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Core and double bond functionalisation of  
cyclopentadithiophene-phosphaalkenes†‡

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The heterofulvenoid cyclopentadithiophene-phosphaalkene is a versatile building block for opto-electronic tuning with donor and acceptor moieties. Both the annulated thienyl rings and the phosphaalkene bond can be functionalised using a variety of chemical transformations, *e.g.* forming C–C, C–E (E=Si, Br) bonds, or oxidation and metal coordination, respectively. Solid-state structures, optical and electronic properties are probed theoretically and experimentally, illustrating the opto-electronic tailoring opportunities at this motif.

## Introduction

Combining electron donor and electron acceptor units to tailor the HOMO and LUMO of dye molecules has been an extensively used concept in the design of dyes for sensors, solar cells, *etc.*<sup>1</sup> In the cyclopentadithiophene (CPDT) motif two thiophene units are linked using a methylene bridge improving the orbital delocalisation by removing a torsional degree of freedom, *i.e.* planarising the conjugated system.<sup>2</sup> Moreover, functionalisation at the bridgehead positions allowed addressing of the solubility (Fig. 1, I) and electronic nature of this annulated bithiophene framework.<sup>2,3</sup> In particular, electron deficient bridgehead substituents, *e.g.* ketone (II) and dicyanoalkenes (III) have led to interesting opto-electronic materials and building blocks for donor–acceptor polymers, in which the exocyclic fulvenoid repeating unit acts as an outstanding acceptor unit.<sup>4,5</sup> Recently, examples of imine functionalised CPDT units have been incorporated into polymers allowing controlled modifications of the HOMO and LUMO levels of the molecular components.<sup>6</sup> The diagonal relationship of phosphorus and carbon has led to a variety of conjugated compounds in which carbene fragments of double bonds are replaced by isolobal phosphinidenes giving rise to

phosphaalkenes.<sup>7–9</sup> The introduction of this group 15 hetero- element as part of conjugated frameworks has noticeable stabilising effects on the LUMO levels, while the HOMO remains almost unaltered. In these building blocks, the low-lying heteroalkene  $\pi^*$  orbitals provide an acceptor group resulting in properties applicable to opto-electronic materials.<sup>10,11</sup> The marked difference to its lighter homologue, *i.e.* the imine derivatives, hold promising developments for the heavier pnictaalkene systems. Interesting approaches have already been pursued to translate the molecular phosphaalkene motifs into polymeric materials,<sup>12</sup> both by keeping the P=C motif intact and using it as polymerizable unit.<sup>13–17</sup> Furthermore, the lone pair at the phosphaalkene can easily be functionalised by metal coordination allowing for catalytic applications,<sup>18–20</sup> but also altering the opto-electronic properties.<sup>21,22</sup> A significantly less explored but equally powerful P=C functionalisation is its oxidation with chalcogens<sup>23–25</sup> and coordination by NHCs.<sup>26</sup>

In view of these properties, we have already described the synthesis and excited state behaviour of phospha- and arsaalkene functionalised cyclopentadithiophene derivatives, where the heteroalkenes lead to pronounced acceptor character by stabilizing the heterofulvenoid-type antibonding orbitals. The elemental substitution (phosphorus *vs.* arsenic) and metal

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† This paper is dedicated to all people whose background, ethnicity, sexuality, or gender identity are underrepresented STEM subjects, and who suffer from the systemic flaws preventing them from flourishing.

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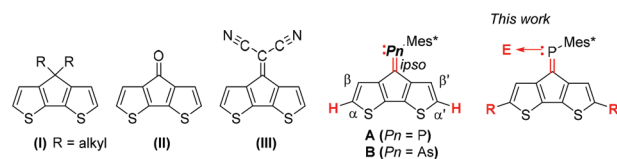


Fig. 1 Literature known CPDT-derivatives (I–III), phospha-/arsaalkene-CPDT-derivative A and B, and novel derivatives studied in this work.



coordination (AuCl) allowed us to address the energy and dynamics of the lowest energy transitions.<sup>27</sup>

In this manuscript, we address the further functionalisation of the CPDT-phosphaalkene by metal coordination or oxidation at the heteroalkene and chemical functionalisation at the thiophene rings achieving stabilisation of the LUMO and destabilisation of the HOMO levels, respectively. The impact of these functionalisations is illustrated by comparing electronic spectra and electro-chemistry data.

## Results and discussion

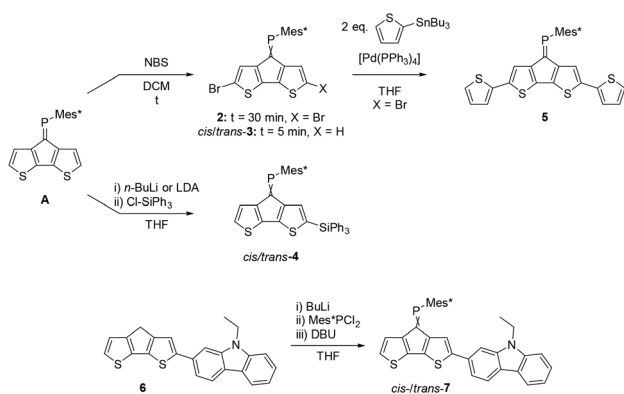
Cyclopentadithiophene phosphaalkene **A** is synthesised according to published procedures from readily available starting materials.<sup>27</sup> In order to study the versatility of this building block we initially investigated the derivatisation of the thiophene rings using standard organic transformations (Scheme 1).

Expectedly, the bromination with *N*-bromo succinimide (NBS) occurs selectively at the  $\alpha,\alpha'$ -positions under ambient conditions and ultrasonication. Chromatographic workup afforded the dibrominated derivative **2** in approx. 75% isolated yields. Attempts to synthesise the corresponding mono-brominated derivatives *cis*-**3** and *trans*-**3** using strictly one equivalent of NBS and low reaction temperatures suffer from unselective reactivities, affording a statistical mixture of **A**, **2** and both isomers of **3**. The two mono-brominated isomers can be separated chromatographically, however isomerisation within a couple of hours establishes the mixture of *cis/trans*-**3** isomers. The conditions which trigger phosphaalkene *E*-/*Z* isomerisation (e.g. light etc.) seem highly dependent on the steric and electronic nature of the substituent and are less explored and understood<sup>28</sup> compared to the diphosphene *E*-/*Z* isomerisation.<sup>29</sup> A possible explanation for the isomerisation of **3** could be an increased phosphonium character promoted by the electron withdrawing Br-substituents.

