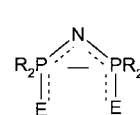
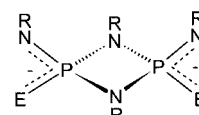
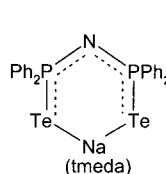
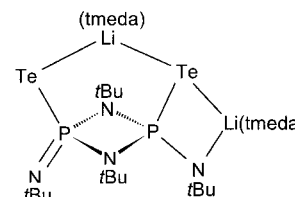


- [12] The structures of racemic **4**, **13**, and **14** have been determined by X-ray crystallography. CCDC-184691 (**4**), CCDC-184693 (**13**), and CCDC-184694 (**14**) contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/conts/retrieving.html (or from the Cambridge Crystallographic Data Centre, 12, Union Road, Cambridge CB21EZ, UK; fax: (+44) 1223-336-033; or deposit@ccdc.cam.ac.uk).
- [13] For examples of other metal-catalyzed asymmetric reactions that show strong solvent effects on enantioselectivity, see: a) Y. Sato, N. Saito, M. Mori, *J. Am. Chem. Soc.* **2000**, *122*, 2371; b) M. Ogasawara, H. Ikeda, T. Nagano, T. Hayashi, *J. Am. Chem. Soc.* **2001**, *123*, 2089.
- [14] CCDC-184692 ((1*R*,2*S*)-**4**) contains the supplementary crystallographic data for this paper.^[12] The single crystal used for the structure determination was obtained from an optically active sample of **4** with 82% *ee*. The small absolute structure parameter of 0.15(12) derived from the structure determination confirms the chirality of the crystal.
- [15] Following the procedures reported by Dauban and Dodd for preparation of iminoiodane $\text{PhI}=\text{NSO}_2(\text{CH}_2)_2\text{SiMe}_3$ from $\text{PhI}(\text{OAc})_2$ and $\text{Me}_3\text{Si}(\text{CH}_2)_2\text{SO}_2\text{NH}_2$ (P. Dauban, R. H. Dodd, *J. Org. Chem.* **1999**, *64*, 5304).
- [16] ^1H NMR (CDCl_3 , 400 MHz) of **19**: δ = 7.89 (d, J = 7.8 Hz, 2H), 7.54 (t, J = 7.4 Hz, 1H), 7.33 (t, J = 7.7 Hz, 2H), 7.16 (m, 4H), 5.19 (m, 1H), 3.06 (m, 4H). The rather high instability of **19** renders it difficult to characterize this compound fully.
- [17] The lower *ee* value in the catalytic reaction than in the reaction between **2** and **19** may arise from a lower loading of **2** in the former case. We once observed that reducing the loading of **2** from 10 to 2 mol% in the amidation of **6** in CH_2Cl_2 , under otherwise the same conditions, led to a decrease in the *ee* value of **11** from 46 to 39%. Notice that some other metal-catalyzed asymmetric reactions also show significant dependence of enantioselectivity on catalyst loading (see for example: H. M. L. Davies, T. Hansen, M. R. Churchill, *J. Am. Chem. Soc.* **2000**, *122*, 3063).

**1** (E = O, S, Se)**2** (E = S, Se)

Despite this intense activity, the tellurium analogues of **1** and **2** are unknown. Anionic tellurophosphinic amides of the type $[\text{tBu}_2\text{P}(\text{Te})\text{NR}]^-$ ($\text{R} = \text{iPr}$, Cy) can be prepared by lithiation of $[\text{tBu}_2\text{P}(\text{Te})\text{NHR}]$ with Li^+Bu^- , and chelate complexes of this anion with Group 12 metals have been investigated as single-source precursors of binary metal tellurides.^[8] Although ditellurides of the type $[\text{R}(\text{Te})\text{P}(\mu\text{-N}t\text{Bu})_2\text{P}(\text{Te})\text{R}]$ ($\text{R} = \text{Me}$, $t\text{Bu}$) have been reported,^[9] our attempts to oxidize $t\text{BuN}(\text{H})\text{P}(\mu\text{-N}t\text{Bu})_2\text{PN}(\text{H})t\text{Bu}$ with an excess of elemental tellurium in boiling toluene produced only the monotelluride $[\text{tBuNH}(\text{Te})\text{P}(\mu\text{-N}t\text{Bu})_2\text{PN}(\text{H})t\text{Bu}]$ in about 5% yield.^[10] Endeavors to generate $(\text{TePPH}_2)_2\text{NH}$ in a similar manner have also been unsuccessful. Consequently, we adopted a different approach to the synthesis of the anionic ligands **1** (E = Te) and **2** (E = Te), which involves metalation of the neutral imido or amido precursor prior to the reaction with tellurium.^[11] Herein, we report the synthesis and X-ray structures of $[[[\text{Na}(\text{tmeda})][(\text{TePPH}_2)_2\text{N}]]_2$ (**3**) and $[\text{Li}(\text{tmeda})_2][\text{Te}(\text{N}t\text{Bu})\text{P}(\mu\text{-N}t\text{Bu})_2\text{P}(\text{N}t\text{Bu})\text{Te}]$ (**4**), (tmeda = tetramethylethylenediamine), which contain the first examples of **1** (E = Te) and **2** (E = Te), respectively.

