Diaryldichalcogenide radical cations†‡

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One-electron oxidation of two series of diaryldichalcogenides [(C6F5S)2] (13a–c) and [(C6F5Se)2] (16a–c) was studied (E = S, Se, Te). The reaction of 13a and 13b with AsF5 and SbF5 gave rise to the formation of thermally unstable radical cations [(C6F5S)2] + (14a) and [(C6F5Se)2] + (14b) that were isolated as [SbF6] and [AsF6] salts, respectively. The reaction of 13c with AsF5 afforded only the product of a Te–C bond cleavage, namely the previously known dication [Te2]2+ that was isolated as [AsF6]− salt. The reaction of (2,6-Mes2C6H3E)2 (16a–c) with [NO][SbF6] provided the corresponding radical cations [(2,6-Mes2C6H3E)2] + (17a–c; E = S, Se, Te) in the form of thermally stable [SbF6]− salts in nearly quantitative yields. The electronic and structural properties of these radical cations were probed by X-ray diffraction analysis, EPR spectroscopy, and density functional theory calculations and other methods.

Introduction

The landmark paper by Gomberg on the stable free triphenylmethyl radical initiated numerous investigations on molecules containing unpaired electrons.1 Several kinds of persistent and stable radicals2 have been described ever since, and several more general classes of (poly)radicals have been developed in recent times.3 As stated recently, “much of the current interest in stable radicals probably arises […] from the fundamental structure and bonding issues that naturally arise with this class of compounds”.4 Most often, light carbon-based or heteroatom radicals have been studied. Investigations on heavier main group radicals are fewer in number, although the last decades witnessed spectacular discoveries in the chemistry of the heavier main group elements.5 The design of novel synthetic strategies, particularly the use of very bulky substituents, has led to the isolation of a wide range of compounds, including main group radicals, an area which has been reviewed recently.6 The interest in the one-electron oxidation of diorganodichalcogenides dates back to 1868 when it was observed that (PhS)2 (1a) dissolves in conc. H2SO4 to give intensively coloured solutions of radical cations (Scheme 1).7 Nowadays, it is understood that the formation of the latter begins with the one-electron oxidation of (PhS)2 (1a) providing the intermediate radical cation [(PhS)2]•+ (2a), which upon loss of another electron gives rise to the intermediate dication [(PhS)2]2+ (3). Charge repulsion (“Coulomb explosion”) leads to the dissociation into two sulfenyl cations [PhS]+ (4), which undergo mutual electrophilic substitution of their phenyl rings

[Diagram of radical cations and their reactions]

Scheme 1 Oxidation of diphenylsulfide in conc. sulfuric acid.

† Dedicated to Professor Hubert Schmidbaur on the occasion of his 80th birthday.
in ortho-position to produce thianthrene (5). Another one-electron oxidation eventually yields the thianthrene radical cation (6) and unaccounted products (Scheme 1). The one-electron oxidation of (PhS)₂ (1a) and its Se-congener (PhSe)₂ (1b) in the confined voids of the acidic pentasil zeolite allowed the tentative characterization of the radical cations [(PhS)₂]⁺ (2a) and [(PhSe)₂]⁺ (2b) by EPR spectroscopy. Previous attempts to prepare a persistent dialkyldisulfide radical cation using (NeoS)₂ [7; Neo = neopentyl] and nitrosyl triflate [NO][O₂SCF₃] afforded a dialkyldisulfide nitrosium adduct [(NeoS)₂NO]⁺ (8) comprising a four-membered ring structure (Scheme 2). The related dianionic dicationic rings [(MeSe)₄]²⁻ (9) and [(EtTe)₄]²⁻ (10) were similarly obtained by the one-electron oxidation of heavier group 16 dialkyldichalcogenides (MeSe)₂ (11) and (EtTe)₂ (12) with [NO][O₂SCF₃]. Compounds 9 and 10 can be regarded as dimers of persistent radical cations, which dimerise by π→π* interactions. The dissociation energies of these rings were estimated to be in the order of magnitude of 50 kcal mol⁻¹, which prompted us to investigate if persistent or even stable dialkyldichalcogenide radical cations [(RE)₂]⁺ can be prepared using fluorinated (R = C₆F₅) or bulky aromatic substituents (R = 2,6-Mes₂C₆H₄).

Results and discussion

In an initial foray, we investigated the one-electron oxidation of bis[pentafluorophenyl]dichalcogenides with antimony pentafluoride and arsenic pentafluoride. The reaction of (C₆F₅S)₂ (13a) and (C₆F₅Se)₂ (13b), respectively, with SbF₅ or AsF₅ gave an immediate color change to dark blue and dark green upon contact, which suggested that paramagnetic species had formed (Scheme 3). Unfortunately, almost all attempts to isolate crystalline products by crystallisation from SO₂ClF at low temperatures using co-solvents such as aHF, F₁₁₄ and CFCl₃ were precluding any detailed spectroscopic characterisation. It is noted that the reaction of (C₆F₅S)₂ (13a) with AsF₅ in liquid SO₂ was studied previously at room temperature and provided (C₆F₅S)₂SCF₃AsF₅. The reaction of (C₆F₅Te)₂ (13c) with SbF₅ or AsF₅ under similar conditions gave no indication for the formation of radicals. The isolation of small amounts of the previously known [Te₄]²⁺ (15; counterion [AsF₆]⁻) provided evidence that Te–C bonds were cleaved.

