluroether ligands and bromides between two Cu(II) centers. There are, however, marked differences in the conformations of the ditelluroether ligands and in the packing of the complexes.

The revPBE/def2-TZVPP calculations of the reaction profiles indicate that 1 and 2 are formed concurrently. While both reactions are spontaneous, the total Gibbs energy change for the formation of 1 is somewhat less favourable than for that of 2. The driving force for the reactions is the formation of solid LiBr in the case of 1 and the salt 2 itself in the latter case. While at room temperature the activation energy in the formation of 1 is higher than that of 2, at low temperatures the situation is reversed. It can therefore be concluded that at low temperatures the formation of 1 from (C10H7)Te− and Br(CH2)4Br is faster than that of 2, but at the room temperature the telluronium salt 2 is somewhat favoured over the ditelluroether 1. While the calculations indicate that the decomposition of 1 to 2 upon interaction with Br− is energetically possible, the activation energy of the reaction is somewhat higher than in the concurrent reactions, which renders the decomposition of 1 less likely. Furthermore, a separate experiment has shown that there is indeed little or no evidence for the formation of 2 from 1.

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Notes and references