**3****4**

A New Approach to Metalated Imido and Amido Tellurophosphoranes**

Glen G. Briand, Tristram Chivers,* and Masood Parvez

Monoanionic ligands of the type $[\text{R}_2\text{P}(\text{E})\text{NP}(\text{E})\text{R}_2]^-$ **1** have been investigated extensively as ligands for both main group elements^[1] and transition metals.^[2] This widespread interest stems from their potential uses as lanthanide shift reagents,^[2] industrial catalysts,^[3] luminescent materials,^[4] or in metal extraction processes.^[5] Recently we^[6] and Stahl et al.^[7] reported the first ambidentate dianionic ligands $[\text{RN}(\text{E})\text{P}(\mu\text{-NR})_2\text{P}(\text{E})\text{NR}]^{2-}$ **2**, which adopt a variety of bonding modes, that is *N,E*, *N,N'*, or *E,E'*, with metal centers.

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The reaction of $\text{Na}[\text{Ph}_2\text{PNPPh}_2]$ with tellurium powder in hot toluene in the presence of TMEDA produced **3** as moisture-sensitive, yellow crystals in 33% yield. The molecular structure of **3** (Figure 1) was determined by X-ray diffraction^[12] on crystals obtained from hexane. The ditelluroimido diphosphinate ligand **1** ($\text{R} = \text{Ph}$, $\text{E} = \text{Te}$) is *Te,Te'* chelated to sodium and forms a centrosymmetric dimer through $\text{Na}-\text{Te}$ interactions. This is the first example of *Te,Te'* chelation to an alkali metal. The coordination sphere of the Na^+ ions is completed by one *N,N'* chelating tmeda ligand. A similar structure has been reported for the sodium salt of a monothioimido diphosphinate $[[\text{Na}(\text{thf})_2][(\text{OPPh}_2)(\text{SPPH}_2)\text{-N}]]_2$.^[13] The central Na_2Te_2 ring in **3** is almost square-planar (bond angles at Na1 and Te1 are $87.51(5)^\circ$ and $92.49(5)^\circ$, respectively) with $\text{Na}-\text{Te}$ distances of 3.143(2) and 3.181(2) Å, which are close to the value of 3.16 Å predicted from the ionic radii^[14] and much shorter than the weak $\text{Na}-\text{Te}$ interactions (3.494(3) Å) in the tellurolate $[\text{Na}(\text{tmeda})_2][\text{Te}(2,4,6\text{-Me}_3\text{C}_6\text{H}_2)]$.^[15] The $\text{P}-\text{Te}$ bond lengths of 2.383(1) and 2.403(1) Å are only slightly longer than the values of about 2.37 Å determined for $t\text{Bu}_3\text{P}=\text{Te}$ ^[16] and amino-substituted tellurophosphoranes.^[9c,17] The shorter $\text{P}-$