The oxidation of the bulky bis(m-terphenyl)dichalcogenides were studied by cyclic voltammetry first. At a stationary Pt electrode, electrochemical oxidation of 16a–c in CH₂Cl₂/0.1 M [n-Bu₄N][BF₄] within the potential sweep range 0 < E < 1.5 V is characterized by a one-electron quasi-reversible peak (Ep* - Epox = 0.07 – 0.08 V, Epox - Epoxp = 0.06 V, Epox/Epoxp ≈ 0.8 – 0.9). The observed quasi-reversibility of the peak indicates relative stability of the radical cations. Significant differences in the peak currents for 16a–c can be tentatively attributed to the differences in their diffusion coefficients. The anodic potentials with respect to a saturated calomel electrode of (2,6-Mes₂C₆H₄)₂ (16a, E = S) over 1.09 V (16b, E = Se) to 0.79 V (16c, E = Te), respectively, which suggested nitrosonium salts to be suitable one-electron oxidizers. Indeed, the reaction of the bis(m-terphenyl)dichalcogenides (2,6-Mes₂C₆H₄)₂ (16a, E = S; 16b, E = Se; 16c, E = Te) with [NO][SbF₆] in propionitrile provided the corresponding radical cations [(2,6-Mes₂C₆H₄)₂]⁺ (17a, E = S; 17b, E = Se; 17c, E = Te) as dark blue crystals in very high yields, which showed no signs of decomposition for several months when isolated from the mother liquor and stored under argon (Scheme 3).

The molecular structures of [(C₆F₅S)₂]⁺ (14a, E = S; 14b, E = Se) and [(C₆F₅Se)₂]⁺ (17a, E = S; 17b, E = Se; 17c, E = Te) are shown in Fig. 1. Selected bond parameters are collected in Table 1 together with those of the neutral parent compounds. The radical cations 14a, 14b, 17a and 17b containing S and Se atoms adopt nearly C₂ symmetry conformations. The phenyl rings comprising C10–C15 are almost coplanar with the C10–E₁–E₂ plane pointing to delocalization of unpaired electron spin density along the aromatic π-system, whereas the phenyl rings including C20–C25 are perpendicular to the C20–E₁–E₂ plane (E = S, Se). The radical cation 16c containing Te atoms is centrosymmetric and possesses C₂ symmetry. Consequently, only one crystallographically independent Te atom is present. In the radical cations [(C₆F₅S)₂]⁺ (14a) and [(2,6-Mes₂C₆H₄)₂]⁺ (17a) the delocalization is also reflected in the S–C bond lengths; S1–C10 (14a, 1.727(8) Å; 17a, 1.762(5) Å) is significantly shorter than S1–C20 (14a, 1.764(8); 17a, 1.799(5) Å) pointing to a quinoid structure of the coplanar phenyl ring (C10–C15). Indeed, the quinoid character of the coplanar phenyl rings of 14a (Q = 0.036 (26%)), 14b (Q = 0.012 (9%)), 17a (Q = 0.019 (14%)) and 17b (Q = 0.019 (14%)) is substantially higher than that of the perpendicular phenyl rings of 14a (Q = 0.024 (17%)), 14b (Q = 0.001 (<1%)), 17a (Q = 0.001 (<1%)) and 17b (Q = 0.002
Fig. 1 Molecular structures of the radical cations 14a,b and 17a–c; thermal ellipsoids are set at 30% probability.

Comparison of the parent compounds with the corresponding radical cations reveals a shortening of the E–E bonds by 0.008 Å for 13a/14a, 0.030 Å for 13b/14b, 0.075 Å for 16a/17a, 0.050 Å for 16b/17b and 0.049 Å for 16c/17c (E = Te), respectively. The E–E bond shortening unambiguously suggests that the bond order increased as electron density from π*-orbitals of the chalcogens has been depleted upon oxidation. The increase of the E–E bond orders should be also reflected by an increase of the E–E stretching vibrations, however, all attempts to obtain reasonable Raman spectra failed due to the intense colour of the compounds. The most striking structural difference upon going from the parent compounds to the radical cations is the dramatic increase of the C–E–C torsion angles from 84.6(2)° to 175.8(4)° for 13a/14a, 127.2(1) to 174.6(3)° for 16a/17a (E = S), 75.3(1) to 178.1(1)° for 13b/14b, 128.3(2) to 172.6(5)° for 16b/17b (E = Se) and 123.1(1) to 155.5(3)° for 16c/17c (E = Te), respectively. The positive charges of 17a–c seem to be compensated by intramolecular Menshutkin interactions between chalcogen atoms and mesityl groups of the m-terphenyl substituents (E=Zn ca. 3 Å; Zn = centroid of the phenyl ring). The neutral parent compounds exhibit interactions which are substantially longer (E=Zn ca. 3.4 Å). Presumably for the same reason, 14a and 14b possesses a short intramolecular S–F (2.712(6) Å) and Se–F (2.770(2) Å) contacts. These structural changes upon oxidation were satisfactorily reproduced by DFT calculations on two series of parent compounds, namely (PhE)2 (1a–c) and (Cp3E)2 (13a–c), and radical cations, namely [PhE]3+] (2a–c)] and [(Cp3F3E)2]3+ (14a–c) for E = S, Se, Te. The adiabatic ionization energies of 1a–c (6.98–7.38 eV) are lower than those of 13a–c