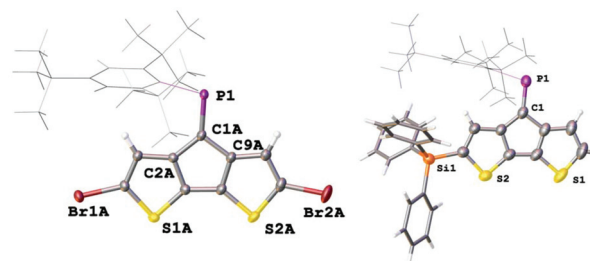
The electron withdrawing effect of the bromide substituent on the remote P=C bond is clearly visible in the <sup>31</sup>P NMR

spectra, which show resonances shifted to higher frequencies. ( $\delta^{31}\text{P}$  for **2**: 269.6 ppm; *cis*-**3**: 262.9 ppm; *trans*-**3**: 263.1 ppm). Compared to **A**, which has a resonance at 256.6 ppm, the addition of each Br-atom shields the resonance by approx. 6 ppm. The assignment of the two isomers is based on the coupling constants of the shielded  $\beta'$  protons positioned below the Mes\*-ring current; for *cis*-**3** this characteristic proton (4.36 ppm) shows as a doublet with a small  $^4J_{\text{PH}}$  coupling of 1.7 Hz. In contrast, *trans*-**3** shows a doublet of doublets (dd) splitting pattern with a  $^3J_{\text{HH}} = 5.1$  Hz and a  $^4J_{\text{PH}} = 1.5$  Hz (4.44 ppm) coupling constants to the neighbouring hydrogen and phosphorus atoms, respectively. Single crystals obtained from slow solvent evaporation of pentane/DCM solutions allowed us to perform single crystal X-ray diffraction experiments (Fig. 2). The solid-state structure of **2** gives a P=C bond length of 1.687(2) Å, which is slightly longer than that of **A**, while the C-P=C angle of 102.5(1)° is almost identical. These metrics suggest that the electronic variation has a larger influence on the bond length than the “hybridisation” and steric hindrance, which is more reflected in the angles around the P-centre. The packing is dominated by weak slipped  $\pi$ - $\pi$  interactions (C-centroid distance 3.711(5) Å) and  $\pi$ ...Br interactions (3.782(1) Å).

We further studied the possibility to deprotonate the acidic  $\alpha$ -positions using a Li-base and reacting the *in situ*-formed anion with an electrophile. Compound **A** was thus reacted with two equivalents of LDA in THF at low temperatures and then allowed to reach r.t., followed by the addition of triphenyl chlorosilane as an electron accepting substituent. Surprisingly, two new resonances in the <sup>31</sup>P-NMR spectra are observed indicating formation of two major products. Despite being separable by column chromatography, slow isomerisation in solution precluded detailed studies of the isomers. Selective crystallisation from the mixture afforded single crystals suitable for X-ray diffraction analysis. Indeed, the solid-state structure of one monosilylated product was confirmed as *cis*-**4**. This isomer (*cis*-**4**) crystallises in the monoclinic space group *P*2<sub>1</sub>/*c* (no. 14). The P1-C1 double bond is 1.678(2) Å, slightly longer



**Scheme 1** Functionalisation of parent CPDT phosphaalkene **A** giving dibrominated (**2** and *cis*-/*trans*-**3**), silylated (**4**), dithienyl (**5**) and carbazole (**7**) derivatives. Full experimental details are provided in the ESI.†



**Fig. 2** ORTEP plot of the solid-state structure of **2** and *cis*-**4**. Selected atoms are drawn at 50% probability levels of thermal ellipsoids, while the Mes\* and phenyl groups are depicted as wireframe and capped sticks. Selected bond lengths [Å] and angles [°]: **2**: P1–C1A 1.687(2), P1–C10A 1.841(2), C1A–P1–C10A 102.54(11). *cis*-**4**: P1–C1 1.678(2), Si1–C9 1.852(2), Si1–C<sub>ipso</sub> 1.862(2), 1.867(2), 1.872(2), C1–P1–C10(Mes\*) 103.1(1). Compound **2** shows a positional disorder (0.92:0.08); distances are given for the molecule with the higher occupancy.



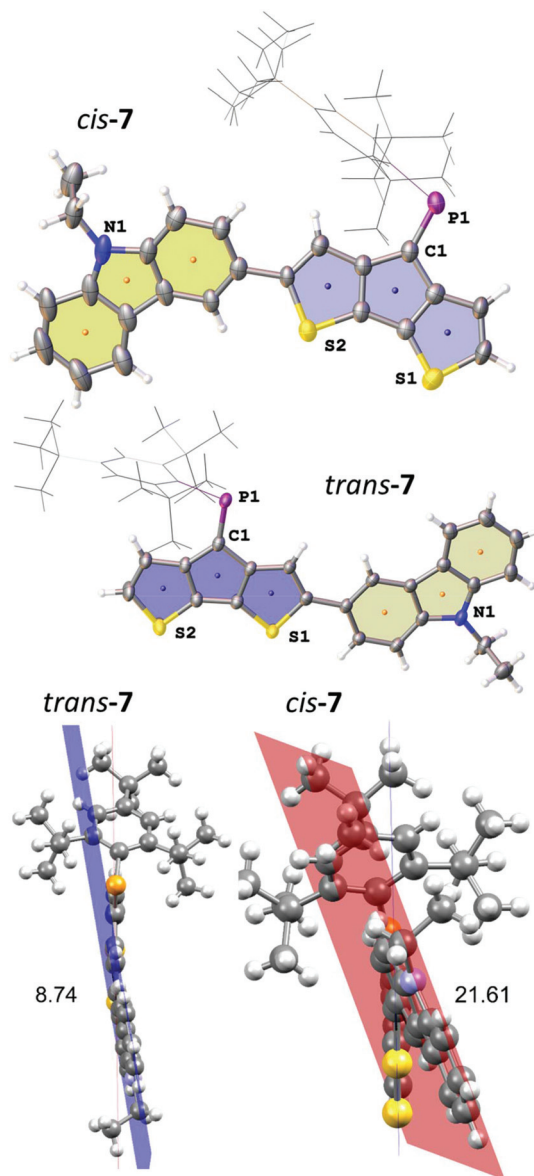
than in the parent system (**A**: 1.667(5) Å). Despite the *cis*-orientation of the sterically demanding triphenylsilyl substituent, the C10–P1–C1 angle increases only marginally from 102.6(2)° in **A** to 103.1(1)° for *cis*-**4**. The Si–C9 bond is significantly shorter (1.852(2) Å) compared to the Si–C<sub>phenyl</sub> bonds (1.862(2) to 1.872(2) Å) indicative of negative hyper-conjugation of the CPDT-core with the silyl substituent. The second product is suspected to be the corresponding *trans*-**4**, based on proton NMR data, indicating that despite the use of two equivalents of LDA or *n*-BuLi the double-deprotonation was unsuccessful.