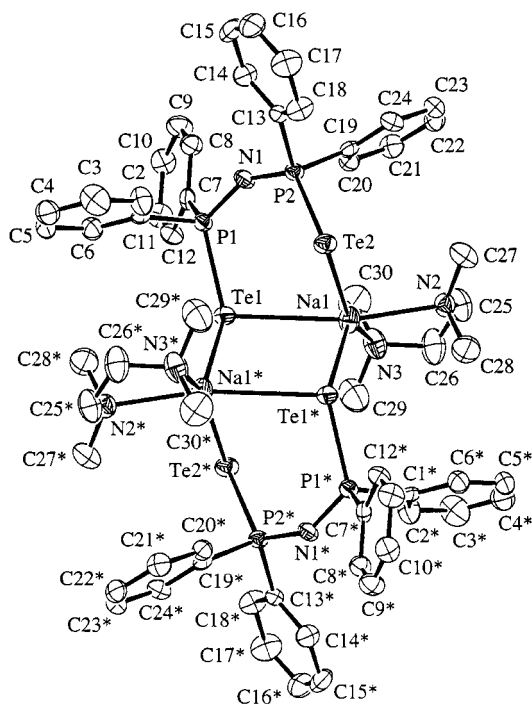


Figure 1. X-ray crystal structure of **3**. Selected bond lengths [Å] and angles [°]: P1–N1 1.591(4), P2–N1 1.589(4), P1–Te1 2.4029(13), P2–Te2 2.3827(13), Na1–Te1 3.143(2), Na1–Te2 3.259(2), Na1–Te1* 3.181(2); P1–N1–P2 143.3(3), N1–P1–Te1 122.18(15), N1–P2–Te2 120.88(15), P1–Te1–Na1 104.20(5), P2–Te2–Na1 93.10(5), Te1–Na1–Te2 84.69(5), Te1–Na1–Te1* 87.51(5), Na1–Te1–Na1* 92.49 (5). Symmetry transformations used to generate equivalent atoms: * $-x, -y, -z + 2$.

Te bond is associated with the two-coordinate tellurium center, which also participates in the longer Te–Na bond (3.259(2) Å) in the six-membered $\text{NP}_2\text{Te}_2\text{Na}$ ring. The ^{31}P NMR and ^{125}Te NMR spectrum of **3** in $[\text{D}_8]\text{THF}$ at 235 K reveal single environments for both the phosphorus and tellurium atoms. This observation may be attributed to a fast exchange process involving the Na and Te sites or to dissociation into a monomer in which the Na^+ ion is solvated by THF. The coupling constant $^1J_{(\text{P-Te})} = 1619 \text{ Hz}$ is, as expected, significantly smaller than the values of 1663–2095 Hz reported for neutral tellurophosphoranes.^[18]

The treatment of $[\text{Li}(\text{thf})]_2[\text{tBuNP}(\mu\text{-NtBu})_2\text{PNtBu}]$ with tellurium powder in toluene at 80 °C in the presence of TMEDA produces **4** as yellow, moisture-sensitive crystals in 19% yield. The molecular structure of **4** (Figure 2) was determined on single crystals obtained from hexane.^[19] The structure reveals a unique bonding mode for ligands of the type **2** in which one Li^+ ion is coordinated in a Te, Te' fashion, while the second Li^+ ion is N, Te chelated by the dianion. By contrast, the sulfur analogue, **4** (E = S), exhibits bis- N, S chelation to two Li^+ centers.^[6b] The X-ray structures of $\text{Li}[\text{tBu}_2\text{P}(\text{Te})\text{NR}]$ (R = *i*Pr, Cy) were not determined, but chelate complexes of the anion with transition metals are N, Te bonded.^[8] The P–Te bond length involving the two-coordinate Te center in **4** is 0.027 Å longer than that involving the three-coordinate Te atom, presumably because the former is associated with the shorter exocyclic P–N bond (1.534(2) versus 1.574(2) Å). The mean value of $d(\text{P}–\text{Te})$ in **4** is, as expected, significantly longer by about 0.07 Å than the value