Table 1 Selected bond parameters [Å, °] of the parent diarylchalcogenides 13a–b, 16a–c and the corresponding radical cations 14a–b, 17a–c (italics)

<table>
<thead>
<tr>
<th></th>
<th>13a (ref. 18)/14a, E = S, R = C6F6</th>
<th>13b (ref. 18)/14b, E = Se, R = C6F6</th>
<th>16a/17a, E = S, R = m-Ter</th>
<th>16b (ref. 19)/17b, E = Se, R = m-Ter</th>
<th>16c (ref. 20)/17c, E = Te, R = m-Ter</th>
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</thead>
<tbody>
<tr>
<td>E–E</td>
<td>2.022(2)</td>
<td>2.319(4)</td>
<td>2.073(1)</td>
<td>2.339(2)</td>
<td>2.711(1)</td>
</tr>
<tr>
<td>C–E</td>
<td>2.014(3)</td>
<td>2.289(1)</td>
<td>1.998(2)</td>
<td>2.289(7)</td>
<td>2.662(1)</td>
</tr>
<tr>
<td>C–E–E</td>
<td>1.785(6)</td>
<td>1.96(2)</td>
<td>1.790(2)</td>
<td>1.926(6)</td>
<td>2.144(3)</td>
</tr>
<tr>
<td></td>
<td>1.976(6)</td>
<td>1.92(1)</td>
<td>1.787(2)</td>
<td>1.926(7)</td>
<td>2.151(3)</td>
</tr>
<tr>
<td></td>
<td>1.727(8)</td>
<td>1.876(2)</td>
<td>1.762(5)</td>
<td>1.94(1)</td>
<td>2.131(8)</td>
</tr>
<tr>
<td></td>
<td>1.764(8)</td>
<td>1.904(2)</td>
<td>1.799(5)</td>
<td>1.947(8)</td>
<td></td>
</tr>
<tr>
<td>C–E–E</td>
<td>104.5(2)</td>
<td>98.7(3)</td>
<td>104.8(1)</td>
<td>102.3(2)</td>
<td>103.1(3)</td>
</tr>
<tr>
<td></td>
<td>106.2(2)</td>
<td>98.9(6)</td>
<td>104.7(1)</td>
<td>102.3(3)</td>
<td>103.0(1)</td>
</tr>
<tr>
<td></td>
<td>96.2(3)</td>
<td>93.7(7)</td>
<td>95.4(2)</td>
<td>95.9(3)</td>
<td>98.5(2)</td>
</tr>
<tr>
<td></td>
<td>107.5(3)</td>
<td>103.9(1)</td>
<td>110.4(2)</td>
<td>105.6(3)</td>
<td></td>
</tr>
<tr>
<td>Q(C10–C13)a</td>
<td>0.036 (26%)</td>
<td>0.012 (9%)</td>
<td>0.019 (14%)</td>
<td>0.019 (14%)</td>
<td>0.037 (27%)</td>
</tr>
<tr>
<td>Q(C20–C25)a</td>
<td>0.024 (17%)</td>
<td>0.001 (&lt;1%)</td>
<td>0.001 (&lt;1%)</td>
<td>0.002 (&lt;1%)</td>
<td></td>
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<tr>
<td>E–Zn</td>
<td>—</td>
<td>—</td>
<td>3.439(1)</td>
<td>3.452(2)</td>
<td>3.377(1)</td>
</tr>
<tr>
<td></td>
<td>—</td>
<td>2.975(2)</td>
<td>3.01(1)</td>
<td>3.481(1)</td>
<td>3.161(2)</td>
</tr>
<tr>
<td>E···F</td>
<td>2.712(6)</td>
<td>2.770(2)</td>
<td>—</td>
<td>—</td>
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</table>

a Quinoid character is defined as Q(C20–C25) = \( \frac{d_{C12–C13} + d_{C12–C14} + d_{C12–C15} + d_{C20–C25}}{4} \) – \( \frac{d_{C21–C24} + d_{C21–C25} + d_{C22–C24} + d_{C22–C25}}{4} \) and 0 for a perfectly delocalized hexagonal benzene structure and 0.138 for a perfect quinoid structure where \( d_{C12–C13} = d_{C12–C14} = d_{C12–C15} = 1.455 \text{ Å} \) and \( d_{C21–C24} = d_{C21–C25} = d_{C22–C24} = 1.317 \text{ Å} \).