The usefulness of **2** as a synthon towards extended conjugated systems was probed using a Stille coupling protocol with 2-tributylstannyl thiophene. The reaction proceeds in moderate yield to give the desired disubstituted product **5** with a <sup>31</sup>P-NMR resonance of 265.5 ppm, which is quite similar to those of the brominated derivatives.

Further attempts to use the brominated derivatives **2** and **3** as starting materials in Suzuki type C–C coupling reactions to target extended conjugated systems has proven difficult. In contrast, the carbazole functionalised ketone **5** can be prepared from readily available iodinated precursors and undergoes facile Wolff–Kishner reduction to the methylene derivative **6**. In analogy to the synthesis of **A**, a two-step, one-pot reaction afforded the desired phosphalkene as an inseparable isomeric mixture of *cis*/*trans*-**7** in an approximately 1:1 ratio. Unlike the brominated derivative **3**, no *cis/trans* isomerisation of **7** was observed in solution.

The two isomers crystallised as slightly differently shaped dark purple blocks. Compound *trans*-**7** crystallises in the triclinic space group *P* $\bar{1}$  (no. 2) as a hexane solvate (Fig. 3). The disordered solvent molecule is treated by diffuse contribution to the overall scattering using the solvent masking function implemented in OLEX2.<sup>30</sup> Notably, the P=C double bond in *trans*-**7** is even longer (1.684(2) Å) compared to that of *cis*-**4**, while the C–P=C angle (102.6(1)°) is more similar to that found in **A**. The *cis*-isomer crystallises in the monoclinic space group *C2/c* (no. 15) as a hexane solvate (removed also by solvent masking). The P=C bond lengths in both isomers are comparable (P=C: 1.683(3) Å), but the C–P=C angle is wider for *cis*-**7** (103.4(1)°) reflecting the steric demand of the carbazole substituent on the Mes\* position. The most pronounced effect in the different isomers is observed for the orientation of the carbazole moiety and its torsion with respect to the CPDT core. The smaller torsion angle between the two chromophores' planes in *trans*-**7** (8.74°) gives rise to efficient delocalisation, while steric interaction of the large Mes\* with the carbazole unit leads to an increased torsion angle (21.61°) in *cis*-**7**, with the *N*-ethyl group pointing in the same and opposite direction of the P=C bond, respectively.

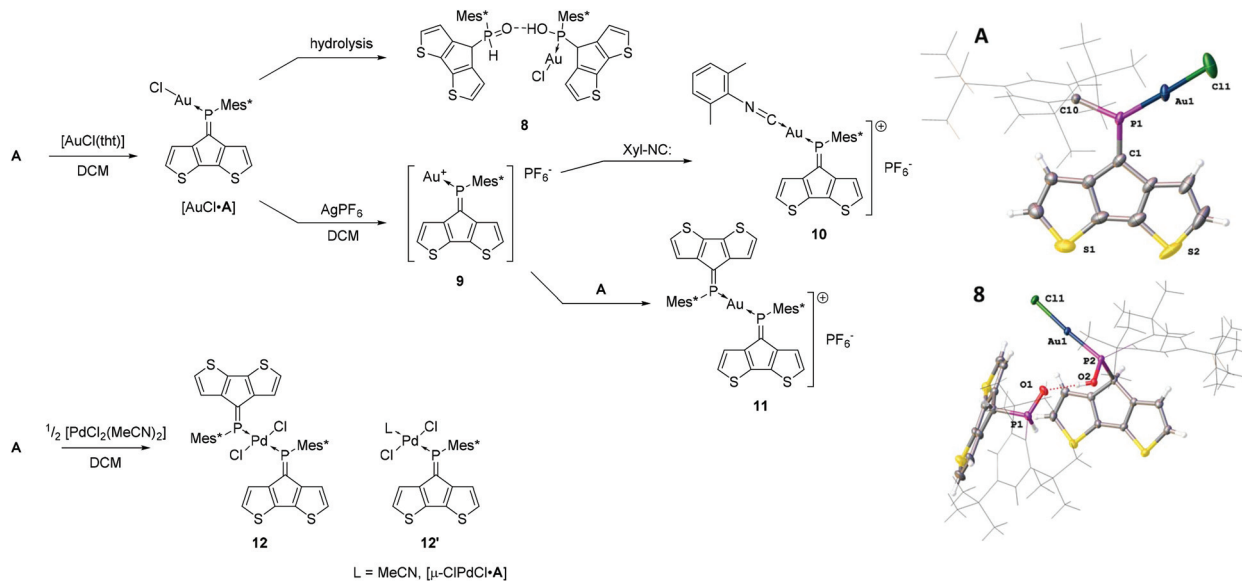
The previously reported gold(i) complex [AuCl·**A**]<sup>27</sup> is only moderately stable under ambient conditions and slowly decomposes in the presence of water (Scheme 2). We report the solid-state structure of [AuCl·**A**] together with one of the decomposition products (**8**) using single crystal X-ray diffraction analysis (Scheme 2). Complex [AuCl·**A**] shows fairly short P=C and P–Au bonds lengths of 1.668(7) and 2.213(2) Å,