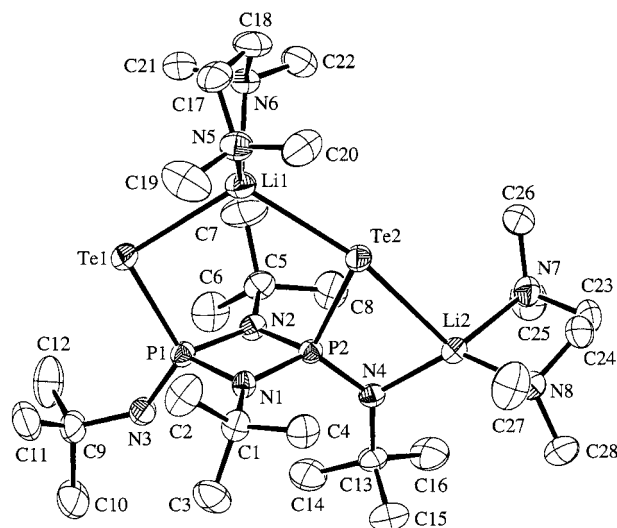


Figure 2. X-ray crystal structure of **4**. Selected bond lengths [Å] and angles [°]: Li1–Te1 2.842(5), Li1–Te2 2.791(5), Li2–Te2 2.865(5), Li2–N4 1.973(5), P1–Te1 2.4515(8), P2–Te2 2.4243(7), P1–N3 1.534(2), P2–N4 1.574(2); Te1–Li1–Te2 113.35(17), N4–Li2–Te2 80.93 (18), N3–P1–Te1 119.94(11), N4–P2–Te2 104.89(9), P2–Te2–Li2 65.59 (10), P2–Te2–Li1 96.51(10), Li1–Te2–Li2 160.06(14), P1–Te1–Li1 97.15(11).

of 2.370(1) Å found for the monotelluride $[\text{tBuNH}(\text{Te})\text{P}(\mu\text{-NtBu})_2\text{PN}(\text{H})\text{tBu}]$.^[10] The NMR data for **4** in $[\text{D}_8]\text{toluene}$ demonstrate that the novel, asymmetric coordination of the two Li^+ ions to the dianion **2** observed in the solid state is maintained in solution at 235 K. Thus, the ^7Li NMR spectrum of **4** in $[\text{D}_8]\text{toluene}$ exhibits two well-separated resonance signals at $\delta = 0.70$ and 3.88 ppm, while the ^{31}P NMR spectrum shows two singlets (with ^{125}Te satellites) at $\delta = -113.7$ and -75.3 ppm . Consistently, the ^{125}Te NMR spectrum displays two doublets at $\delta = -289$ and -87 ppm . The smaller values of $^1J_{(\text{P-Te})}$, 1309 and 1467 Hz, are consistent with the longer P–Te bond lengths in **4** compared to those in **3**.

In summary, the discovery of a new approach to the previously inaccessible ditelluroimidodiphosphinate anion **1** (E = Te) and the ditellurodiimidocyclodiphosph(v)azane dianion **2** (E = Te) paves the way for extensive investigations of the coordination chemistry of these tellurium-centered ligands. Semiempirical PM3 calculations indicate that the success of this method can be attributed to an increase in the nucleophilicity of the phosphorus(III) centers in $(\text{PPh}_2)_2\text{NH}$ and $\text{tBuN}(\text{H})\text{P}(\mu\text{-NtBu})_2\text{PN}(\text{H})\text{tBu}$ upon metalation. Details of this study, as well as additional examples of the successful application of this approach to the synthesis of novel $\text{N}(\text{v})\text{Te}$ anions, will be discussed in a full account of this work.

Experimental Section

All manipulations were carried out under anaerobic and anhydrous conditions. NMR spectra were obtained on $[\text{D}_8]\text{THF}$ (**3**) or $[\text{D}_8]\text{toluene}$ (**4**) solutions at 235 K using a Bruker DRX 400 MHz spectrometer.

3·0.25 C_6H_{14} : A mixture of $\text{Na}[\text{Ph}_2\text{PNPPH}_2]$ ^[20] (0.100 g, 0.245 mmol), tellurium powder (0.063 g, 0.49 mmol), and TMEDA (0.114 g, 0.982 mmol) in toluene (5 mL) was heated to 80 °C for 3 h. After cooling to 23 °C, the mixture was centrifuged and decanted to remove unreacted tellurium. Yellow crystals of **3**·(C_6H_{14})_{0.25} (0.065 g, 33%) were obtained after one day at 23 °C. Elemental analysis calcd for $\text{C}_{31.5}\text{H}_{39.5}\text{N}_3\text{NaP}_2\text{Te}_2$: C 47.27, H 4.97, N

5.25; found: C 48.42, H 5.35, N 5.32; $^{31}\text{P}\{^1\text{H}\}$ NMR: $\delta = 9.5$ ppm (s, $^1J_{\text{P(Te)}} = 1632$ Hz); ^{125}Te NMR: $\delta = -403.6$ ppm (d, $^1J_{\text{P(Te)}} = 1619$ Hz).