b Element distance to the centroid of the phenyl ring E–Zn.
(7.64–8.16 eV) and follow the same trend as the cathodic peak potentials of series \( \{2,6\text{-Mes}\text{C}_6\text{H}_3\text{E}\}_2 \) for E = S, Se, Te. In contrast to the neutral (PhE)\( _2 \) (1a–c) that all have a C\( _2 \) \( (\text{H}_2\text{O}_2 \) type) structure with dihedral angles around 90°, all radical cations are calculated having much larger dihedral angles. The phenyl-substituted radical cations [PhE]\( _2 \)^{+} (2a–c), can be considered as essentially freely rotating around the E–E bonds, with a minimum energy at dihedral angles around 160° and a C\( _2 \)-symmetric structure. The experimentally found C\( _2 \) symmetric structure is less than 1 kcal mol\(^{-1} \) higher in energy and corresponds to a calculated transition state (one imaginary frequency \( > 10 \text{ cm}^{-1} \)). The size of the butterfly shaped \( m \)-terphenyl groups easily explains why the radical cations \([2,6\text{-Mes}\text{C}_6\text{H}_3\text{E}]_2\)\(^{+} \) (17a–b; E = S, Se) have C\( _2 \) symmetry, simply for reasons of s terical crowding. In the pentafluorophenyl substituted radical cations \([\text{CF}_3\text{F}_2\text{E}]_2\)\(^{+} \) (14a–c) with E = S and Se, the experimentally observed C\( _2 \) symmetry is the ground state, and the C\( _2 \) symmetric structure is about 1.5 kcal mol\(^{-1} \) higher in energy. Interestingly, \([\text{CF}_3\text{F}_2\text{Te}]_2\)\(^{+} \) (14c) behaves differently. Its ground state is C\( _2 \) symmetric, completely flat, with a dihedral angle of 180°. The C\( _2 \) symmetric structure is again only <0.5 kcal mol\(^{-1} \) higher in energy. It is unclear whether this peculiar structure is the reason why it has not been possible to isolate it. The HOMO of (\( \text{CF}_3\text{F}_2\text{Se} \))\(_2 \) (13b) is an admixture of components situated at the Se atoms and the \( \pi \)-system of the pentafluorophenyl groups, whereas in the SOMO of the corresponding radical cation \([\text{CF}_3\text{F}_2\text{Se}]_2\)\(^{+} \) (14b), the unpaired electron is strongly distributed over the coplanar phenyl group (Fig. 2).

The Wiberg bond indices (WBIs) of the E–E bonds increase from about 1 from 1 for the parent compounds 1a–c and 13a–c to values between 1.216 and 1.315 for the [PhE]\( _2 \)^{+} (2a–c; E = S, Se, Te) series and between 1.176 and 1.282 for the \([\text{CF}_3\text{F}_2\text{E}]_2\)\(^{+} \) (14a–c; E = S, Se, Te) series of radical cations. In the absence of electron delocalization across the coplanar phenyl rings the WBIs of the radical cations would have been expected to be close to 1.5.