**Fig. 3** Top: solid-state structures of *cis*- and *trans*-**7**. Bottom: planes through the CPDT and carbazole rings and the torsion angles between both planes (bottom). Selected bond lengths [Å] and angles [°] *cis*-**7**: P1–C1 1.683(4), P1–C24 1.835(4), C9–C10 1.478(5) C1–P1–C24 103.38 (16). *trans*-**7**: P1–C1 1.683(3), P1–C24 1.845(2), C9–10 1.467(3), C1–P1–C24 102.59(11).

respectively. On the other hand, the C10–P1–C1 angle is fairly large indicating an increase of s-character of the coordinating lone pair. Notably, the almost linear P–AuCl fragment (178.16 (8)°) is strongly twisted out of the CPDT plane (angle between the CPDT least squares plane (l.s.pl.) and the Cl–P vector of 24.15(3)°). In the solid-state structure of decomposition product **8**, the tautomeric hydroxyphosphine and phosphine oxide of AuCl·**A** form a hydrogen bonded quasi-dimer where the two tautomers act as a hydrogen bond donor and a hydrogen bond acceptor, respectively. The colourless crystals solved in the space group *P* $\bar{1}$  (no. 2). The gold coordination is barely





**Scheme 2** Left: metal coordination of the CDPT-phosphaalkene. The previously reported complex  $[\text{AuCl}\cdot\text{A}]$  slowly decomposed hydrolytically with partial metal loss giving the hydrogen bonded monometallic complex **8**. Formation of unstable cationic complex **9** by adding slight excess of  $\text{AgPF}_6$  to a DCM solution of **A**. More stable derivatives are obtained by adding 2,6-dimethylphenyl isocyanide giving **10**, or using two equivalents of **A** giving homoleptic complex **11**. Coordination to palladium proceeds less cleanly giving the mononuclear complex **12** as crystallographically identified product. Reactions in THF and different stoichiometries let us assume that alternative products such as **12'** could be present in solution. Right: solid-state-structures of  $[\text{AuCl}\cdot\text{A}]$  (top) and product of its partial hydrolysis (**8**, bottom) forming the metal coordinated hydroxyphosphine hydrogen bonded to the demetallated phosphine oxide subunit. Selected bond lengths [Å] and angles [°].  $[\text{AuCl}\cdot\text{A}]$ : P1–C1 1.668(7), P1–C10 1.800(6), P1–Au1 2.2130(15), C1–P1–C1 108.3(3), P1–Au1–C1 178.16(8). **8**: P1–O1 1.489(3), P2–O2 1.578(3), P2–Au1 2.232(1), Au1–C11 2.298(1), O1...H–O2 164, O1...O2 2.550(4).

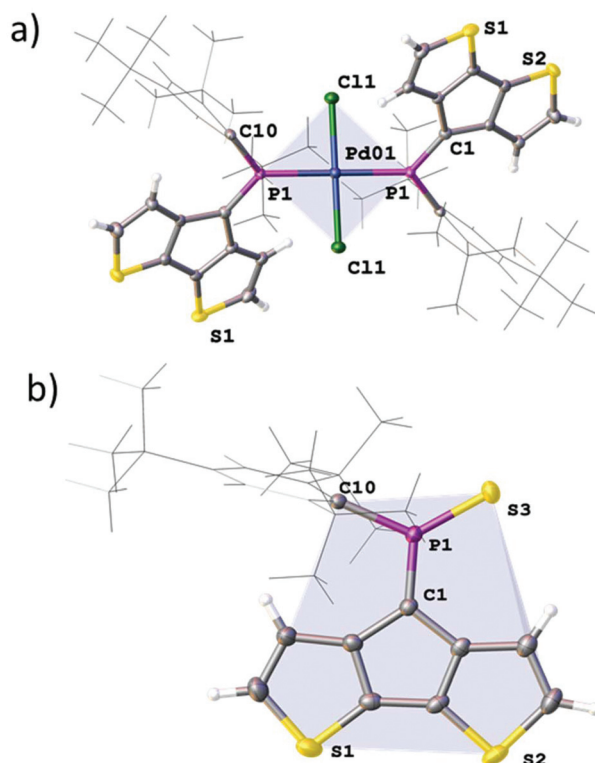
affected by the hydrolysis of the P=C bond with typical P–Au and Au–Cl distances of 2.232(1) and 2.298(1) Å, respectively. The phosphorus oxygen distances are in the typical range of P–O single (1.578(3) Å) and P=O double bonds (1.489(3) Å). The hydrogen bond donor...acceptor distance is 2.550(4) Å and the O–H...O angle 164° indicating strong hydrogen bonding.<sup>31,32</sup> These findings are in line with previous observations of changing the tautomeric forms upon coordination<sup>33–36</sup> or shifting the equilibrium towards the hydroxyphosphine tautomer using electron deficient P-substituents.<sup>37,38</sup>

Complex  $[\text{AuCl}\cdot\text{A}]$  can also be converted into a cationic gold (i) species **9** upon reaction with  $\text{AgPF}_6$  characterised by a broad shielded  $^{31}\text{P}$ -NMR resonance at *ca.* 172 ppm. However, the low coordinate gold(i) species decomposes. Overnight, trace amounts of water and/or reactivity with the  $\text{PF}_6^-$  counter ion give a cationic phosphonium species (based on the deshielded resonances around 320 ppm in the  $^{31}\text{P}$ -NMR spectra), which is followed by further decomposition to untraceable products and formation of a clearly visible gold mirror. Noteworthy, during this process the counter ion decomposes to trifluorophosphine oxide (see ESI†). The cationic intermediate can be stabilised using 2,6-dimethylphenyl isocyanide<sup>39</sup> yielding the heteroleptic complex **10**, which is characterised by a very broad  $^{31}\text{P}$ -NMR resonance at 203.8 ppm (FWHM = 316 Hz) and a sharp septet for the counter ion at –143.9 ppm. Similarly, the reaction of two equivalents of **A** with  $[\text{AuCl}(\text{tht})]$  in the presence of  $\text{AgPF}_6$  gives the moderately stable homoleptic complex **11** with a broad  $^{31}\text{P}$ -NMR resonance at 197.9 ppm (FWHM *ca.*

