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Controlled scrambling reactions to polyphosphanes *via* bond metathesis reactions†

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Triphosphanes $R'_2PP(R)PR'_2$ (**9a,c**: $R = \text{Py}$; **9b** $R = \text{BTz}$), 1,3-diphenyl-2-pyridyl-triphospholane **9d** and pentaphospholanes $(\text{RP})_5$ (**13**: $R = \text{Py}$; **18**: $R = \text{BTz}$) are obtained in high yield of up to 98% from the reaction of dipyrazolylphosphanes RPpyr_2 (**5**: $R = \text{Py}$; **6**: $R = \text{BTz}$; $\text{pyr} = 1,3\text{-dimethylpyrazolyl}$) and the respective secondary phosphane ($R'_2\text{PH}$, $R' = \text{Cy}$ (**9a,b**), $t\text{Bu}$ (**9c**); $\text{PhPH}(\text{CH}_2)_2\text{PHPh}$ (**9d**)). The formation of derivatives **9a–d** proceeds *via* a condensation reaction while the formation of **13** and **18** can only be explained by a selective scrambling reaction. We realized that the reaction outcome is strongly solvent dependent as outlined by the controlled scrambling reaction pathway towards pentaphospholane **13**. In our further investigations to apply these compounds as ligands we first confined ourselves to the coordination chemistry of triphosphane **9a** with respect to coinage metal salts and discussed the observation of different *syn*- and *anti*-isomeric metal complexes based on NMR and X-ray analyses as well as quantum chemical calculations. Methylation reactions of **9a** with MeOTf yield triphosphan-1-ium $\text{Cy}_2\text{MePP}(\text{Py})\text{PCy}_2^+$ (**10**⁺) and triphosphane-1,3-dium $\text{Cy}_2\text{MePP}(\text{Py})\text{PMeCy}_2^{2+}$ (**11**²⁺) cations as triflate salts. Salt **11**[OTf]₂ reacts with pentaphospholane **13** in an unprecedented chain growth reaction to give the tetraphosphane-1,4-dium triflate salt $\text{Cy}_2\text{MePP}(\text{Py})\text{P}(\text{Py})\text{PMeCy}_2^{2+}$ (**19**[OTf]₂) *via* a P–P–P bond metathesis reaction. The latter salt is unstable in solution and rearranges *via* a rare [1,2]-migration of the Cy_2MeP -group followed by the elimination of the triphosph-2-en-1-ium cation [$\text{Cy}_2\text{MePPPMeCy}_2^+$ (**20**⁺) to yield a novel 1,4,2-diazaphospholium salt (**21**[OTf]).

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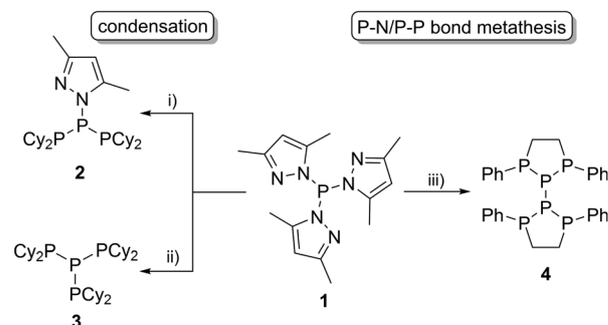
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Introduction

A wide range of polyphosphorus compounds, comprising diverse bonding motifs, have been synthesised in the last few decades.^{1,2} Many of the observed compounds are often found to be labile in solution and their rearrangement or decomposition reactions are discussed to proceed *via* intermolecular scrambling reactions.³ Even though there are early reports on controlling such scrambling reactions,⁴ the selective and high yielding synthesis of certain polyphosphanes is still difficult, hampering their subsequent chemistry. Besides classical approaches for the generation of P–P bonds such as salt metathesis reactions,^{2a,b} dehalosilylation reactions^{2c} or condensation reactions,^{2d} our group focusses on the development of methodologies using tripyrazolylphosphane **1** or derivatives thereof (*vide infra*) as a P₁-building unit for P–P bond forming reactions *via* condensation or P–N/P–P bond

metathesis reactions (Scheme 1).^{5,6} In this regard, the reaction of **1** with Cy_2PH to give triphosphane **2** or *iso*-tetraphosphane **3** depending on the applied stoichiometry has been reported.⁵ The reaction of **1** with 1,2-bis(phenylphosphanyl)ethane yields hexaphosphane **4** in a sequence of condensation and P–N/P–P bond metathesis reactions, showing an example of a controlled scrambling reaction.⁶ In our effort to further control these metathesis reactions and driven by our general interest in azole-



Scheme 1 Application of tripyrazolylphosphane **1** in P–P bond formation reactions; (i) 2 Cy_2PH , –2 pyrH , CH_3CN ; (ii) 3 Cy_2PH , –3 pyrH , and CH_3CN ; (iii) 5 $\text{PhPH}(\text{CH}_2)_2\text{PHPh}$, 4 Ppyr_3 , – $\text{PhP}(\text{pyr})(\text{CH}_2)_2\text{P}(\text{pyr})\text{Ph}$, –10 pyrH , CH_3CN .

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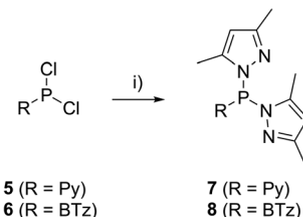
† Electronic supplementary information (ESI) available. CCDC 1950469–1950486. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c9sc04501e



and azine-substituted phosphanes, we explored the synthesis of phosphanes with a pyridyl- or benzothiazolyl-substituent bound to phosphorus *via* the carbon atom. This limits the number of P–N bound pyrazolyl-substituents and therefore the possible reaction pathways by P–N/P–P bond metathesis. For a better overview and in thought of a systematic approach we organised our general types of phosphanes with respect to their number of P–N bound substituents as type I to type IV (Scheme 2). In type I phosphanes all three substituents are bound *via* P–N bonds whereas a type II phosphane is defined by two P–N bonds and one P–C bond. This order is maintained in type III phosphanes with one P–N and two P–C bonds and is finalised in type IV phosphanes featuring only P–C bonds. Importantly, all substituents at the phosphorus atom have an additional sp² hybridised donor atom such as nitrogen, being essential as a further reaction or coordination site and for the possible formation of resonance stabilised reaction intermediates. In this contribution, we report the synthesis of the hitherto unknown type II phosphanes carrying either a pyridyl- or benzothiazolyl-substituent and their use for the generation of triphosphanes, a triphospholane and pentaphospholanes and selected one triphosphane to explore its coordination chemistry with respect to coinage metal salts.

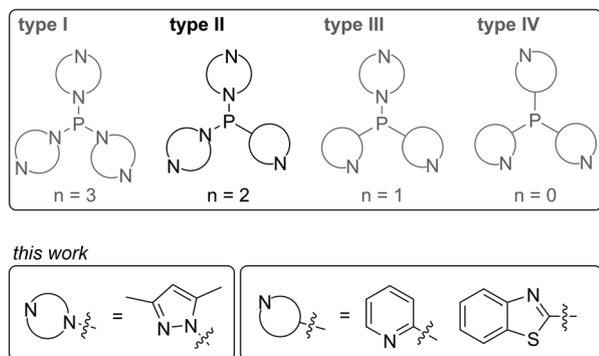
Results and discussion

The targeted type II phosphanes are synthesised by the reaction of the dichlorophosphanes R₂PCl₂ (R = Py (5); R = BTz (6))^{7,8} with two equivalents of 3,5-dimethyl-1-(trimethylsilyl)pyrazole (Scheme 3).⁹ The released chloro(trimethyl)silane can be removed under reduced pressure after 16 hours and no further purification is necessary. Both phosphanes 7 and 8 are conveniently prepared on a multi gram scale in excellent yields (97% (7); 96% (8)). The ³¹P NMR spectra of the dipyrazolylphosphanes show a singlet resonance in each case (7: δ(P) = 44.8 ppm; 8: δ(P) = 38.0 ppm) being significantly shifted to a higher field compared to the corresponding dichlorophosphanes (5: δ(P) = 138.7 ppm; 6: δ(P) = 132.0 ppm). Suitable crystals for X-ray analysis of phosphanes 7 and 8 are obtained by slow vapour diffusion of *n*-pentane into a saturated CH₂Cl₂ solution of 7 and



Scheme 3 Synthesis of dipyrazolylphosphanes 7 and 8; R = Py; (i) 2 eq. pyrSiMe₃, neat, –2 Me₃SiCl; R = BTz; (i) 2 eq. pyrSiMe₃, Et₂O, –2 Me₃SiCl.