The stability of 17a–c in solution dramatically depends on the solvents and in \( \text{CH}_2\text{Cl}_2 \) on the nature of the chalcogen. Inkblue solutions of 17a and 17b in \( \text{CH}_2\text{Cl}_2 \) are stable for months, whereas purple-blue 17c decomposes within a few days. The blue colour arises from very broad absorptions in the near IR region, which stretch into the visible range. Only 17c shows an absorption maximum (\( \text{CH}_2\text{Cl}_2 \)) at \( \lambda_{\text{max}} \) = 583 nm in the visible range that is only slightly shifted compared to that of 16c (553 nm) and responsible for the purple tinge. The absorption is tentatively assigned to a n(\( \text{Te} \)) \( \rightarrow \) \( \sigma^* \) transition.\(^{22} \) Dark blue solutions of 17a–c in acetonitrile and propionitrile are stable only for a few days. Notably, after some time the parent compounds 16a–c slowly form back and eventually precipitate. Electrospray mass spectra (MeCN, positive mode) of 17a–c show prominent mass clusters at \( m/z \) = 690.4, 786.3 and 882.2, which were unambiguously assigned to the radical cations on the basis of the correct isotopic patterns. The molecular conductivities (MeCN, \( c = 3 \times 10^{-2} \text{ mol L}^{-1} \) ) of \( \lambda \) = 1800, 600 and 340 \( \Omega^{-1} \text{ cm}^{-1} \) confirm a high concentration of electrolytes in solutions of 17a–c. The one-electron oxidation upon going from \([2,6\text{-Mes}\text{C}_6\text{H}_3\text{E}]_2 \) (16b, E = Se; 16c, E = Te) to the radical cation \([2,6\text{-Mes}\text{C}_6\text{H}_3\text{E}]_2\)\(^{+} \) (17b, E = Se; 17c, E = Te) is well reflected by multinuclear NMR spectroscopy. The 77Se NMR spectra show signals at \( \delta = 426.2 \) (CDCl\(_3 \)) for 16b and at \( \delta = 1362.3 \) (CDCl\(_3 \), 223 K) and 1362.4 (MeCN) for 17b, respectively. The 125Te NMR spectra exhibit signals at \( \delta = 322.2 \) (CDCl\(_3 \)) for 16c and at \( \delta = 1703.8 \) (CDCl\(_3 \)) and 1698.7 (MeCN) for 17c, respectively. These values are in good agreement with those calculated for (PhSe)\(_2 \) (1b, \( \delta = 451.5 \)), [(PhSe)\(_2\)\(^{+} \) (2b, \( \delta = 1237.2 \) and 1493.0; average 1365.1), (PhTe)\(_2 \) (1c, \( \delta = 185.1 \)) and [(PhTe)\(_2\)\(^{+} \) (1c, \( \delta = 1346.8 \) and 1778.9; average 1562.9). \(^1\)H and \(^{13}\)C NMR spectra gave expectedly broad signals. The paramagnetism of the radical cations was unambiguously confirmed by EPR spectroscopy and SQUID magnetometry. X- and Q-band field-sweep EPR spectra measured in frozen \( \text{CH}_2\text{Cl}_2/\text{THF} \) solution (1:1) show that all three species have \( g \) values that deviate significantly from the free-electron value. All radicals exhibit a rhombic g-matrix with the same \( g \)-value ordering, inferring the electronic structure of the three radicals is very similar. The \( g \) anisotropy for the radical cations \([2,6\text{-Mes}\text{C}_6\text{H}_3\text{E}]_2\)\(^{+} \) increases in the order S (17a) < Se (17b) < Te (17c), as expected from the increasing spin–orbit coupling constant and for radicals where the spin density is predominately located on the central dichalcogen moiety. The EPR spectra of the \([2,6\text{-Mes}\text{C}_6\text{H}_3\text{E}]_2\)\(^{+} \) radical cations feature a single proton hyperfine coupling resolved at \( g_1 \) and \( g_2 \) and this coupling was fully characterized by \(^1\)H Davies ENDOR spectra recorded at Q-band (Fig. 3, see ESI† for details). Simulation of the data yielded the hyperfine matrix (Table 2, see ESI† for details), which reveals a small but significant isotropic component of \( |a_{1\text{a\text{o}}} | = 7.1 \text{ MHz} \) \( (a_{1\text{a\text{o}}} = (a_1 + a_2 + a_3)/3) \), proving a small amount of spin density delocalises onto one of the \( m \)-terphenyl ligands.\(^{23} \) The findings were supported by spin density calculation (B3LYP/def2-TZVP) showing noticeable negative spin density on the para-H atom of one \( m \)-terphenyl ring (Fig. 4). The EPR spectra of the \([2,6\text{-Mes}\text{C}_6\text{H}_3\text{Se}]_2\)\(^{+} \) (17b) at X- (Fig. 3 top) and Q-band (Fig. 3 bottom) exhibit two distinct Se hyperfine couplings along \( g_1 \) (\( 77\text{Se}, S = 1/2 \), 7.6% abundance), consistent with a radical containing a Se–Se moiety with a small asymmetry in the electronic structure. Along \( g_2 \) and \( g_{1\text{a\text{o}}} \), \( 77\text{Se} \) hyperfine splittings were partially resolved and enabled an estimation of the remaining principal values of the two hyperfine interactions. Q-band Davies ENDOR was used to investigate the proton hyperfine interactions (see ESI† for details) which revealed (again) one largest resolved proton hyperfine interaction. The latter is, however, smaller than in the case of 17a, with \( |a_{1\text{a\text{o}}} | = 4.2 \text{ MHz} \). This result is consistent with the small
corresponding simulations. 

The EPR spectra of $^{17b}$ has a structure intermediate between equivalent but relatively small delocalisation of the spin constituent have essentially the same orientation with respect to the $^{17c}$ carrying the unpaired electron and less spin delocalisation. For orientation and thus there is poor overlap with the $^{35S}$ moiety and thus facilitates spin density delocalization. In contrast, the phenyl group of the second $m$-terphenyl substituent is approximately at 90° to this orientation and thus there is poor overlap with the $S$-$S$ moiety and thus facilitates spin density delocalization. In contrast, the phenyl group of the $m$-terphenyl substituent is well orientated to allow overlap with the $\pi$-type orbitals carrying the unpaired electron on the $S$-$S$ moiety and thus facilitates spin density delocalization. In contrast, the phenyl group of the $m$-terphenyl substituent is approximately at 90° to this orientation and thus there is poor overlap with the $S$-$S$ moiety and thus facilitates spin density delocalization.

