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Tuning the selectivity of P₄ reduction at alkalineearth metal centres†

Stefan Thum, Doliver P. E. Townrow, Dolern Langer Doliver P. E. Townrow, Doliver Doliver Doliver P. E. Townrow, Doliver Dolive

Reduction of P_4 with β -diketiminate Mg^1 complexes, (BDI)MgMg(BDI), depends strongly on the bulk of the ligand. Whereas superbulky BDI ligands gave selective reduction to ${P_4}^{2-}$ in a butterfly conformation, reduction with a less bulky ligand gave various products among which P_8^{4-} had a realgar-type structure. The selectivity of P_4 reduction can also be controlled by metal choice. Reduction of P_4 with Ca^1 synthons of general type (BDI*)Ca–X–Ca(BDI*) in which BDI* is a superbulky ligand and X is a bridging dianion ($C_6H_6^{2-} < p$ -xylene $^{2-} < N_2^{2-}$) led to reduction of P_4 to the very common, stable Zintl anion P_7^{3-} . Monitoring this process with ^{31}P NMR shows that $cyclo-P_4^{2-}$ is an intermediate en route to P_7^{3-} . Conversion rates increase with increasing reducing power: $X = C_6H_6^{2-} < p$ -xylene $^{2-} < N_2^{2-}$. A complex with the weakly reducing DBA $^{2-}$ dianion led to selective P_4 reduction to $cyclo-P_4^{2-}$ (DBA = 9,10-dimethyl-diboraanthracene). DBA inhibits P_4 -to- P_7 conversion, most likely by capturing the electron needed for further P_4 reduction by radical processes. Experimental investigations are supported by crystal structure determinations and a computational DFT study which also shows that the nature of metal- P_4 bonding (covalent or ionic) determines the preference for formation of butterfly-shaped P_4^{2-} or planar 6π -electron aromatic $cvclo-P_4^{2-}$.

Introduction

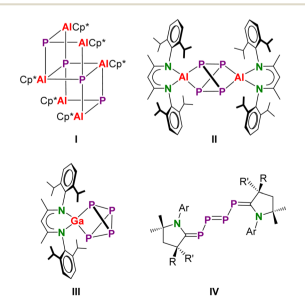
White phosphorus (P_4) is a commodity reagent for the production of industrially relevant P-containing products. While traditional bulk processes convert P_4 with highly corrosive Cl_2 to PCl_3 for further functionalization with polar organometallic reagents, current research initiatives aim for catalytic protocols to directly convert P_4 to organophosphorus compounds. In this light, the activation and chemical breakdown of P_4 is an important research field. Being a highly strained molecule, P_4 can be easily oxidized or reduced and shows diverse reactivity. Reacting either as an electrophile, nucleophile, or as an e-donor/acceptor, it could be seen as a chameleon in P-chemistry.

Although P_4 is inherently highly reactive, reaction pathways often remain unclear and selective conversions are difficult to achieve. Numerous groups have reported on P_4 activation using the rich redox reactivity of the transition metals.³⁻⁶ Recent developments in low-valent p-block chemistry stimulated P_4 activation with reagents that, due to small HOMO-LUMO gaps,

Inorganic and Organometallic Chemistry, Friedrich-Alexander-Universität Erlangen-Nürnberg, Egerlandstraße 1, 91058 Erlangen, Germany. E-mail: sjoerd. harder@fau.de

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show transition metal-like reactivity. ⁷⁻⁹ Earlier highlights of this work include the insertion of $(Me_3Si)_3CGa$ in three P–P bonds of P_4 by Uhl and coworkers ¹⁰ or the complete reduction of P_4 by Cp*Al to give a P^{3-} containing cluster $(Cp*Al)_6(P)_4$ (I, Scheme 1) by the Schnöckel group. ¹¹ In contrast, the bulkier β -diketiminate complex (BDI)Al reacted to give a complex of the P_4^{4-} anion



Scheme 1 Selected products of P_4 activation by low-valent main group compounds.

(II), formed by 4e-reduction and cleavage of two edges in the P_4 tetrahedron; BDI is herein defined as $HC[C(Me)-N(DIPP)]_2$ (DIPP = 2,6-diisopropylphenyl). Reaction with the softer reducing agent (BDI)Ga led only to 2e-reduction and cleavage of one P–P edge (III).¹² The valence isoelectronic cyclic alkyl amino carbenes (CAACs) have also been shown to activate P_4 (IV),¹³ whereas silylenes show either 4e-reduction giving P_4^{4-} , like in II, or 2e-reduction resulting in P_4^{2-} , like in III.¹⁴