4: A mixture of $[\text{Li}(\text{thf})_2][\text{tBuNP}(\mu\text{-NbBu})_2\text{NbBu}]^{[21]}$ (0.200 g, 0.396 mmol), tellurium powder (0.101 g, 0.79 mmol), and TMEDA (0.368 g, 3.17 mmol) in toluene (5 mL) was heated at 80°C for 3 h. The mixture was centrifuged and the supernatant was decanted from unreacted tellurium. After removal of solvent under vacuum, the product was redissolved in *n*-hexane (ca. 1 mL). Yellow crystals of **4** (0.062 g, 19%) were deposited after one day at 23°C . Elemental analysis calcd for $\text{C}_{28}\text{H}_{68}\text{Li}_2\text{N}_8\text{P}_2\text{Te}_2$: C 39.66, H 8.08, N 13.22; found: C 38.06, H 8.23, N 12.25; ^1H NMR ($[\text{D}_8]\text{toluene}$, 235 K): $\delta = 1.62$ (s, 9H; NbBu), 1.97 (s, 9H; NbBu), 2.18 (s, 18H; $\mu\text{-NbBu}$), 2.20 (s, 12H; NMe_2), 2.30 ppm (s(br), 8H; NCH_3); $^{31}\text{P}\{^1\text{H}\}$ NMR: $\delta = -113.7$ (s, $^1J_{\text{P(Te)}} = 1467$ Hz), -75.3 ppm (s, $^1J_{\text{P(Te)}} = 1309$ Hz); ^7Li NMR: $\delta = 0.70$ (s), 3.88 ppm (s); ^{125}Te NMR: $\delta = -289$ (d, $^1J_{\text{Te,P}} = 1352$ Hz), -87 ppm (d, $^1J_{\text{Te,P}} = 1486$ Hz). The values of $^1J_{\text{Te,P}}$ obtained from the ^{31}P NMR spectrum are more reliable than those obtained from the ^{125}Te NMR spectrum owing to the broad line widths ($\Delta\nu_{1/2} \sim 225$ Hz) in the latter spectrum.

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- [19] Crystal data for **4**: $\text{C}_{28}\text{H}_{68}\text{Li}_2\text{N}_8\text{P}_2\text{Te}_2$, $M_r = 847.92$, monoclinic, space group $P2_1/c$, $a = 17.8810(3)$, $b = 9.7701(2)$, $c = 24.0003(5)$ Å, $\beta = 91.6451(6)^\circ$, $V = 4191.10(14)$ Å³, $Z = 4$, $\rho_{\text{calcd}} = 1.344 \text{ g cm}^{-3}$, $F(000) = 1728$, $T = 170(2)$ K. Data were collected on a Nonius Kappa CCD diffractometer on a yellow plate ($0.20 \times 0.15 \times 0.08 \text{ mm}^3$) coated with Paratone 8277 oil and mounted on a glass fiber using ω and ϕ scans. Of the 16 883 reflections collected 9529 were unique ($R_{\text{int}} = 0.028$) and 7266 were observed [$I > 2.00\sigma(I)$] and used to refine 379 parameters. Structure solution and refinement followed the procedures described above for **3**. Refinement by least-squares calculations converged at $R_1 = 0.034$ and $wR_2 = 0.072$. CCDC-184064 and CCDC-184065 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/contents/retrieving.html (or from the Cambridge Crystallographic Data Centre, 12, Union Road, Cambridge CB2 1EZ, UK; fax: (+44) 1223-336-033; or deposit@ccdc.cam.ac.uk).
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Metallabenzenes and Valence Isomers. Synthesis and Characterization of a Platinabenzene**

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Michael M. Haley*

*Dedicated to Professor Gottfried Huttner
on the occasion of his 65th birthday*

In our group we have developed a convenient method for the preparation of iridabenzene starting from the C_5 synthetic equivalent *Z*-3-(2-iodovinyl)-1,2-diphenylcyclopropene (**1**).^[1] Using this precursor, iridabenzene synthesis can also be carried out in a stepwise manner via an iridabenzvalene intermediate, which could be characterized and converted into the corresponding iridabenzene by thermal treatment.^[2] In addition to varying the coordination pattern at the iridium center and the substituents on the vinylcyclopropene precursor,^[3] we are currently trying to show this concept to be generally applicable for the synthesis of metallabenzene and

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