Table 2 Experimental and DFT (in parentheses) EPR parameters for $^{17a}$–c. Hyperfine values are in MHz

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Principal values</th>
</tr>
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<tbody>
<tr>
<td>$^{17a}$</td>
<td>g 2.0014, 2.0115, 2.0285 (2.0022, 2.0107, 2.0213) A(\text{H}) $^{17b}$</td>
</tr>
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asymmetry in the spin density inferred from the two inequivalent $^{77}$Se hyperfine couplings. The EPR spectra of $[\text{2,6-Mes}_2\text{C}_6\text{H}_3\text{Te}]^+$ ($^{17c}$) established principal values for the rhombic $g$-matrix (Fig. 3 and Table 2). Because of the low natural abundance of $^{125}$Te ($S = 1/2, 7\%$ abundance), the broad EPR spectrum and small background signals of the data only allowed estimates for the hyperfine couplings and were not of sufficient quality to resolve a potential inequivalence in the two Te nuclei. Along $g_1$, the hyperfine coupling is resolved with $|A(^{125}\text{Te})| = 1000$ MHz; along $g_2$ and $g_3$, the $^{125}$Te hyperfine couplings were partially resolved and allowed the estimates shown in Table 2. Davies ENDOR was used to investigate the proton hyperfine interactions and showed that the largest proton hyperfine interaction has further decreased in comparison to $^{17a}$ and $^{17b}$. Along $g_2$, the coupling is $|A| = 3.8$ MHz, whereas for $^{17b}$: $|A| = 6.4$ MHz, and for $^{17a}$: $|A| = 11.0$ MHz. The experimental EPR data are very well modelled by DFT calculations (Fig. 4 and ESi† for details). The principal $g$-values are well reproduced as well as the two inequivalent hyperfine couplings for $^{77}$Se. The DFT data are able to unambiguously assign the largest proton coupling to the para-H atom of the central phenyl group of the one of the $m$-terphenyl substituents (Fig. 4). The trend to smaller $^1\text{H}$ hyperfine values observed experimentally in the series $^{17a}$, $^{17b}$, and $^{17c}$ follows the change in the orientation of two $m$-terphenyl substituents with respect to the central dichalcogen moiety: for $^{17a}$, the phenyl group $\pi$ system of one $m$-terphenyl substituent is well orientated to allow overlap with the $\pi$-type orbitals carrying the unpaired electron on the $S$-$S$ moiety and thus facilitates spin density delocalization. In contrast, the phenyl group of the second $m$-terphenyl substituent is approximately at 90° to this orientation and thus there is poor overlap with the $S$-$S$ orbitals carrying the unpaired electron and less spin delocalisation. For $^{17c}$, the $\pi$-orbitals of both phenyl rings of the $m$-terphenylsubstituent have essentially the same orientation with respect to the $\pi$-type orbitals of the Te–Te moiety carrying the unpaired electron but with relatively poor orbital overlap, resulting in an equivalent but relatively small delocalisation of the spin densities onto the two $m$-terphenyl substituents. As expected, $^{17b}$ has a structure intermediate between $^{17a}$ and $^{17c}$ and this is reflected in the experimental $^1\text{H}$ hyperfine coupling assigned to one para-H of the $m$-terphenyl group. The summed chalcogen-based spin densities from DFT calculations are 0.698, 0.734 and 0.778 for $^{17a}$, $^{17b}$, and $^{17c}$, respectively. These findings in conjunction with the EPR data leave no doubt that all three radical cations are characterized best as chalcogen-centred (i.e. ca. 70–80% spin density on the chalcogen atoms).
The effective magnetic moments ($\mu_{\text{eff}}$) of 17a-c at 300 K are 1.61, 1.48 and 1.41 $\mu_B$, respectively, and reasonably close to the theoretically expected value of 1.744 $\mu_B$, 1.7880 $\mu_B$, and 1.8625 $\mu_B$, respectively, for a system of uncoupled paramagnetic centers with spins $S = 1/2$; the $\mu_{\text{eff}}$ decreases slightly with lowering temperature, that implies weak antiferromagnetic interactions in the solid state (see ESI† for details).

Conclusion

The stable group 16 radical cations $[(RE)_2]^+$ (14a-b, 17a-c; E = S, Se, Te) were prepared and fully characterized by various methods for the first time. Note that these radical cations are isoelectronic with the previously described group 15 radical anions $[(\text{RTE})_2]^-$ (E = P, As, Sb; R = 2,6-[Me$_3$Si$_2$]$_2$CH)$_3$-4-[Me$_3$Si$_2$C]-C$_6$H$_5$). Bearing in mind that the aryltellurenyl fragment RTe is isolobal with atomic iodine I, the paramagnetic $[(\text{RTE})_2]^+$ (17c) and the diamagnetic $[(\text{RTe})_2]^2+$ (10, R = Et) also resemble the paramagnetic $[\text{I}_2]^+$ (18) and the diamagnetic $[\text{I}_2]^+$ (19). In the same notion, the radical cations $[(\text{RS})_2]^+$ (14b, R = CaF$_2$; 17b, R = 2,6-Mes$_2$Ca$_2$H$_5$) and $[(\text{RS})_2]^-$ (14a, R = CaF$_2$; 17a, R = 2,6-Mes$_2$Ca$_2$H$_5$) are closely related to the known $[\text{Br}_2]^+$ (20) and the elusive $[\text{Cl}_2]^+$ (21), which forms $[\text{Cl}_4]^+$ (22) with Cl$_2$ (Scheme 2). The reported findings leave no doubt that the radical cations 14a-b, 17a-c contain chalcogen-centred odd-electron π-bonds. It is noteworthy that recently also odd-electron S–S and Se–Se σ-bonded 1,8-bis(arylcyclohexyl)naphthalene were investigated. In view of the fact that many odd-electron species exist as (reactive) intermediates in various chemical reactions and play an important role in bond formation and cleavage processes, alongside many applications that have been reported or envisaged for stable radicals, the successful isolation of the title compounds provide a suitable entry point for an in-depth exploration of these and related species. Accordingly, the isolation and characterization of other examples for odd-electron bonding, which is of both fundamental and practical interest, is currently under way in our laboratories.