Given the plethora of important breakthroughs in lowoxidation state s-block metal chemistry,15-17 it is remarkable that there is a complete lack of research on P4 reduction with these highly reactive early main group metal complexes. The reduction chemistry of one of the first MgI complexes, (BDI) MgMg(BDI),18 has been extensively investigated15 but we are unaware of reactivity studies with P4. However, this mild reducing agent has been reported to reduce the P5 ring in Cp*Fe(P5).19 The far majority of P4 activation studies with sblock metal reagents exploit their superb nucleophilicity. Classical examples include P-P bond cleavage in P4 by nucleophilic addition of RLi or Grignard reagents.20 An interesting case of nucleophilic activation of P4 is its reaction with nucleophilic hydride reagents like $[(BDI)Ca(\mu_2-H)]_2$ which after subsequent H2 release gave the reduction-like product [(BDI) $Ca_{3}(P_{7})$, containing the Zintl P_{7}^{3-} ion. In this example, the calcium hydride complex reacts as a synthon for the hitherto unknown Ca^I complex (BDI)CaCa(BDI). Such reductive reactivity of group 2 metal hydride complexes is well-established.²¹

We herein report a systematic study on the reduction of P_4 with Mg^I complexes or Ca^I synthons, *i.e.* Ca^{II} complexes containing electron-rich ligands that can react like the corresponding Ca^I species. $^{22-24}$ We demonstrate that selectivity is largely dependent on metal choice, ligand bulk or the nature of the electron-rich ligand delivering the electrons for P_4 reduction.

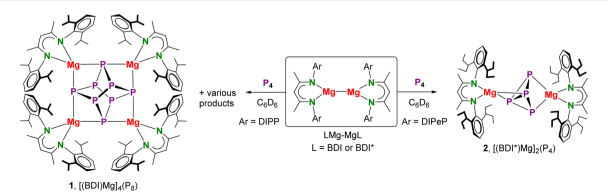
Results and discussion

P₄ activation with Mg^I complexes

The direct reduction of P_4 with β -diketiminate Mg^I complexes of type (BDI)MgMg(BDI) has so far not been described in the literature. This could be because 1H and $^{31}P\{^1H\}$ NMR monitoring of an equimolar mixture of $[(BDI)Mg]_2$ and P_4 in C_6D_6 at

room temperature showed a highly unselective conversion (Scheme 2 and Fig. S37/S38†). Various side-reactions may originate from the poor solubility of the reactants and the heterogeneity of the reaction mixture. However, we found that gently heating the mixture for three days at 60 °C leads to further conversion and selective crystallization of a most insoluble reaction product [(BDI)Mg]₄(P₈) (1) in 10% yield (Fig. 1a). This minor product shows 31P NMR resonances at +68.3 and +145.0 ppm which were not observed in the crude product of the room temperature [(BDI)Mg]₂/P₄ conversion. This means that 1 was formed after thermal treatment. The P₈⁴⁻ unit is isostructural and valence isoelectronic to α-P₄S₄, which is of the realgar-type,25 and flanked by four [(BDI)Mg]+ fragments at the corners with each Mg atom bound to two P atoms. Despite the high symmetry of the P₈⁴⁻ anion, the complex shows no crystallographic symmetry. Although this structure is unprecedented in s-block metal chemistry, a few examples for P₈⁴⁻ formation are known from transition metal,26-28 lanthanide,29 and gallium mediated P₄ activation.³⁰ Similar realgar-type polystibides Sb₈ have been isolated as the corresponding [(BDI) Mg]₄(Sb₈) complexes with different BDI ligands.³¹ The Mg-P distances in 1 are in the narrow range of 2.599(1)-2.692(1) Å with P-Mg-P bite angles varying from 73.92(3)° to 74.79(3)°. In contrast to [(BDI)Mg]₄(Sb)₈ which exhibits an almost linear Mg-Sb-Mg arrangement, the Mg-P-Mg angles in 1 deviate slightly from linearity: $164.26(3)-168.05(3)^{\circ}$. Within the P_8^{4-} anion in 1 there are two types of P-P bonds. The P-P bonds between threecoordinate P atoms, P2-P6 (2.2868(8) Å) and P4-P8 (2.2751(8) Å), are slightly elongated compared to the remaining P-P bonds ranging from 2.1999(7) to 2.2188(8) Å. The P-P-P angles range from 92.60(3)° to 102.67(3)°. These structural features are in agreement with those reported for the SmIII cluster $(Cp*_2Sm)_4(P_8).^{29}$

In contrast, reaction of P_4 with the more sterically hindered Mg^I complex $[(BDI^*)Mg]_2$, 32 featuring a considerably elongated Mg–Mg interatomic distance, 33 gave at room temperature overnight highly selective conversion; BDI^* is defined as HC $[C(Me)-N(DIPeP)]_2$ ($DIPeP=2,6-(Et_2CH)$ -phenyl). In contrast to the unselective $[(BDI)Mg]_2/P_4$ conversion (Fig. S37/S38†), the crude product of the $[(BDI^*)Mg]_2/P_4$ conversion showed a ^{31}P ^{1}H NMR spectrum with only two triplet resonances in a 1:1 ratio (Fig. S41†). Despite the high selectivity of this reaction, the



Scheme 2 Activation of P_4 by β -diketiminate Mg^I complexes.