8. Their molecular structures are depicted in Fig. 1. The P–C bond lengths (7: P1–C1 1.827(1) Å; 8: P1–C1 1.828(2) Å) are in the expected range of a P–C single bond (1.83 Å),¹⁰ involving a tri-coordinate phosphorus atom. For both phosphanes the P–N bond lengths (7: P1–N4 1.721(1) Å, P1–N2 1.724(1) Å; 8 P1–N2/N2' 1.719(1) Å) are in accordance with those reported for tri(1*H*-pyrazol-1-yl)phosphane (1.714(4) Å).¹¹ In continuation, we carried out the reaction of dipyrazolylphosphanes 7 and 8 with secondary phosphanes R'₂PH (R' = Cy, *t*Bu or R'₂PH = PhP(H)C₂H₄P(H)Ph) to give triphosphanes 9a–c and triphospholane 9d with concomitant release of two equivalents of 3,5-dimethylpyrazole (Scheme 4). Compounds 9a–d readily precipitate from the reaction mixture and are isolated by filtration. Subsequent washing with CH₃CN and drying *in vacuo* give 9a–d as colourless solids in good to excellent yield (96% (9a); 93% (9b); 77% (9c); 69% (9d)). The ³¹P NMR spectra of all compounds show AX₂ spin systems which are characteristic of triphosphanes and 1,2,3-triphospholanes (Fig. 2). The NMR spectroscopic parameters are listed in Table 1. Single crystals for X-ray analysis are obtained by slow vapour diffusion of *n*-pentane into saturated CH₂Cl₂ solutions of 9a–d. The molecular structures of 9a–d are depicted in Fig. 3. Their structural parameters compare very well with those of similar triphosphanes¹² and are thus not discussed in detail. Only the bond angle of the P–P–P moiety in compound 9c is approximately 10° wider which can be explained by the sterically demanding *tert*-butyl substituents. In recent years numerous examples of the coordination chemistry of anionic¹³ and zwitterionic¹⁴ oligophosphorus compounds towards transition metals have been



Scheme 2 Classification of *N*-heterocyclic substituted phosphanes into type I to type IV phosphanes (*n* is the number of P–N bonds) with an emphasis on the type II phosphanes presented in this contribution.

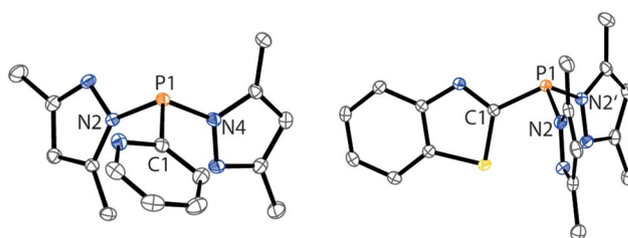
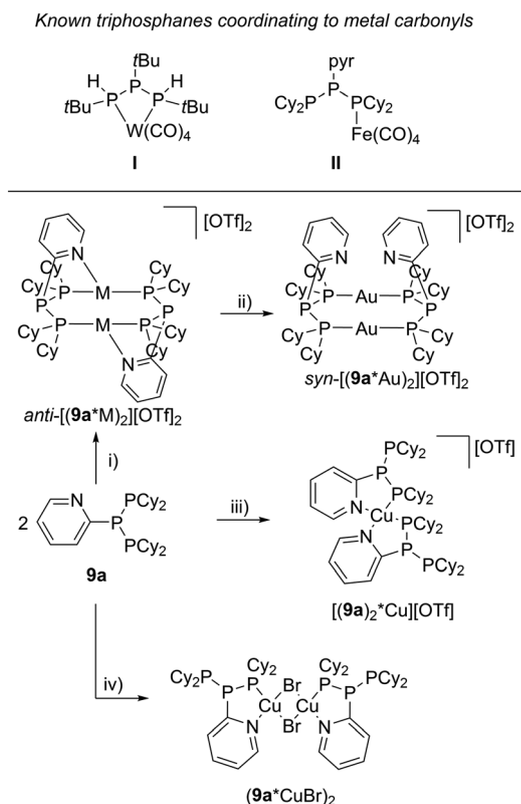


Fig. 1 Molecular structures of 7 (left) and 8 (right) (hydrogen atoms are omitted for clarity and thermal ellipsoids are displayed at 50% probability). Selected bond lengths (Å) and angles (°): (7) P1–C1 1.827(1), P1–N4 1.721(1), P1–N2 1.724(1), C1–P1–N4 101.28(5), N4–P1–N2 103.26(5), N2–P1–C1 100.18(5); (8) P1–C1 1.828(2), P1–N2 1.719(1), C1–P1–N2 101.26(5), N2–P1–N2' 103.84(7).





Scheme 5 Known transition metal complexes featuring a neutral triphosphane ligand (top); synthesis of coinage metal coordination complexes starting from triphosphane **9a** (bottom); (i) 2 eq. Ag[OTf], PhF (M = Ag); 2 eq. [(MeCN)₄Cu][OTf], CH₂Cl₂, -8 MeCN (M = Cu); (ii) 2 eq. (tht)AuCl, MeCN, -2 tht, -2 AgCl; (iii) 1 eq. [(MeCN)₄Cu][OTf], CH₂Cl₂, -4 MeCN; (iv) 2 eq. (tht)CuBr, THF, -2 tht.

been investigated by variable temperature NMR experiments and theoretical calculations (*vide infra*). The difference in the arrangement in the solid state causes a widening of the P–M–P angles in the copper complex [(**9a***Cu)₂][OTf]₂ (P3′–Cu1–P1 136.52(2)°) compared to the silver complex [(**9a***Ag)₂][OTf]₂ (P1–Ag1–P3′ 143.96(3)°) to an almost linear geometry around the gold atoms in [(**9a***Au)₂][OTf]₂ (P1–Au1–P3′ 171.52(3)°). The distances between the silver atoms in [(**9a***Ag)₂][OTf]₂ and the gold atoms in [(**9a***Au)₂][OTf]₂ indicate the presence of argentophilic and aurophilic interactions, respectively.¹⁸ As the distance between the copper atoms in [(**9a***Cu)₂][OTf]₂ is slightly larger than the sum of the van der Waals radii of two copper atoms (2.80 Å),¹⁹ significant van der Waals interactions can be excluded. The observed P–M bond lengths in [(**9a***M)₂][OTf]₂ as well as in [(**9a**)₂*Cu][OTf] and (**9a***CuBr)₂ are in good agreement with bond lengths reported for the structurally similar [M₂(dcpm)₂]²⁺ cations (M = Cu(I), Ag(I) or Au(I); dcpm = bis(dicyclohexylphosphanyl)methane).²⁰ For (**9a***CuBr)₂ the formation of the typical Cu₂X₂ (X = halogen) geometry is found, only marginally differing from the typical planar arrangement reported for many Cu₂X₂-type structures.²¹ Compared to free ligand **9a**, the P–P bond lengths and the P–P–P angles in the coinage metal complexes are only marginally altered. The room temperature NMR spectra of the dinuclear complexes [(**9a***Au)₂][OTf]₂ and [(**9a***Ag)₂][OTf]₂ show broadened resonances due to dynamic processes in solution.²² The low temperature ³¹P NMR spectrum of [(**9a***Au)₂][OTf]₂ shows two separate AA′XX′X″X″′ spin systems at δ(P_A) = −34.2 ppm, δ(P_X) = 52.6 ppm and δ(P_A) = −37.8 ppm, δ(P_X) = 50.3 ppm corresponding to the respective *syn*- and *anti*-isomers (Fig. 5). Details of the coupling constants are included in the ESI.† For [(**9a***Ag)₂][OTf]₂ two isomers with two separate AA′XX′X″X″′ spin systems at δ(P_A) = −48.5 ppm, δ(P_X) = 36.1 ppm and δ(P_A) = −51.5 ppm, δ(P_X) = 36.1 ppm are