Experimental

Synthesis of (C$_6$F$_5$S)$_2$SbF$_{11}$ ([14a][SbF$_{11}$])

SbF$_5$ (220 mg, 1.0 mmol) was added to solid 13a (40 mg, 0.1 mmol) in a PFA tube. 13a directly changed the colour to dark blue when SbF$_5$ reached the solid. The mixture has been kept 3 h at room temperature until the dissolving of 13a in SbF$_5$ completed. The reaction mixture was then evacuated and SO$_2$ClF (1.0 mL) was condensed into the PFA tube at –196 °C. The mixture was allowed to slowly warm up to –80 °C using a dry ice/ethanol bath. After the melting of the solvent (1 h) completed, the reaction mixture was allowed to slowly warm up to room temperature (4 h) to ensure the complete dissolution of the reaction mixture. Afterwards the PFA tube was sealed under reduced temperature and pressure. Dark blue needles of [14a][SbF$_{11}$] formed at –30 °C after several careful cycles of cooling and warming between –50 °C and to –78 °C.

Synthesis of (C$_6$F$_5$Se)$_2$As$_2$F$_{11}$ ([14b][As$_2$F$_{11}$])

SO$_2$ClF (0.20 mL) was slowly condensed onto solid 13b (25 mg, 0.05 mmol) in a PFA tube at –196 °C. The mixture was slowly warmed up to room temperature for complete dissolution. Then the mixture was cooled again to –196 °C and AsF$_3$ (0.04 mL, 0.5 mmol) was condensed into the PFA tube. The solidified mixture directly changed the colour to dark violet. Afterwards the mixture was slowly warmed up to –78 °C where the mixture melts and partial dissolution of the solid gave a deep green colour. The reaction was allowed to further warm up until reaching –30 °C after 3 h, when additional SO$_2$ClF (1.0 mL) was added. Afterwards the PFA tube was sealed under reduced temperature and pressure. Black needles of [14b][As$_2$F$_{11}$] formed after careful cooling to –78 °C over two days.

Attempts to oxidize (C$_6$F$_5$Te)$_2$ (13c)

At –196 °C, SO$_2$ (0.20 mL) was condensed onto 13c (30 mg, 0.05 mmol) in a PFA tube. After warming up to room temperature, the reaction mixture was cooled down to –196 °C a second time to add AsF$_3$ (0.04 mL, 0.5 mmol). The colour of the reaction changed to dark violet during the process of condensation. Afterwards the mixture was slowly warmed up to –78 °C, to remove the excess of AsF$_3$ under reduced pressure. After a third cooling process (–196 °C), further SO$_2$ (1.0 mL) was added and the reaction mixture was allowed to slowly warm up to –30 °C over a period of 3 h. Afterwards F114 (0.3 mL) has been condensed onto the solidified reaction mixture at –196 °C and the PFA tube sealed afterwards. [Te$_4$][As$_2$F$_{11}$] crystallized as dark violet plates after a careful cooling process from room temperature to –30 °C over two days.

Synthesis of 2,6-Mes$_2$C$_6$H$_5$SH

A solution of 2,6-dimesitylidobenzene (6.50 g, 14.7 mmol) in n-hexane (150 mL) was treated with a 2.5 M solution of n-BuLi (6.60 mL, 14.7 mmol) at room temperature and stirred overnight. The volume of the suspension has been reduced to 20 mL under reduced pressure and the white solid has been filtered off. The product, 2,6-dimesitylphenyl lithium (3.33 g, 10.4 mmol) was dissolved in THF (50 mL), cooled to –78 °C and sulfur powder (0.40 g, 12.5 mmol) was cautiously added. The red suspension was stirred overnight and allowed to warm up to room temperature during that period. Hydrochloric acid (40 mL, 10%) was added to the suspension and the stirring has been extended for another 2 h. The organic phase was extracted with CHCl$_3$ (3 × 50 mL), filtered and the solvent removed under reduced pressure. The crude product was recrystallized from CH$_3$Cl$_2$ to yield pale yellow crystals of 2,6-Mes$_2$C$_6$H$_5$SH (3.02 g, 8.72 mmol, 84%; Mp. 192–194 °C).

$^1$H-NMR (CDCl$_3$): $\delta$ = 7.27 (1H, $J$ = 8.1 Hz), 7.06 (d, 2H, $J$ = 7.5 Hz), 7.02 (s, 4H), 3.07 (s, 1H, SH), 2.38 (s, 6H), 2.07 (s, 12H)
Synthesis of (2,6-Mes2C6H3SeH)2 (16b)

A solution of 2,6-Mes2C6H3SH (2.00 g, 5.77 mmol) in toluene (80 mL) was treated with a solution of ethyl nitrite in ethanol (15%, 25 mL, 40 mmol). The solution was kept at room temperature for 4 h before stirring it at 76 °C over a period of 72 h. The resulting yellow solid was filtered off and dried under vacuum. The crude product was recrystallized from CHCl3 to afford yellow crystals of 16a (1.80 g, 2.60 mmol, 90%; Mp. >230 °C).