125 Hz). The  $^{31}\text{P}$ -NMR resonances of the cationic gold complexes **10** and **11** are shifted to higher frequencies compared to the gold(i) chloride complex  $[\text{AuCl}\cdot\text{A}]$  ( $\delta^{31}\text{P} = 175.6$  ppm) indicating reduced electron density. The broadening of the  $^{31}\text{P}$ -NMR resonances can be tentatively attributed to a dynamic behaviour in solution and is commonly observed for phosphalkene coinage metal complexes.<sup>40,41</sup> Interestingly, the deshielded  $\beta$  proton changes from 8.05 ppm in  $[\text{AuCl}\cdot\text{A}]$  to 7.58 and 7.67 ppm in **10** and **11**, respectively, illustrating the different electrostatic situations proximal to the  $\text{Au}(\text{i})^+$  centre, whereas the shielded  $\beta'$  proton is barely influenced. The low-energy absorption band in the UV/vis spectrum of the gold complexes is red-shifted by *ca.* 30 nm upon formation of **9**. This band is detected as a broad feature after coordination of the isocyanide (**10**) and barely visible in the homoleptic complex **11**. Based on previous calculations this band is assigned to a HOMO ( $\pi_{\text{CPDT}}$ ) to LUMO ( $\pi^*_{\text{heterofulvene+metal}}$ ) transition of very low probability.<sup>27</sup>

Addition of 0.5 equivalents of  $[\text{PdCl}_2(\text{MeCN})_2]$  to a dichloromethane solution of **A** immediately results in the formation of a dark green-brown solution. The  $^{31}\text{P}$  NMR spectrum of the reaction mixture contains a major peak at 194 ppm which can be ascribed to  $[\text{PdCl}_2(\text{A})_2]$  (**12**). Additionally, a broad peak can be observed at 162 ppm as well as unreacted **A**. We attribute these observations to the dynamic dissociation of **12** in solution to a monosubstituted palladium complex  $[\text{PdCl}_2(\text{A})\text{L}]$  (**12'**; L = solvent) and free **A**. The species corresponding to the peak at 162 ppm becomes the major product when excessive





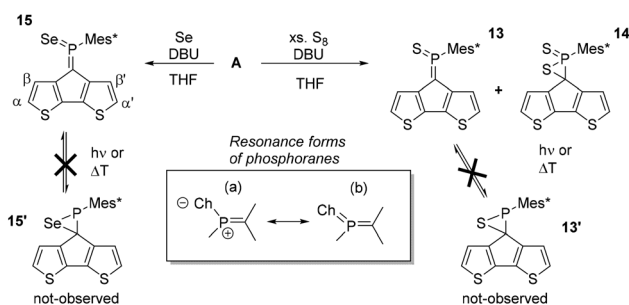
**Fig. 4** ORTEP plot (50% probability ellipsoids) of the solid-state structure of palladium complex **12** (a) and thiaphosphorane **13** (b). (a) Light blue: least squares plane of the square planar Pd-coordination environment including (Pd1, P1, Cl1). Selected bond lengths [Å] and angles [°] **13**: Cl1–Pd01 2.2902(8), P1–Pd01 2.3088(8) P1–Pd01–Cl01 91.64(3), P1–C1 1.674(3), C1–P1–C(Mes\*) 106.8(2). Light blue: least squares plane of the CPDT-PCS subunit. Selected bond lengths [Å] and angles [°] **15**: P1–C1 1.669(4), P1–S3 1.917(1), C1–P1–C10 111.0(2), C1–P1–S3 124.2(1).

amounts of  $[\text{PdCl}_2(\text{MeCN})_2]$  are used, supporting the idea that the **A**:Pd ratio is closer to 1:1 in this species. Formation multi-/di-nuclear complexes with bridging chloride ligands (**12'**;  $L = \mu\text{-ClPdCl}\cdot\text{A}$ ) is also possible (ESI $^\ddagger$ ). The homoleptic complex **12** crystallises as dark green needles from dichloromethane in the centrosymmetric space group  $P\bar{1}$  with half a molecule in the asymmetric unit (Fig. 4). The observed Pd–Cl and Pd–P distances are 2.2902(8) and 2.3088(8) Å, respectively which form a regular square planar palladium environment (P–Pd–Cl 91.64(3)°). The resulting *trans*-complex shows an *anti*-arrangement of the ligand **A** minimising the steric interaction of the bulky Mes\* substituents. Quite interestingly, the P=C bond is only slightly elongated to 1.674(3) Å upon metal coordination, while the C–P=C angle is significantly wider (106.8(2)°), quite similar to that of the gold complex  $[\text{AuCl}\cdot\text{A}]$ .

A significantly less explored route to functionalise the phosphalkene bonds is its oxidation with elemental chalcogens (*e.g.*  $\text{O}_3$ ,  $\text{S}_8$ ,  $\text{Se}_{\text{grey}}$ ), the reactivity of which decreases down the group.<sup>42–44</sup> While methyleneoxophosphoranes are reported to be very unstable, methylenethia- and seleno-phosphoranes have been spectroscopically characterised.<sup>24</sup> The oxidation products of  $\text{Mes}^*\text{P}=\text{C}(\text{TMS})_2$  are all isolable and partially stable

in air.<sup>42</sup> Importantly, the oxidation of the P=C moiety extends the acceptor orbitals including the chalcogen atoms, effectively lowering the LUMO levels (*vide infra*), resulting in a dramatic change of the electronic situation. The reaction of **A** with excess  $\text{S}_8$  in benzene or chloroform did not afford the anticipated methylenethiaphosphorane even after refluxing for 12 hours (Scheme 3). Similarly, **A** does not react with grey selenium in refluxing chloroform or dichloromethane. Interestingly, some reports describe the reaction in neat triethylamine or use small amounts of a non-nucleophilic base to promote the desired oxidation. Addition of diazabicycloundecene (DBU, 2% in the respective solvent), results in immediate consumption of the starting material in cases of sulfur, and initiated the reaction of grey selenium. The latter gives a new species (**15**), with a  $^{31}\text{P}$ -NMR resonance of 144.2 ppm and the typical  $^{77}\text{Se}$  satellites. The coupling constant of 915.7 Hz is indicative of very high *s*-character in the P–Se bond compared to *e.g.* phosphine selenides, which typically have coupling constants of around 750 Hz. Previous work by Appel and co-workers already hinted at this situation with a large  $^1J_{\text{PSe}}$  of 890 Hz. Some examples of selenoiminophosphirenes are currently still the representatives with the largest P–Se coupling constants (960 Hz).<sup>42</sup>