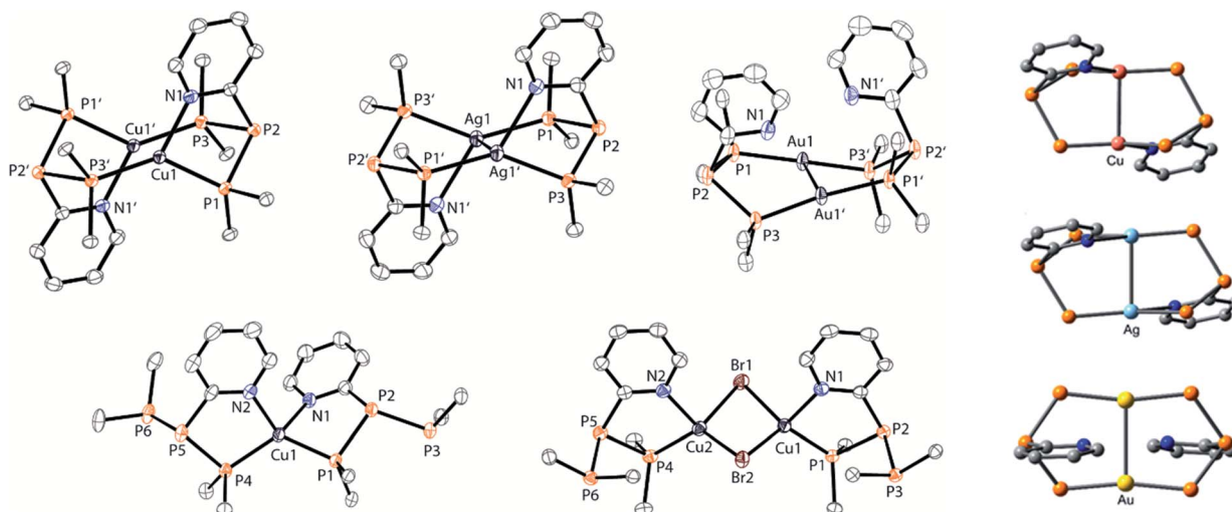


Fig. 4 Molecular structures of the coinage metal complexes [(**9a***Cu)₂][OTf]₂, [(**9a***Ag)₂][OTf]₂, [(**9a***Au)₂][OTf]₂, [(**9a**)₂*Cu][OTf] and (**9a***CuBr)₂ (hydrogen atoms, anions and solvate molecules are omitted for clarity, cyclohexyl groups are represented only by their phosphorus bound carbon atoms, and thermal ellipsoids are displayed at 50% probability; left). Selected geometrical parameters are given in Table 2; Optimised geometries of [(**9a***M)₂][OTf]₂ using the BP86–D3/def2–TZVP functional, showing the different coordination arrangement of the pyridyl-moieties towards the metal atoms (M = Cu, Ag, and Au; right).



Table 2 Selected geometrical parameters of crystallographically characterised coinage metal complexes $[(9a^*Cu)_2][OTf]_2$, $[(9a^*Ag)_2][OTf]_2$, $[(9a^*Au)_2][OTf]_2$, $[(9a)_2^*Cu][OTf]$ and $(9a^*CuBr)_2^a$

	$[(9a^*Cu)_2][OTf]_2$	$[(9a^*Ag)_2][OTf]_2$	$[(9a^*Au)_2][OTf]_2$	$[(9a)_2^*Cu][OTf]^a$	$(9a^*CuBr)_2^a$
M1–M1' in Å	2.8225(5)	2.9013(4)	2.880(3)	—	2.942
M1–P1 in Å	2.2755(4)	2.3818(8)	2.3033(8)	2.243	2.191
M1–P3' in Å	2.2176(5)	2.4468(7)	2.3198(8)	—	—
M–N in Å	2.048(1)	2.404(3)	2.948(3)	2.106	2.116
P–P ^a in Å	2.225	2.224	2.226	2.212	2.214
P1–M1–P3 in °	136.52(2)	143.96(3)	171.52(3)	—	—
P–P–P in °	98.26(2)	100.16(4)	98.29(5)	104.43	102.43

^a Average bond lengths and angles are given.

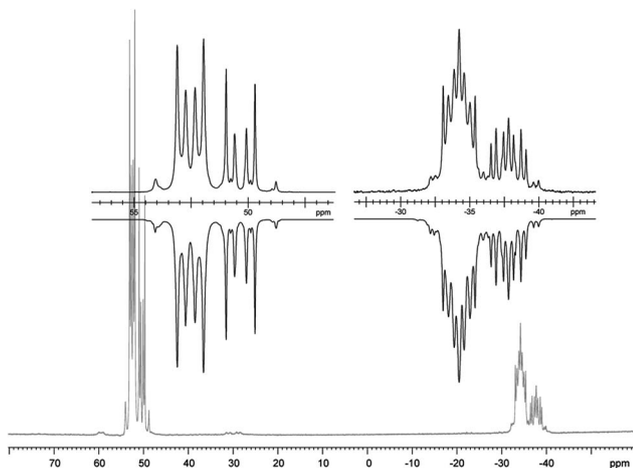


Fig. 5 ^{31}P NMR spectrum of $[(9a^*Au)_2][OTf]_2$ (CD_2Cl_2 , 243 K, insets show the two $AA'XX'X''X'''$ spin systems of the experimental (upwards) and fitted spectra (downwards)).

also observed. However, further line splitting can be observed as a result of the complexation with the Ag^{107}/Ag^{109} nuclei, which made us refrain from iteratively fitting the corresponding ^{31}P NMR spectrum. The ^{31}P NMR spectrum of the dinuclear copper complex $[(9a^*Cu)_2][OTf]_2$ shows only the resonances of one $AA'XX'X''X'''$ spin system at $\delta(P_A) = -39.1$ ppm and $\delta(P_X) = 15.4$ ppm. An additional broadened AX_2 spin system ($\delta(P_A) = -37.5$ ppm and $\delta(P_X) = 15.3$ ppm ($^1J(P_AP_X) = -168$ Hz)) gives rise to the formation of a monomeric species in solution which was not further investigated. A comparable AX_2 spin system is observed for $[(9a)_2^*Cu][OTf]$ ($\delta(P_A) = -44.8$ ppm and $\delta(P_X) = -8.9$ ppm ($^1J(P_AP_X) = -275$ Hz)) as well as for $(9a^*CuBr)_2$ ($\delta(P_A) = -46.6$ ppm and $\delta(P_X) = -6.2$ ppm ($^1J(P_AP_X) = -253$ Hz)). For both mononuclear copper complexes the resonances in the ^{31}P NMR spectrum are noticeably broadened due to the fast quadrupole relaxation of the ^{63}Cu nucleus.^{22,23} Theoretical calculations performed at the BP86-D3/def2-TZVP level of theory support the isomerisation in the dinuclear metal complexes $[(9a^*M)_2]^{2+}$. For $[(9a^*Cu)_2]^{2+}$ we found a relative energy difference between the *syn*- and the *anti*-isomer of 5.0 kcal mol⁻¹ which explains that only one isomer is observed in the ^{31}P NMR spectrum. While the two isomers of $[(9a^*Ag)_2]^{2+}$ are isoenergetic with a difference of 0.1 kcal mol⁻¹, the *syn*-isomer of

$[(9a^*Au)_2]^{2+}$ is favored by 1.7 kcal mol⁻¹ compared to its *anti*-isomer. The optimised geometries also show the different coordination behaviour of the pyridyl-moieties in the three metal complexes $[(9a^*M)_2]^{2+}$. While the free electron pair of the pyridyl nitrogen atom is pointing towards the metal centre in $[(9a^*Cu)_2]^{2+}$ and $[(9a^*Ag)_2]^{2+}$ it is pointing towards the middle of the Au–Au bond in $[(9a^*Au)_2]^{2+}$ (Fig. 4, right). Our calculations furthermore indicate that the isomerisation is not driven by rotation but proceeds *via* dissociation of the M(I)–N coordination bond or *pseudo*-coordination in the case of $[(9a^*Au)_2]^{2+}$ and subsequent association of the ligand on the opposite side (Fig. 6). This interconversion is the highest in energy for $[(9a^*Cu)_2]^{2+}$ (14.7 kcal mol⁻¹), followed by $[(9a^*Ag)_2]^{2+}$ (13.2 kcal mol⁻¹) and finally $[(9a^*Au)_2]^{2+}$ (9.6 kcal mol⁻¹).²³