1H-NMR (CDCl3): δ = 7.19 (t, 2H, J = 7.4 Hz), 6.88 (d, 4H, J = 7.5 Hz), 6.80 (8H), 2.35 (12H), 1.70 (24H) ppm. 

13C{1H}-NMR (CDCl3) for C48H50Se2Sb (1105.59): C, 56.49; H, 4.94. Found C, 56.57; H, 5.12.

syn-NMR (CDCl3): δ = 0 signal. 77Se-NMR (CDCl3, r.t.): δ = 1362.3 ppm. 125Te-NMR (CDCl3): δ = 1362.4 ppm. ESI MS (CH3CN, positive mode): m/z = 786.3 [C48H50Te2] + for 17b. UV-vis (CH2Cl2, c = 1 × 10−3 mol L−1): λmax = 710 nm. Molar conductivity (CH2Cl2, c = 5 × 10−3 mol L−1): λm = 600 Ω−1 cm2 mol−1. SQUID: μeff (300 K) = 1.48 μB. Anal. calcld for C48H50F6NTe2Sb: C, 56.49; H, 4.94. Found C, 56.57; H, 5.12.

Synthesis of 2,6-Mes2C6H3SeH (2.00 g, 5.77 mmol) was dissolved in THF (50 mL), cooled to −78 °C and selenium powder (0.990 g, 12.3 mmol) was cautiously added. The black suspension was stirred overnight and allowed to warm up to room temperature during that period. Hydrochloric acid (40 mL, 5%) was added to the suspension and the stirring was extended for another 2 h. The organic phase has been extracted with CHCl3 (3 × 50 mL), filtrated and the solvent removed under reduced pressure. The remaining solid was dissolved in THF (50 mL) and slowly added to a suspension of LiAlH4 (0.79 g, 20.8 mmol) in THF (10 mL) at 0 °C. The suspension was stirred for 1 h, cautiously poured onto ice-water and extracted with CHCl3 (3 × 50 mL). The crude product was recrystallized from n-hexane to afford yellow crystals of 2,6-Mes2C6H3SeH (1.76 g, 4.47 mmol, 43%; Mp. recrystallized from n-hexane 16a, 180 °C).

1H-NMR (CDCl3): δ = 7.19 (t, 2H, J = 7.4 Hz), 6.82 (d, 4H, J = 7.5 Hz), 6.74 (8H), 5.72 (12H), 1.72 (24H) ppm. 


Synthesis of (2,6-Mes2C6H3Se)2SbF6 ([17a][SbF6])

Solid 16a (193 mg, 0.28 mmol) was added to a stirred solution of [NO][SbF6] in propionitrile (15 mL). After 18 h of stirring, the solvent was removed under reduced pressure affording a deep blue solid that was recrystallized from CH2Cl2/n-hexane (1:1) at room temperature to yield [17a][SbF6] (250 mg, 0.027 mmol, 96%).

ESI MS (CH3CN, positive mode): m/z = 690.4 [C48H50Sb2] + for 17a. Molar conductivity (CH2CN, c = 5 × 10−7 mol L−1): λ = 1800 Ω−1 cm2 mol−1. SQUID: μeff (300 K) = 1.61 μB. Anal. calcld for C48H50Se2Sb: C, 52.38; H, 4.62; N, 1.24. Found C, 52.38; H, 4.62; N, 1.24.

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Notes and references


2 Stable radicals can be isolated and handled as a pure compound, whereas radicals that are sufficiently long-lived to be observed using conventional spectroscopic methods but cannot be isolated are classified as persistent. D. Griller and K. U. Ingold, Acc. Chem. Res., 1976, 9, 13.


4 R. G. Hicks, Org. Biomol. Chem., 2007, 5, 1321. However, apart from these fundamentally important questions, many applications of stable radicals have been reported or envisaged such as spin labelling, spin trapping, EPR imaging, radical polymerization, or catalysis, just to mention a few. In particular, the development of new materials with technologically relevant properties, such as magnetism or conductivity is in the current focus of interest. A very appealing idea in this context is to use stable radicals as both a charge carrier and magnetic coupler. Selected reviews: (a) Stable Radicals – Fundamentals and Applied Aspects of Odd-Electron Compounds, ed. R. G. Hicks, John Wiley & Sons Ltd, 2010; (b) A. Rajca, Chem. Rev., 1994, 94, 871; (c) J. S. Miller, Adv. Mater., 2002, 14, 1105; (d) Special issue on, Molecular Conductors: Chem. Rev., 2004, 104; (e) J. M. Rawson, A. Alberola and A. Whalley, J. Mater. Chem., 2006, 16, 2560.


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23 The spin density is asymmetric since the largest hyperfine couplings from the second Ar ligand is significantly smaller, Amax < 3.5 MHz.