In case of the sulfur reaction, the crude reaction mixture presents three new resonances in the  $^{31}\text{P}$ -NMR spectra at around 160, 80, and 0 ppm. The latter signal is tentatively assigned to the di-oxidised species **14** based on 2D-NMR data (see ESI $^\ddagger$ ) and literature references.<sup>45,46</sup> Chromatographic workup of the crude mixture on silica gel (DCM/pentane) allowed isolation of the methylenethiaphosphorane (**13**) with a characteristic  $^{31}\text{P}$ -NMR resonance at 157.2 ppm. The more deshielded  $^{31}\text{P}$ -NMR signal for **13** compared to **15** agrees with the slightly higher electronegativity of sulfur, favouring an ionic resonance structure (a, Scheme 3). The  $\beta'$  protons are again strongly shielded (**13**: 4.48 and 7: 4.53 ppm) suggesting similar orientation of the Mes\* group. The close proximity to



**Scheme 3** Top: functionalisation of the CPDT phosphalkene using  $\text{S}_8$  and  $\text{Se}_{\text{grey}}$  as oxidants. Thiaphosphorane **13** and selenophosphorane **15** could be isolated using column chromatography as ambiently stable compounds. Rearrangement of the thiaphosphorane into the thiaphosphirane was not observed under various conditions. Bottom: relevant resonance structures of the phosphoranes. The compound with a  $^{31}\text{P}$ -NMR resonance of –1.26 ppm was isolated by column chromatographic work-up. 2D NMR analysis indicates a P–C(CPDT) single bond, and the  $^{31}\text{P}$ -NMR resonance agrees with the formation of **14**.



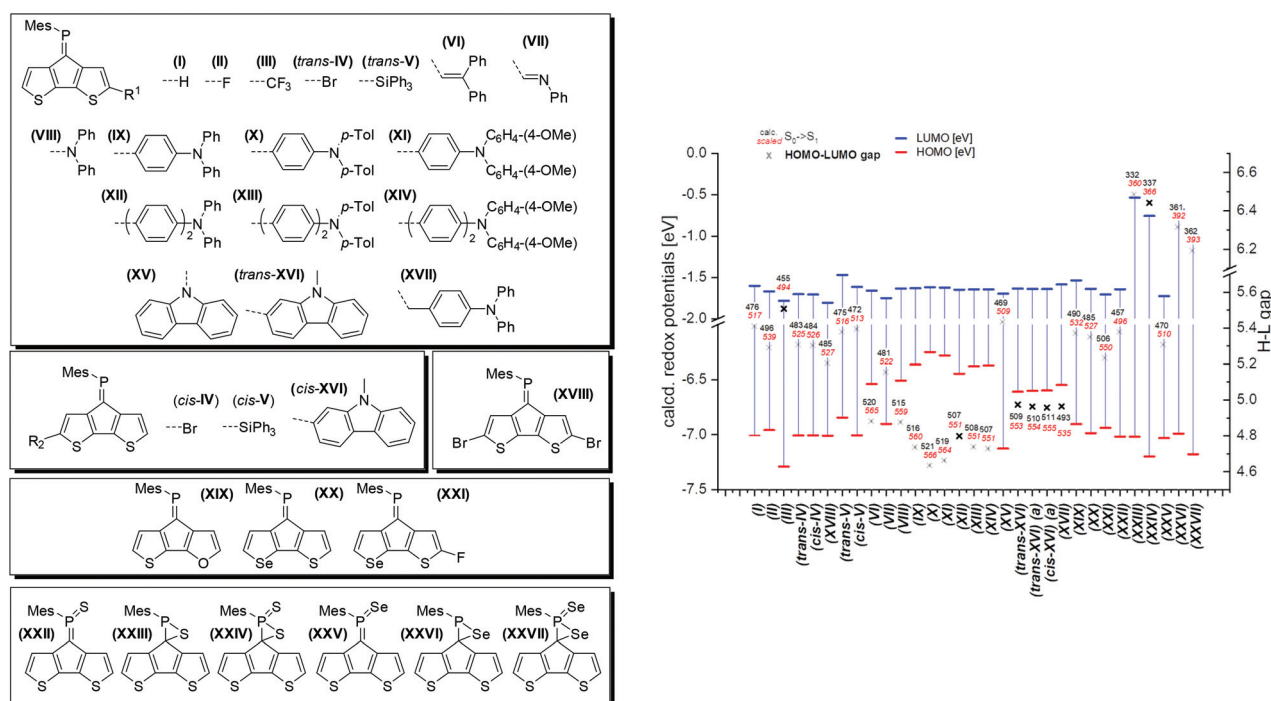
the chalcogen atoms has also a strong impact on the  $\beta$ -proton resonances shifting them to higher frequencies (**13**: 7.88 and **15**: 8.09 ppm). Yoshifuji and co-workers have described the thermal conversion of the methylenethiaphosphorane into the thiaphosphirane, however we were unable to trigger this conversion to give **13'**.<sup>45,46</sup>

The formation of **13** was structurally confirmed by SC-XRD; **13** crystallises in the hexagonal space group  $P6_5$  as dark red blocks. The CPDT core displays the expected co-planarity with the exocyclic unsaturated CPS fragment. The PC (1.669(4) Å) and PS (1.917(1) Å) bonds are in the typical range of strong double bonds. The angle of the P=C unit towards the Mes\* and S are 111.0(2) and 124.2(1); the former being much wider compared to the parent compound as well as the metallated derivatives **12** and [AuCl-A]. The thiaphosphorane unit is coplanar with the CPDT core, efficiently extending the  $\pi$ -conjugated framework. The NBO analysis carried out on these thia- and selenophosphoranes support the strong contribution of the ionic resonance forms in Scheme 3(a) having strong P( $\delta^+$ )Ch( $\delta^-$ ) character and no d-orbital contributions from phosphorus in agreement with the common views of hypervalency in p-block compounds. The significantly larger polarisation in the thiaphosphorane compared to the selenophosphorane is evidenced by the relatively higher negative NBO charge at sulfur (-0.52) compared to Se (-0.45). (*vide infra* and ESI†).