Further reactivity studies on triphosphane **9a** revealed that this compound readily rearranges in solution by means of a P–P/P–P bond metathesis reaction. After 24 h the CH_2Cl_2 solution of **9a** shows an additional singlet resonance at $\delta(P) = 21.0$ ppm which we assign to tetracyclohexyldiphosphane (**12**)²⁴ and an $AA'BB'C$ spin system at $\delta(P) = 14.0$ – 25.2 ppm being characteristic of pentaphospholanes of type $(RP)_5$.^{24,25} As this scrambling reaction is in a state of equilibrium the isolation of **13** is hampered. However, the reaction of pure **13** with Cy_4P_2 yielding **9a** illustrates that pentaphospholane **13** might be used as a PyP-synthon as it inserts into the P–P bond of Cy_4P_2 . The selective synthesis of pentaphospholane **13** is achieved by the equimolar reaction of **7** with Cy_2PH in Et_2O as a non-polar solvent (Scheme 6). The formed precipitate is filtered and

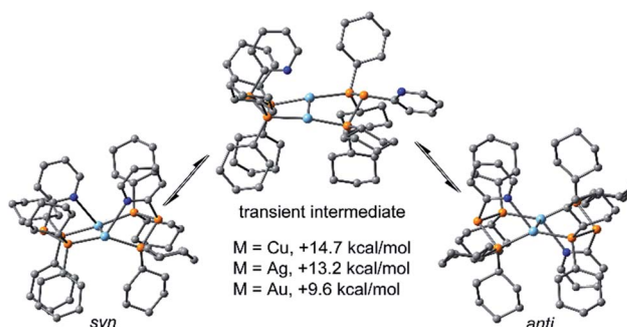
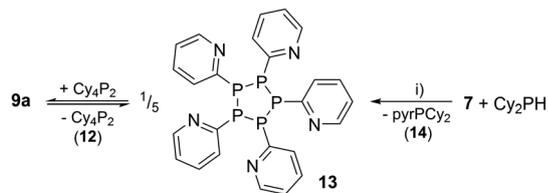


Fig. 6 Interconversion mechanism of the *syn*- and *anti*-isomers of $[(9a^*M)_2]^{2+}$; (M = Cu, Ag, and Au).





Scheme 6 Equilibrium reaction of **9a** with **13** and **12** and selective synthesis of **13** by the reaction of **7** with Cy_2PH ; (i) Et_2O , $-\text{pyrH}$.

dried *in vacuo* giving **13** in 98% yield as a colourless powder which shows the AA'BB'C spin system in the ^{31}P NMR spectrum (Fig. 7). Dicyclohexyl(3,5-dimethylpyrazolyl)phosphane (**14**) is identified as a side product in the supernatant solution by its singlet resonance in the ^{31}P NMR spectrum ($\delta(\text{P}) = 58.4$ ppm) indicating a P-N/P-P bond metathesis reaction.⁶ To further understand the formation of **13** we studied the reaction in CD_2Cl_2 by means of ^{31}P NMR spectroscopy (Fig. 8). After 3.5 h compound **7** is still present in the solution while Cy_2PH is fully consumed by the condensation reaction with **7** to form triphosphane **9a** (*vide supra*) and diphosphane **15** which is identified by an AX spin system ($\delta(\text{P}_\text{A}) = 10.2$ ppm and $\delta(\text{P}_\text{X}) = 33.4$ ppm; $^1J(\text{P}_\text{A}\text{P}_\text{X}) = -281$ Hz). The AMX spin system at $\delta(\text{P}_\text{A}) = -35.6$ ppm, $\delta(\text{P}_\text{M}) = -7.2$ ppm and $\delta(\text{P}_\text{X}) = 29.9$ ppm is assigned to triphosphane **16** which is formed *via* a P-N/P-P bond metathesis reaction of two equivalents of **15** and *via* a P-N/P-P bond metathesis reaction of **7** and **9a** (Scheme 7). In both cases phosphane **14** is liberated. We propose a further chain growth in which triphosphane **16** reacts with **15** with the release of phosphane **14** in a P-N/P-P oligomerisation reaction which ultimately yields pentaphospholane **13**. We furthermore propose the formation of tetraphosphetane **17** ($\delta(\text{P}) = -49.4$ ppm) which can be formed *via* the reaction of two equivalents **16** with concomitant release of **14** (Scheme 7). Similar to the known ring expansion reactions of certain cyclophosphanes,²⁶ **17** might react to form pentaphospholane **13**, being the

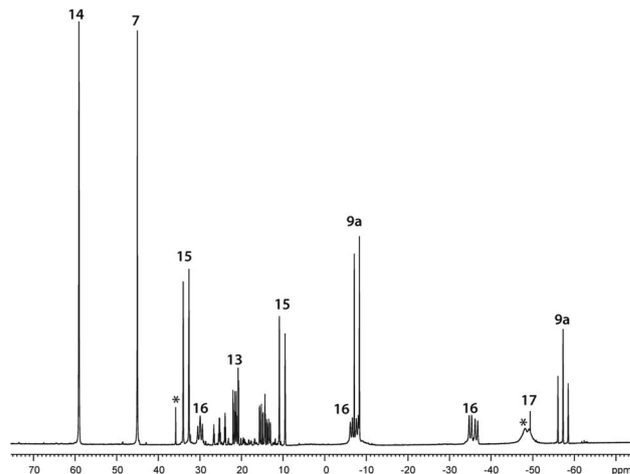


Fig. 8 ^{31}P NMR spectrum of the reaction of **7** with Cy_2PH in a 1 : 1 ratio in CD_2Cl_2 after 3.5 h; unidentified products are marked with asterisks.

thermodynamically favoured product. In analogy to **13** we are able to isolate the corresponding benzothiazolyl substituted pentaphospholane **18**.²⁷ Crystals suitable for X-ray analysis are obtained for both compounds by the slow diffusion of *n*-pentane into saturated CH_2Cl_2 solutions at -30 °C (Fig. 9).

Both compounds reveal the typical envelope conformation of the P_5 -ring featuring the substituents in the all-*trans* position. The P-P bond lengths observed for both pentaphospholanes **13** (av. P-P 2.218 Å) and **18** (av. P-P 2.246 Å) are in a range known for pentaphospholanes such as $(\text{PhP})_5$ (av. 2.217 Å); also the P-P-P angles are in good accordance with the reported data for $(\text{PhP})_5$.²⁸ We subsequently envisioned the use of **13** as a PyP-synthon for insertion reactions into P-P bonds which was already indicated by the aforementioned scrambling reaction between **9a**, Cy_4P_2 and **13** (see Scheme 6).

As similar scrambling reactions might be expected for triphosphonium and triphosphanedium salts, we methylated triphosphane **9a** with different amounts of MeOTf. The ^{31}P

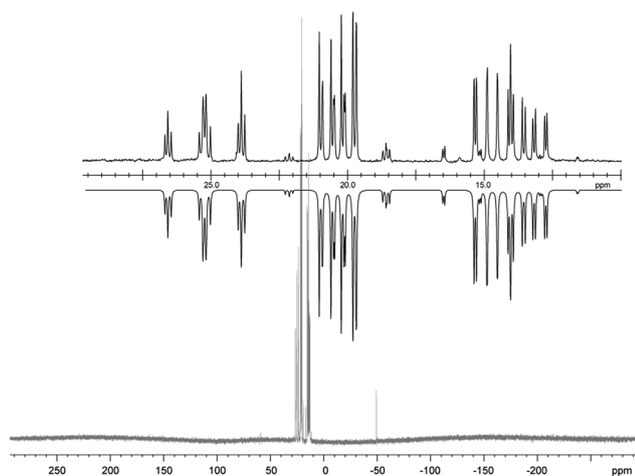
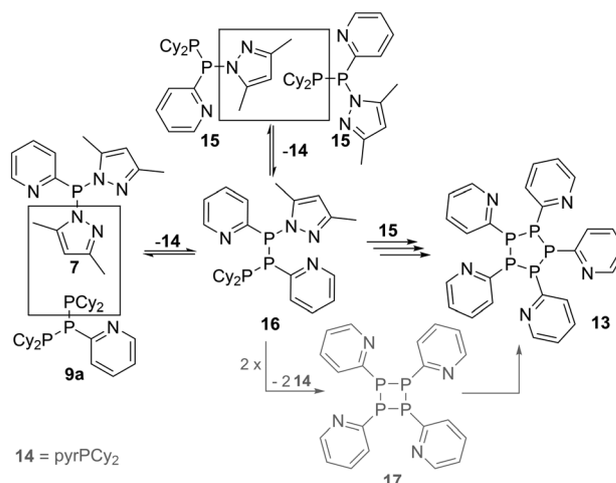


Fig. 7 ^{31}P NMR spectrum of pentaphospholane **13** (CD_3CN , 300 K, the inset shows the AA'BB'C spin system of the experimental (upwards) and fitted spectra (downwards)).



Scheme 7 Possible formation of **16** *via* P-N/P-P bond metathesis reactions.



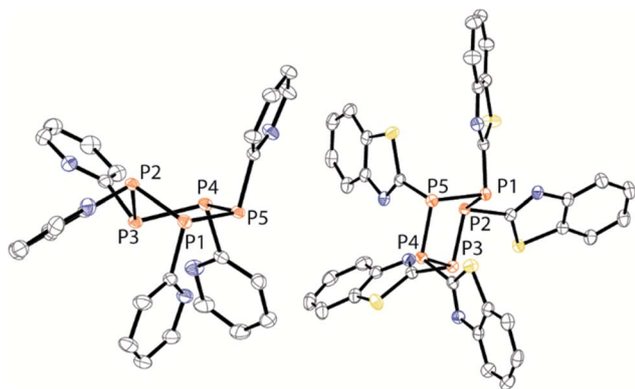
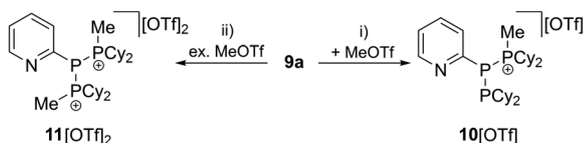


Fig. 9 Molecular structures of the pentaphosphanes **13** (left) and **18** (right) (hydrogen atoms and solvate molecules are omitted for clarity and thermal ellipsoids are displayed at 50% probability). Selected bond lengths (Å) and angles ($^{\circ}$) (**13**): P1–P2 2.2222(4), P2–P3 2.2200(5); P3–P4 2.22200(5), P4–P5 2.2078(4), P1–P2–P3 97.25(2), P2–P3–P4 95.59(2), P3–P4–P5 102.27(2), P4–P5–P1 106.77(2), P5–P1–P2 97.14(2) (**18**): P1–P2 2.2336(6), P2–P3 2.2232(6), P3–P4 2.338(6), P4–P5 2.2069(6), P5–P1 2.2288(6), P1–P2–P3 93.54(2), P2–P3–P4 100.29(2), P3–P4–P5 107.27(2), P4–P5–P1 96.32(2), P5–P1–P2 96.85(2).

NMR spectrum of the 1 : 1 reaction of **9a** with MeOTf shows an AMX spin system ($\delta(\text{P}_A) = -46.2$ ppm, $\delta(\text{P}_M) = -12.9$ ppm and $\delta(\text{P}_X) = 34.2$ ppm ($^1J(\text{P}_A\text{P}_M) = -290$ Hz, $^1J(\text{P}_A\text{P}_X) = -281$ Hz, and $^2J(\text{P}_M\text{P}_X) = 58$ Hz)), suggesting the methylation of the phosphorus atom rather than the nitrogen atom of the pyridyl-substituent. After the removal of all volatiles *in vacuo* triphosphan-1-ium triflate salt **10**[OTf] can be isolated in quantitative yield as a colourless powder. A second methylation is achieved when **9a** is reacted with an excess of MeOTf under solvent free conditions (Scheme 8). Similar to the aforementioned reaction, only the phosphorus atoms are methylated giving triphospane-1,3-diumtriflate salt **11**[OTf]₂. As both dicyclohexylphosphanyl moieties are methylated, the ^{31}P NMR spectrum of **10**[OTf]₂ shows an AX₂ spin system with resonances at $\delta(\text{P}_A) = -67.9$ ppm and $\delta(\text{P}_X) = 44.0$ ppm ($^1J(\text{P}_A\text{P}_X) = -315$ Hz). The molecular structures of cation **10**⁺ and dication **11**²⁺ are depicted in Fig. 10. Note that a small number of triphosphanediium salts are reported in the literature and compound **10**[OTf] extends the library of cationic, catenated phosphorus compounds as a structurally characterised triphosphan-1-ium salt.^{29–31} The P1–P2 bond length in cation **10**⁺ (P1–P2 2.1956(5) Å) is shortened whereas the P2–P3 bond (P2–P3 2.2362(5) Å) is comparable to those in **9a** (P1–P2 2.2301(3) Å, P1–P3 2.2371(3) Å) or similar diphosphanium cations (compare [Ph₂P–PPh₃]⁺:



Scheme 8 Methylation reactions of **9a**; (i) 1 eq. MeOTf, Et₂O; (ii) 5 eq. MeOTf, neat.

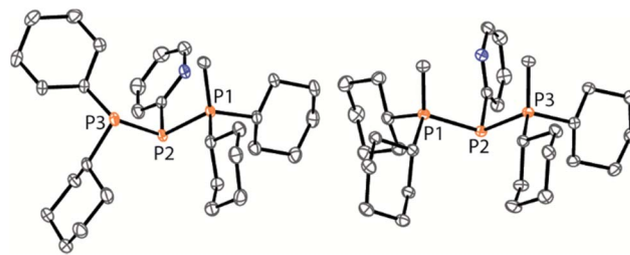
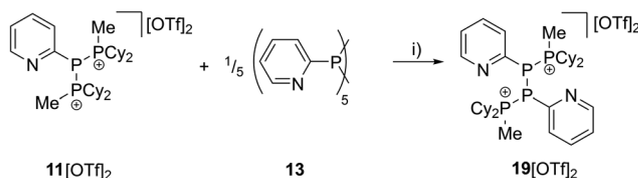


Fig. 10 Molecular structures of **10**⁺ in **10**[OTf] (left) and **11**²⁺ in **11**[OTf]₂ (hydrogen atoms and anions are omitted for clarity and thermal ellipsoids are displayed at 50% probability). Selected bond lengths (Å) and angles ($^{\circ}$) (**10**⁺): P1–P2 2.1956(5), P2–P3 2.2362(5), P1–P2–P3 97.89(2) (**11**²⁺): P1–P2 2.2152(4), P2–P3 2.2182(4), and P1–P2–P3 107.43(1).

2.230(1) Å).^{29b} The P–P–P bond angle in **10**⁺ (97.89(2) $^{\circ}$) is similar to that in compound **9a** (97.16(6) $^{\circ}$). The P–P bond lengths in **11**²⁺ (P1–P2 2.2152(4) Å and P2–P3 2.2182(4) Å) are marginally shorter compared to those in **9a** but in accordance with bond lengths reported for similar triphospane-1,3-dium cations (compare [Me₃P–P(Cy)–PMe₃]²⁺: 2.1979(5) Å and 2.1976(6) Å).^{29b} The P–P–P bond angle in **11**²⁺ (107.43(1) $^{\circ}$) is significantly wider compared to those in **9a** and [Me₃P–P(Cy)–PMe₃]²⁺ (103.11(2) $^{\circ}$). In order to investigate the envisioned PyP-transfer into a P–P bond, we reacted **13** with 5 equivalents of **11**[OTf]₂ in CH₃CN (Scheme 9). Upon addition of **13** to a colourless CH₃CN solution of **11**[OTf]₂ the reaction mixture turns from deep red to pale yellow. The ^{31}P NMR spectrum of the reaction mixture after 24 h shows two AA'XX' spin systems which are attributed to two isomers of tetraphospane-1,4-dium cations **19**²⁺ (Fig. 11), indicating the successful PyP-transfer *via* a P–P/P–P bond metathesis reaction. The A part of the prominent AA'XX' spin system at $\delta(\text{P}_A) = -55.1$ ppm is assigned to the inner pyridyl-substituted P nuclei and the X part at $\delta(\text{P}_X) = 40.7$ ppm to the tetra-coordinate phosphorus atoms which is similar to known tetraphospane-1,4-dium salts.³² The minor spin system resonates at $\delta(\text{P}_A) = -40.1$ ppm and $\delta(\text{P}_X) = 39.8$ ppm, respectively. After work-up compound **19**[OTf]₂ can be isolated analytically pure in 51% yield. X-ray analysis revealed that the centrosymmetric *meso*-isomer **19**²⁺ crystallized showing a central torsion angle of 180 $^{\circ}$ (Fig. 11). The P–P bond lengths in **19**²⁺ (P1–P2 2.234(1) Å and P2–P2' 2.232(2) Å) are in good agreement with the reported data for comparable tetraphospane-1,4-dium cations (compare [Ph₃P–(PPh)₂–PPh₃]²⁺: P1–P2 2.258(1) Å, P2–P2' 2.221(1) Å).³² Similar to the reported PhP-transfer from (PhP)₅ to a NHC,³³ **13** can be used as a PyP-synthon, thus featuring an



Scheme 9 Synthesis of tetraphospane-1,4-dium triflate salt **19**[OTf]₂; (i) MeCN.