In order to elucidate the impact of the modifications of the parent CPDT phosphalkene **A** we performed a detailed theoret-

tical study including the newly synthesised and further *in silico* derivatives. For all calculated systems the Mes\* substituent has been replaced by the electronically similar, but computationally simpler mesityl (Mes) substituent (Fig. 5). We have focused on the electronic impact of the derivatisations in the ground state and its consequences for the excited states. The structures are optimised in their ground state using a continuum solvation model for DCM at the DFT cam-B3LYP/6-311G\*\* level of theory. At least 30 singlet and triplet transitions are calculated using time-dependent DFT (TD-DFT). As previously noted<sup>27</sup> the charge transfer character complicates computation of the transition energies, thus we scaled the low-energy transitions to the experimentally observed absorption maximum of **A** with respect to the calculated value of (**I**).

Substitution of the core with electron withdrawing halogen substituents (**II**, **III**, *trans*-**IV**, and **XVIII**) has a notable stabilising influence on the LUMO energies compared to the parent model compound (**I**). No significant effects on the HOMO levels have been noted, with the exception of fluoride which also results in the lowest energy absorption. Introducing a silyl group (*cis*- and *trans*-**V**) leads to a destabilisation of both HOMO and LUMO levels, however with no net effect on the absorption energies (or  $E_g$  = HOMO-LUMO gap). The dibromo derivative **2** shows a slightly milder reduction potential compared to the parent compound, which is in line with the calculated LUMO levels of **I** and **XVIII**. Noteworthy, the introduction of these substituents has almost no influence on the extinction coefficients of the low energy transitions. Both, alkene (**VI**) and



**Fig. 5** Left: depiction of all theoretically studied systems using the simplified mesityl (1,3,5-trimethylphenyl) phosphorus protecting group at the TD-DFT B3LYP/6-311G\*\* level of theory using the continuum solvation model for DCM. Right: summarised results of the optical and electronic data of the studied systems. The calculated lowest energy absorption energy/wavelength (grey number) is scaled with respect to the experimental wavelength for **A** (red numbers). The theoretical  $E_g$  (cross) is depicted on the secondary axis (right). HOMO and LUMO energy levels are provided in eV.



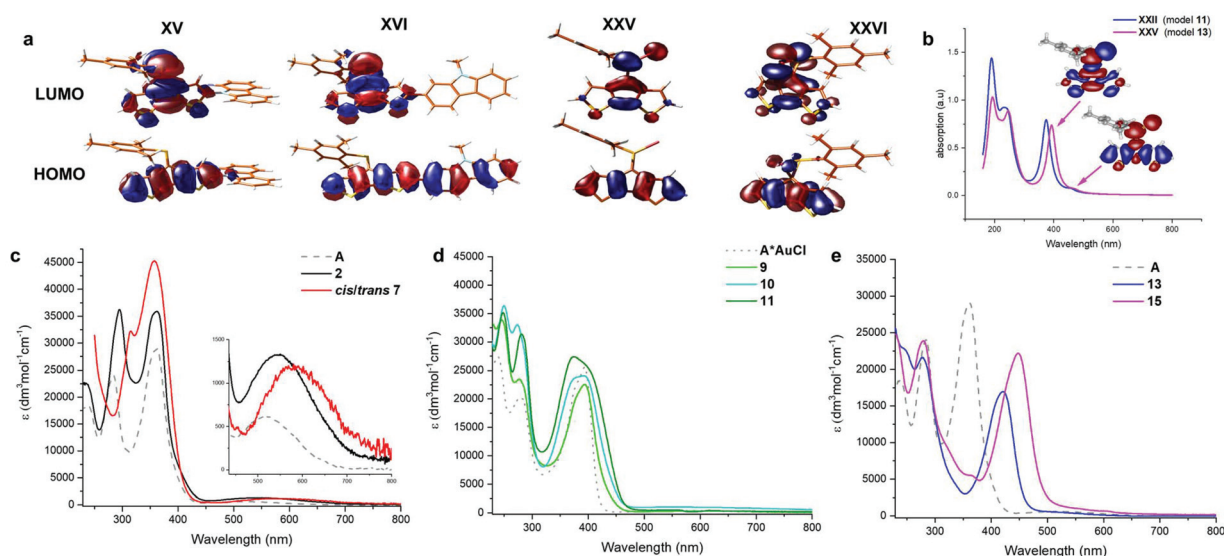
imine (**VII**) derivatives lead to an increased transition probability (by a factor of three to five), however only the imine (**VII**) substituent also causes a pronounced red-shift. Similar effects are obtained using diphenyl (**VIII**), and triphenylamine (**IX–XI**) substitutions. The electron richer ditolyl(phenyl)amine derivative (**X**) is most red-shifted in this series. Separating the donor and acceptor moieties by another phenyl linker has no further effects on the orbital shapes and transition energies.

Interestingly, the N-linked carbazole derivative (**XV**) displays a blue shifted low-probability absorption (0.04), while the 2-linked N-methyl carbazole (**XVI**) shows red-shifted absorption and increased transition probabilities (0.08). This is explained by the different donor sites within the seemingly similar derivatives. The former has a CPDT localised HOMO, while the latter has a significantly extended HOMO stretching over the carbazole and the CPDT core leading to a pronounced charge-transfer character (Fig. 6). In order to further support these findings, electrochemical analyses of isomeric **7** were carried out. The CPDT-phosphaalkene core gives rise to a reversible reduction at  $E_{1/2} = -1.806$  V (vs. Fc/Fc<sup>+</sup>), which is shifted to more negative potentials compared to the parent compound **A** ( $E_{1/2} = -1.62$  V). In contrast to **A**, we observe a reversible oxidation at 0.277 V and a *quasi*-reversible oxidation 0.650 V, followed by an irreversible oxidation around 1.08 V. The former two are assigned to the (*quasi*)-reversible oxidation of the carbazole moiety, while the latter is attributed to the irreversible oxidation of the CPDT-phosphaalkene core, similar to that observed in **A** (+0.84 V). These experimental values are corroborated by the 0.4 eV lower LUMO levels of **XVI** compared to parent phosphaalkene **I**. The different isomers have almost identical optical properties, and their free energies are within less than 0.5 kcal mol<sup>-1</sup> substantiating that different isomers can be present in solution.

Quite surprising is the small impact of replacing the thiophene for selenophene or furan rings, *i.e.* modifying the core of the donor unit, only slightly shifting the absorption energies and not affecting the transition probabilities. A moderate impact could be achieved combining core modification with fluorine-substitution at the  $\alpha$ -position (**XX** vs. **XXI**).