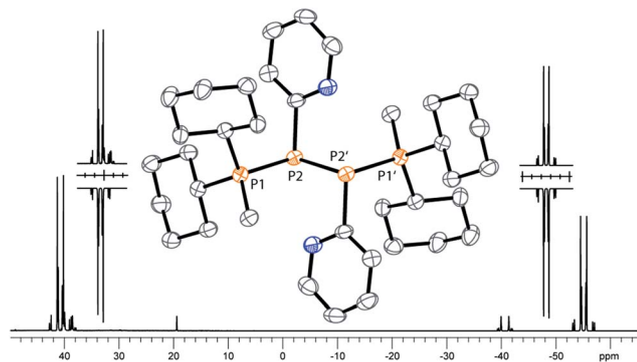


Fig. 11 ^{31}P NMR spectrum of dication $19[\text{OTf}]_2$ (CD_2Cl_2 , 300 K, the inset shows the AA'XX' spin system of the experimental (upwards) and fitted spectra (downwards)); molecular structure of 19^{2+} in $19[\text{OTf}]_2$ (hydrogen atoms and anions are omitted for clarity; thermal ellipsoids are displayed at 50% probability). Selected bond lengths (\AA) and angles ($^\circ$): P1–P2 2.234(1), P2–P2' 2.232(2), and P1–P2–P2' 97.60(6).

additional reaction or coordination site. When compound **19** $[\text{OTf}]_2$ was kept in a CD_3CN solution for 14 days, the ^{31}P NMR spectrum shows three rearrangement products (Fig. 12). Next to an AX_2 spin system ($\delta(\text{P}_\text{A}) = -235.3$ ppm and $\delta(\text{P}_\text{X}) = 38.8$ ppm ($^1J(\text{P}_\text{A}\text{P}_\text{X}) = -501$ Hz)), a *pseudo* triplet resonance ($\delta(\text{P}) = 153.1$ ppm; $^1J(\text{PN}) = 50$ Hz) and an AMXY spin system ($\delta(\text{P}_\text{A}) = -208.2$ ppm, $\delta(\text{P}_\text{M}) = -30.0$ ppm, $\delta(\text{P}_\text{X}) = 31.1$ ppm, and $\delta(\text{P}_\text{Y}) = 39.8$ ppm; $^1J(\text{P}_\text{A}\text{P}_\text{Y}) = -490$ Hz, $^1J(\text{P}_\text{M}\text{P}_\text{X}) = -342$ Hz, $^1J(\text{P}_\text{A}\text{P}_\text{M}) = -298$ Hz, $^2J(\text{P}_\text{M}\text{P}_\text{Y}) = 91$ Hz, $^2J(\text{P}_\text{A}\text{P}_\text{X}) = 45$ Hz, $^3J(\text{P}_\text{X}\text{P}_\text{Y}) = 34$ Hz) are observed. The AX_2 spin system can be attributed to the $3\lambda^5$ -triphosph-2-en-1-ium cation (20^+), as the spectroscopic parameters are in the range of those observed for $[\text{Ph}_3\text{P}-\text{P}-\text{PPh}_3]^+$ ($\delta(\text{P}_\text{A}) = -174$ ppm and $\delta(\text{P}_\text{X}) = 30$ ppm; $^1J(\text{P}_\text{A}\text{P}_\text{X}) = -500$ Hz) which was first reported by Schmidpeter and co-workers.³⁴ A variety of similar derivatives has been synthesised and reported by the group of Macdonald.³⁵ Slow vapour diffusion of Et_2O into the CD_3CN solution yielded crystals suitable for X-ray analysis confirming the structural connectivity of 20^+ (Fig. 13). The

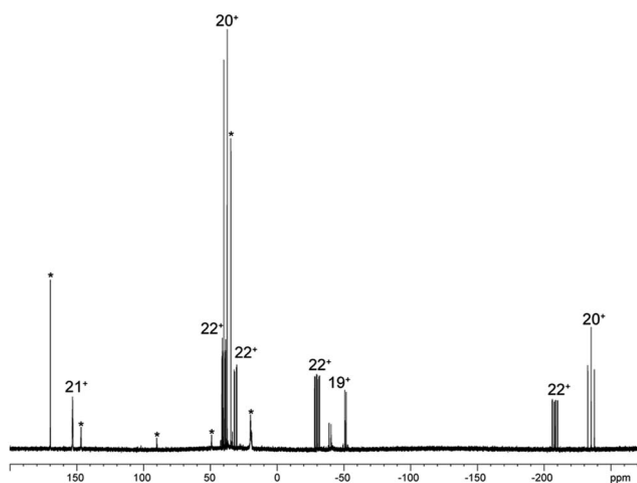


Fig. 12 ^{31}P NMR spectrum of a solution of $19[\text{OTf}]_2$ in CD_3CN after 14 days.

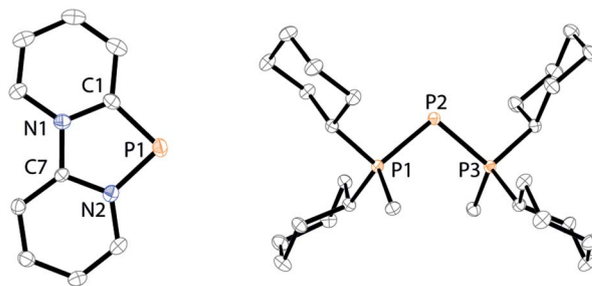


Fig. 13 Molecular structure of 1,4,2-diazaphospholium 21^+ in $21[\text{OTf}]$ (left) and $3\lambda^5$ -triphosph-2-en-1-ium 20^+ in $20[\text{OTf}]$ (right) (hydrogen atoms and anions are omitted for clarity and thermal ellipsoids are displayed at 50% probability). Selected bond lengths (\AA) and angles ($^\circ$) (20^+): P1–C1 1.733(1), P1–N2 1.741(1), C1–P1–N2 87.08(6) (20); P1–P2 2.134(4), P2–P3 2.1382(4), P1–P2–P3 103.85(2).

structural parameters of 20^+ are similar to those observed for the cation $[\text{Ph}_3\text{P}-\text{P}-\text{PPh}_3]^+$, *i.e.* slightly shortened P–P bond lengths (P1–P2 2.134(4), P2–P3 2.1382(4)) and a P–P–P bond angle of $103.85(2)^\circ$.³⁴ By co-crystallisation, a further product was identified as the triflate salt of diazaphospholium cation 21^+ (Fig. 13). We assign this species to the *pseudo* triplet resonance observed in the ^{31}P NMR spectrum. The P–C bond distance of 1.733(1) \AA slightly exceeds the upper limit of typical P=C bond lengths (1.61–1.71 \AA),³⁶ while the P–N bond length of P1–N2 1.741(1) \AA indicates a P–N single bond (P–N: 1.78 \AA).³⁷ Furthermore, the structure of 21^+ reveals an acute angle around the P atom (C1–P1–N2 $87.08(6)^\circ$) and a planar arrangement, suggesting a delocalized π -system. We assign the AMXY spin system to asymmetric tetraphosphanedium dication 22^{2+} , most likely formed by an intramolecular aromatic substitution reaction of 19^{2+} . This can be attributed to be the first step of the rearrangement reaction of 19^{2+} to form 20^+ and 21^+ . We further studied this rearrangement reaction using the TURBOMOLE 7.0 program at the PB86-D3/def2-TZVP level of theory (see ESI for details[†]) and taking into consideration solvent effects by using the Conductor-like Screening Model (COSMO). Our calculations started from 22^+ and show that the cleavage of C_2PMe is energetically quite costly. This is in accordance with the ^{31}P NMR spectrum as C_2PMe is not observed ($\delta(\text{P}) = -19.8$ ppm).³⁸ More likely and energetically favoured is a [1,2]-shift of the C_2PMe group to form *iso*- 22^{2+} with a reaction barrier of +22.0 kcal mol⁻¹. Fig. 14 shows the optimised structures for this [1,2]- C_2PMe shift reaction. It is noteworthy that similar [1,2]- and [1,3]- PR_3 shifts have been reported.³⁹ Compared to 22^{2+} , *iso*- 22^{2+} is almost isoenergetic and

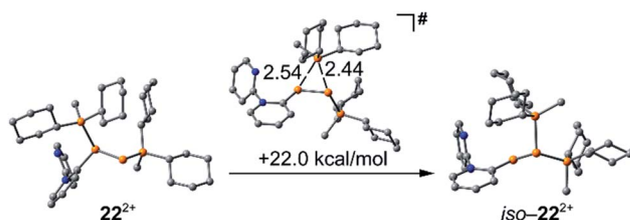
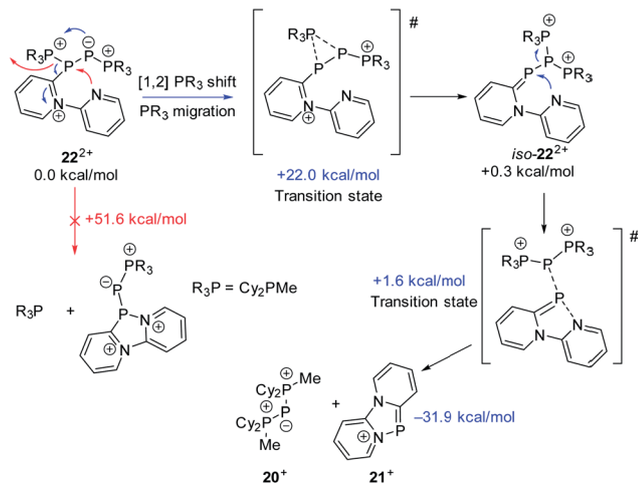


Fig. 14 Optimised structures of 22^{2+} , *iso*- 22^{2+} and the transition state.





Scheme 10 Rearrangement reaction of 22^{2+} to 20^+ and 21^+ calculated at the PB86-D3/def2-TZVP level of theory.

readily reacts in an exergonic step ($-31.9 \text{ kcal mol}^{-1}$) to form cations 20^+ and 21^+ with a low barrier of only $1.9 \text{ kcal mol}^{-1}$ (see Scheme 10 and ESI for details[†]).

Conclusions

In summary we have shown that phosphanes which we denote as type II phosphanes (two P–N bonds and one P–C bond) can be used to build polyphosphorus compounds featuring a further nitrogen donor functionality at the C bound substituent. The reactions of type II phosphanes **7** and **8** with secondary phosphanes of type R'_2PH in a 1 : 2 ratio in MeCN give triphosphanes **9a–c** and triphospholane **9d** via protolysis reactions. With the multi-gram scale synthesis of **9a** we were able to conveniently investigate its coordination chemistry towards coinage metal salts. The synthesised coordination complexes are studied by X-ray analysis, multi nuclear NMR spectroscopy and quantum chemical calculations. Further reactivity studies of **9a** focused on the reaction with the electrophilic MeOTf. Depending on the amount of MeOTf, the reaction yields triphosphane-1,3-dium triflate salt **11**[OTf]₂ or triphosphan-1-ium **10**[OTf] featuring a hitherto unknown structural motif. Pentaphospholanes **13** and **18** are synthesised by reacting type II phosphane **7** or **8** with Cy_2PH in a 1 : 1 ratio in Et_2O or CH_2Cl_2 , respectively. The reaction proceeds via a controlled scrambling reaction showing the susceptible solvent dependency of the reaction of **7** and **8** with Cy_2PH . By reacting **11**[OTf]₂ with **13** we have shown that pentaphospholane **13** is a suitable PyP-synthon in a P–P/P–P bond metathesis reaction to yield tetraphosphane-1,4-dium triflate **19**[OTf]₂. In an interesting rearrangement reaction **19**[OTf]₂ forms triphosph-2-en-1-ium triflate salt **20**[OTf] and 1,4,2-diazaphospholium triflate **21**[OTf] via an unprecedented [1,2]- Cy_2MeP shift. Based on our findings we are certain that controlled P–N(P)/P–P bond metathesis reactions will allow us to access larger acyclic and cyclic polyphosphanes in

a controlled stepwise manner in order to further investigate their fascinating reaction space.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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Notes and references

- 1 M. Donath, F. Hennersdorf and J. J. Weigand, *Chem. Soc. Rev.*, 2016, **45**, 1145; M. H. Holthausen and J. J. Weigand, *Chem. Soc. Rev.*, 2014, **43**, 6639; S. Gomez-Ruiz and E. Hey-Hawkins, *Coord. Chem. Rev.*, 2011, **255**, 1360; M. Scheer, G. Balazs and A. Seitz, *Chem. Rev.*, 2010, **110**, 4236; M. Baudler, *Angew. Chem., Int. Ed. Engl.*, 1987, **26**, 419.
- 2 (a) M. Baudler, *Angew. Chem., Int. Ed. Engl.*, 1982, **21**, 492; M. Baudler and K. Glinka, *Chem. Rev.*, 1993, **93**, 1623. ; M. Baudler and K. Glinka, *Chem. Rev.*, 1994, **94**, 1273. (b) M. Scheer, St. Gremler, E. Herrmann, U. Grünhagen, M. Dargatz and E. Kleinpeter, *Z. Anorg. Allg. Chem.*, 1991, **600**, 203; M. Scheer, St. Gremler, E. Herrmann, M. Dargatz and H.-D. Schädler, *Z. Anorg. Allg. Chem.*, 1992, **619**, 1047. (c) G. Fritz, *Comments Inorg. Chem.*, 1982, **6**, 329; G. Fritz, *Adv. Inorg. Chem.*, 1987, 171. (d) L. Maier, *Helv. Chim. Acta*, 1966, **49**, 1119; E. Niecke, R. Rüger and B. Krebs, *Angew. Chem., Int. Ed. Engl.*, 1982, **21**, 544; A. B. Burg, *J. Am. Chem. Soc.*, 1961, **83**, 2226; K. Jurkschat, C. Mugge, A. Tzschach, W. Uhlig and A. Zschunke, *Tetrahedron Lett.*, 1982, **23**, 1345.
- 3 M. Baudler, G. Reuschenbach and J. Hahn, *Z. Anorg. Allg. Chem.*, 1981, **482**, 27; G. Fritz and T. Vaahs, *Z. Anorg. Allg. Chem.*, 1987, **553**, 85.
- 4 A. H. Cowley and D. S. Dierdorf, *J. Am. Chem. Soc.*, 1965, **87**, 6609; L. R. Avend, R. A. Wolcott, L. V. Cribbs and J. L. Mills, *Inorg. Chem.*, 1989, **28**, 200; L. R. Avens, L. V. Cribbs and J. L. Mills, *Inorg. Chem.*, 1989, **28**, 211.
- 5 K.-O. Feldmann, R. Fröhlich and J. J. Weigand, *Chem. Commun.*, 2012, **48**, 4296.
- 6 K.-O. Feldmann and J. J. Weigand, *J. Am. Chem. Soc.*, 2012, **134**, 15443.
- 7 (a) X. Chen, H. Zhu, T. Wang, C. Li, L. Yan, M. Jiang, J. Liu, X. Sun, Z. Jiang and Y. Ding, *J. Mol. Catal. A: Chem.*, 2016, **414**, 37; (b) I. V. Komarov, A. V. Strizhak, M. Y. Kornilov, E. Zraudnitskiy and A. A. Tolmachev, *Synth. Commun.*, 2000, **30**, 243.