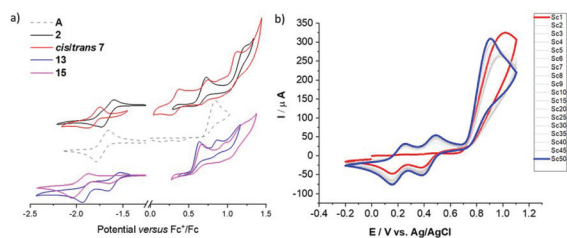
We also studied the chemical functionalisation at the P-centre (and the double bond) using sulfur and selenium oxidations.

The **thiaphosphirane** (**XXIII**) is less stable ( $\Delta G = +1.4$  kcal mol<sup>-1</sup>), than the methylenethiaphosphorane (**XXII**); similarly, the selenophosphirane (**XXV**) is less stable ( $\Delta G = +2.9$  kcal mol<sup>-1</sup>) than the isomeric methyleneselenophosphorane (**XXVI**). The frontier molecular orbitals of the methylenephosphoranes are quite similar to those of unsubstituted **A** (and *e.g.* Au(I) coordinated derivatives) with CPDT localised bonding  $\pi$  orbitals and fulvenoid based anti-bonding orbitals that involve the PS (or Se) moiety. The electronic properties of these two systems (**13** and **15**) have also been probed by means of electrochemical analysis showing clear differences but also a number of similarities. The oxidative side shows two irreversible processes at slightly milder potentials for the sulfur-derivative **13** compared to the selenophosphorane. (**13**: 0.671 and 0.905 V; **15**: 0.707 and 1.012 V). The reductive side is however markedly different. In the case of **13** a mild reversible reduction ( $E_{1/2} = -1.585$  V) is followed by an irreversible reduction (-1.944 V). In contrast, **15** shows an irreversible reduction at -1.658 V, followed by a chemical step (reoxidation at -0.640 V). This initial EC process is followed by a *quasi*-reversible reduction at ( $E_{1/2} = -1.956$  V). The small difference in the oxidative process is in line with the HOMO, being mainly localised on the “organic core”, while the different reductive behaviour agrees with the localisation of the LUMO



**Fig. 6** (a) Calculated Frontier molecular orbitals of selected derivatives. (b) calculated spectra of **XXII** and **XXV** (models of derivatives **13** and **15**), including the EDDMs of the two lowest energy transitions of **XXV**. (c, d & e) optical spectra of compounds **2**, **7**, **9**, **10**, **11**, **13**, **15**, and the literature reported **A**, [**A**\*AuCl]. Solvent: DCM. Due to the facile decomposition of these cationic complexes the extinction coefficients are only approximated.





**Fig. 7** (a) Electrochemical analysis of **2**, **7**, **13**, and **15** in comparison with parent compound **A**. Oxidative and reductive scans at a scan rate of  $100 \text{ mV s}^{-1}$ . Three electrode setup with glassy carbon working electrode (WE), Pt coil counter electrode (CE), Ag wire reference electrode; DCM/ $\text{Bu}_4\text{NPF}_6$  (0.1 M) solution; referenced to the  $\text{Fc}^+/\text{Fc}$  couple. The voltammograms are offset and scaled. (b) Electropolymerisation of **A** (1 mM) by repetitive oxidative cycling ( $-0.2 \text{ V}$  to  $+1.1 \text{ V}$ ) at a scan rate of  $100 \text{ mV s}^{-1}$ . First (red trace) and 50<sup>th</sup> (blue trace) are highlighted indicating the increased current due to material deposition at the WE.

on the heterofulvenoid + chalcogen framework. Model compounds of the radical anions (for S:  $\text{XXII}^{\cdot-}$  and Se:  $\text{XXV}^{\cdot-}$ ) have been investigated showing only small differences in spin and charge localisation. The negative charges on the chalcogen atoms are very similar (S:  $-0.82$ , Se:  $-0.78$ ), however the significantly larger radius of Se implies less nucleophilic character. On the other hand, the spin density at the P atom is higher for  $\text{XXII}^{\cdot-}$  (0.30) compared to  $\text{XXV}^{\cdot-}$  (0.27). In the latter case, spin density is increased at the chalcogen and phosphalkene carbon centres ( $\text{XXV}^{\cdot-}$ : Se 0.14 and C=P 0.31 vs.  $\text{XXII}^{\cdot-}$  S: 0.13 and C=P: 0.28), which might indicate a more facile rearrangement to the three-membered ring structure of the Se-derivative compared to its lighter S-analogue.

The FMOs of the phosphirane have significant bonding and antibonding contributions from the three-membered ring as well as the CPDT core. The calculated  $S_0 \rightarrow S_1$  transitions are slightly blue-shifted for the phosphoranes (**XXII**: 457 (496) nm and **XXV**: 470 (510) nm) compared to the parent motif (**I**: 476 (517) nm), and dramatically blue shifted for the corresponding phosphiranes (**XXIII**: 332(360) nm and **XXVI**: 361(392) nm). These calculations also support the observed inability to trigger conversion of phosphoranes **13** and **15** in the isomeric phosphiranes **13'** and **15'** by illumination or at elevated temperatures.

Re-investigation of the parent phosphalkenes **A**'s electrochemistry revealed that oxidative cycling gives rise to two new reversible redox features. These are accompanied with a gradual shift of the main oxidation peak to milder potential. Based on our previous work we assign this behaviour to an oxidative electro-polymerisation based on the CPDT core (Fig. 7). Detailed investigations of the films for organic electronics applications are currently under way.

## Conclusions

In this manuscript, we have experimentally and theoretically illustrated that the cyclopentadithiophene-phosphalkene core

(**A**) is an interesting building block for optically and electronically tuneable materials. This scaffold offers a large variety of possible functionalisation including the organic core as well as the exocyclic phosphalkene unit. Oxidation of the P-centre with chalcogens and introduction of a donor substituent at the  $\alpha, \alpha'$ -position are most promising. Theoretical studies support these findings and provide further examples of suitable functionalisations. Our current efforts are directed towards using these systems in opto-electronic devices.

## Conflicts of interest

There are no conflicts to declare.

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