- 8 We developed a convenient, salt free synthesis of **5** and **6** which is described in the ESI.†
- 9 (a) S. Fischer, L. K. Peterson and J. F. Nixon, *Can. J. Chem.*, 1974, **52**, 3981; (b) S. Fischer, J. Hoyano and L. K. Peterson, *Can. J. Chem.*, 1976, **54**, 2710.
- 10 H. Goldwhite, *Introduction to phosphorus chemistry*, Cambridge University Press, Cambridge, 1981.
- 11 R. E. Cobblestick and F. W. B. Einstein, *Acta Crystallogr.*, 1975, **B31**, 2731.
- 12 J. J. Daly, *J. Chem. Soc. A*, 1966, 1020.
- 13 S. Gomez-Ruiz and E. Hey-Hawkins, *Coord. Chem. Rev.*, 2011, **255**, 1360; *Phosphorus Chemistry: Catalysis and Material Science Applications*, ed. M. Peruzzini and L. Gonsalvi, Springer, vol. 37, 2011.
- 14 J. W. Dube, C. L. B. Macdonald and P. Ragoonna, *Angew. Chem., Int. Ed.*, 2012, **51**, 13026; J. W. Dube, C. L. B. Macdonald, B. D. Ellis and R. Paul, *Inorg. Chem.*, 2013, **52**, 11438.
- 15 M. Scheer, C. Kuntz, M. Stubenhofer, M. Zabel and A. Y. Timoshkin, *Angew. Chem., Int. Ed.*, 2010, **49**, 188; M. Baacke, S. Morton, G. Johannsen, N. Weferling and O. Stelzer, *Chem. Ber.*, 1980, **113**, 1328; W. S. Sheldrick, S. Morton and O. Stelzer, *Z. Anorg. Allg. Chem.*, 1981, **475**, 232; M. Scheer, S. Gremler, E. Herrmann and P. G. Jones, *J. Organomet. Chem.*, 1991, **414**, 337.
- 16 D. R. Armstrong, N. Feeder, A. D. Hopkins, M. J. Mays, D. Moncrieff, J. A. Wood, A. D. Woods and D. S. Wright, *Chem. Commun.*, 2000, 2483.
- 17 C. E. Averre, M. P. Coles, I. R. Crossley and I. J. Day, *Dalton Trans.*, 2012, **41**, 278.
- 18 For a review on argentophilic interactions see: H. Schmidbaur and A. Schier, *Angew. Chem., Int. Ed.*, 2015, **54**, 746, for aurophilic interactions see: H. Schmidbaur and A. Schier, *Chem. Soc. Rev.*, 2012, **41**, 370.
- 19 A. Bondi, *J. Phys. Chem.*, 1964, **68**, 441.
- 20 C.-M. Che, Z. Mao, V. M. Miskowski, M.-C. Tse, C.-K. Chan, K.-K. Cheung, D. L. Phillips and K.-H. Leung, *Angew. Chem., Int. Ed.*, 2000, **39**, 4084; C.-M. Che, M.-C. Tse, M. C. W. Chan, K.-K. Cheung, D. L. Phillips and K.-H. Leung, *J. Am. Chem. Soc.*, 2000, **122**, 2464; C.-M. Che, W.-F. Wu, K.-C. Chan and K.-K. Cheung, *Chem.-Eur. J.*, 2001, **7**, 4656.
- 21 M. Henary, J. L. Wootton, S. I. Khan and J. I. Zink, *Inorg. Chem.*, 1997, **36**, 796; P. Aslanidis, P. J. Cox, S. Divanidis and A. C. Tsipis, *Inorg. Chem.*, 2002, **41**, 6875.
- 22 H. Friebolin, *Ein- und Zweidimensionale NMR-Spektroskopie*, Wiley VCH, Weinheim, 2006; H. Fujii, M. Tomura, T. Kurahashi and M. Kujime, *Inorg. Chem.*, 2007, **46**, 541.
- 23 (a) The ³¹P NMR spectra of [(**9a***Cu)₂][OTf]₂, [(**9a***Ag)₂][OTf]₂, [(**9a***Au)₂][OTf]₂, [(**9a**)₂*Cu][OTf] and (**9a***CuBr)₂ are depicted in the ESI (Fig. 1–5†); (b) Further information on the theoretical calculations on [(**9a***M)₂][OTf]₂ is also given in the ESI in chapter 4.†
- 24 K. Schwedtmann, R. Schoemaker, F. Hennersdorf, A. Bauzá, A. Frontera, R. Weiss and J. J. Weigand, *Dalton Trans.*, 2016, **45**, 11384.
- 25 (a) J. P. Albrand, D. Gagnaire and J. B. Robert, *J. Am. Chem. Soc.*, 1973, **95**, 6498; (b) J. P. Albrand and J. B. Robert, *J. Chem. Soc., Chem. Commun.*, 1974, 644; (c) L. R. Smith and J. L. Mills, *J. Chem. Soc., Chem. Commun.*, 1974, **20**, 808.
- 26 (a) M. Baudler, J. Hahn and E. Clef, *Z. Naturforsch.*, 1984, **39b**, 438; (b) K. Schwedtmann, J. Haberstroh, S. Roediger, A. Bauzá, A. Frontera, F. Hennersdorf and J. J. Weigand, *Chem. Sci.*, 2019, **10**, 6868.
- 27 For further details see the ESI.†
- 28 J. J. Daly, *J. Chem. Soc.*, 1964, 6147.
- 29 (a) N. Burford, T. S. Cameron and P. J. Ragoonna, *J. Am. Chem. Soc.*, 2001, **123**, 7947; (b) P. A. Gray, Y.-Y. Carpenter, N. Burford and R. McDonald, *Dalton Trans.*, 2016, **45**, 2124.
- 30 For a review on interpnictogen cations see: A. P. M. Robertson, P. A. Gray and N. Burford, *Angew. Chem., Int. Ed.*, 2014, **53**, 6050.
- 31 For cationic, cyclic polyphosphorus compounds see: J. Bresien, K. Faust, A. Schulz and A. Villinger, *Angew. Chem., Int. Ed.*, 2015, **54**, 6926; S. J. Geier, M. A. Dureen, E. Y. Ouyang and D. W. Stephan, *Chem.-Eur. J.*, 2010, **16**, 988; Y.-Y. Carpenter, N. Burford, M. D. Lumsden and R. McDonald, *Inorg. Chem.*, 2011, **50**, 3342; J. J. Weigand, N. Burford, R. J. Davidson, T. S. Cameron and P. Seeheim, *J. Am. Chem. Soc.*, 2009, **131**, 17943; C. A. Dyker, N. Burford, G. Menard, M. D. Lumsden and A. Decken, *Inorg. Chem.*, 2007, **46**, 4277; N. Burford, C. A. Dyker, M. Lumsden and A. Decken, *Angew. Chem.*, 2005, **117**, 6352.
- 32 (a) C. A. Dyker, N. Burford, M. D. Lumsden and A. Decken, *J. Am. Chem. Soc.*, 2006, **128**, 9632; (b) Y.-Y. Carpenter, C. A. Dyker, N. Burford, M. D. Lumsden and A. Decken, *J. Am. Chem. Soc.*, 2008, **130**, 15732.
- 33 (a) A. J. Arduengo III, H. V. R. Dias and J. C. Calabrese, *Chem. Lett.*, 1997, 143; (b) A. J. Arduengo III, J. C. Calabrese, A. H. Cowley, H. V. R. Dias, J. R. Goerlich, W. J. Marshall and B. Riegel, *Inorg. Chem.*, 1997, **36**, 2151; (c) J. H. Barnard, P. A. Brown, K. L. Shuford and C. D. Martin, *Angew. Chem., Int. Ed.*, 2015, **54**, 12083.
- 34 A. Schmidpeter, S. Lochschmidt and W. S. Sheldrick, *Angew. Chem.*, 1985, **97**, 214.
- 35 B. D. Ellis, M. Carlesimo and C. L. B. Macdonald, *Chem. Commun.*, 2003, 1946; B. D. Ellis and C. L. B. Macdonald, *Inorg. Chem.*, 2006, **45**, 6864; E. L. Norton, K. L. S. Szekely, J. W. Dube, P. G. Bomben and C. L. B. Macdonald, *Inorg. Chem.*, 2008, **47**, 1196.
- 36 M. Regitz and O. J. Scherer, *Multiple Bonds and Low Coordination in Phosphorus Chemistry*, Georg Thieme Verlag Thieme Medical Publishers, New York, 1990.
- 37 H. R. Allcock, *Phosphorus-Nitrogen Compounds*, Academic Press, New York, 1972.
- 38 E. Payet, A. Auffrant, X. F. Le Goff and P. Le Floch, *J. Organomet. Chem.*, 2010, **695**, 1499.
- 39 E. Rüba, K. Mereiter, R. Schmid, V. N. Sapunov, K. Kirchner, H. Schottenberger, M. J. Calhorda and L. F. Veiros, *Chem.-Eur. J.*, 2002, **8**, 3948; C. Coletti, L. Gonsalvi, A. Guerriero, L. Marvelli, M. Peruzzini, G. Reginato and N. Rem, *Organometallics*, 2010, **29**, 5982.