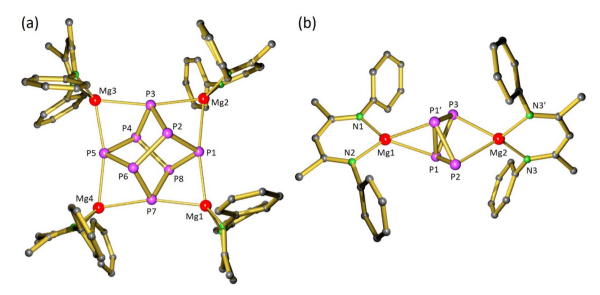


Fig. 1 (a) Crystal structure of [(BDI)Mg]₄(P₈) (1) in which iPr-groups and H atoms are omitted for clarity and a view of the Mg₄P₈ core. (b) Crystal structure of [(BDI*)Mg]₂(P₄) (2); the Et₂CH-groups and H atoms are omitted for clarity. A crystallographic mirror plane runs through the atoms Mg1, Mg2, P2, P3.

very good solubility induced by the Et_2CH -substituents allowed for isolation of crystalline $[(BDI^*)Mg]_2(P_4)$ (2) in only 34% yield.

In agreement with two triplet signals in $^{31}P\{^1H\}$ NMR, inspection of the crystal structure revealed a $P_4{}^{2-}$ dianion in a butterfly conformation which is bridging two (BDI*)Mg^+ units in an unusual η^2,η^2 -fashion. Butterfly-shaped $P_4{}^{2-}$ anions usually bridge metals in η^1,η^1 -fashion. 34 Recently, Aldridge and co-workers isolated an odd example of $P_4{}^{2-}$ bridging between Al and K in η^2,η^1 -fashion. While Hill and co-workers reported bridging in η^2,η^3 -fashion. 36 Complex 2 could also be considered to consist of a magnesate anion (BDI*)Mg(P_4)^- with two polar Mg–P bonds to the two-coordinate P atoms (P2, P3) of 2.573(4)–2.694(4) Å, charge-balanced by a (BDI*)Mg^+ cation that interacts with Mg–P interactions of circa 2.606(4) Å with both three-coordinate P-atoms (P1 and P1'). Due to disorder of the bridging $P_4{}^{2-}$ anion (Fig. S62†) a more accurate discussion of the crystal structure is not possible.

P₄ activation with Ca^I synthons

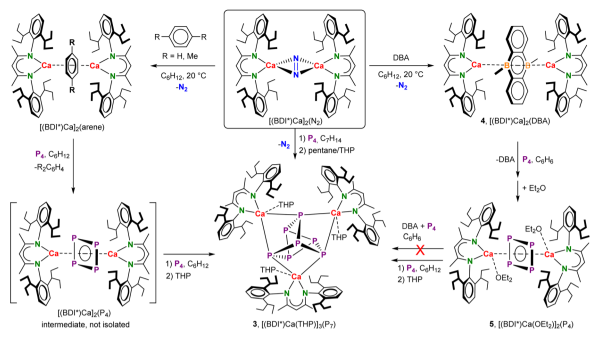
Although β-diketiminate stabilized Ca^{I} complexes are currently unknown, we reported a range of (BDI*)Ca–X–Ca(BDI*) complexes with various bridging X^{2-} anions that react like a low-valent Ca^{I} complex ($X = N_2$, benzene, p-xylene). Their reducing ability is directly related to the 2e-oxidation of the bridging anion, $X^{2-} \rightarrow X + 2e^{-}$, and therefore to the reduction potential of X.

Since the most reducing Ca^I synthon $[(BDI^*)Ca]_2(N_2)$ is unstable in aromatic solvents, 22 the reaction with P_4 was carried out in methylcyclohexane (Scheme 3). Reaction of the Ca^I synthon with a methylcyclohexane solution of P_4 at room temperature led to immediate N_2 evolution and a colour change from dark brown to dark orange. Analysis of the crude reaction mixture by $^{31}P\{^1H\}$ NMR spectroscopy revealed after 30 minutes a sharp low-field singlet at 458.4 ppm and a rather broad high-

field singlet at -85.3 ppm in a ratio of 0.09:0.91, respectively (Fig. S42†). Continued stirring for another 4 hours showed the selective formation of one species corresponding to the broad high-field signal at -85.3 ppm (Fig. S43†). Layering a saturated methylcyclohexane/pentane solution with a few drops of tetrahydropyran (THP) resulted in the isolation of yellow crystals of composition [(BDI*)Ca(THP)]₃(P₇) (3) in a 28% yield.

The crystal structure of 3 (Fig. 2a) revealed that P_4 had been reduced to the polyphosphide P_7^{3-} Zintl ion, encapsulated by three $[(BDI^*)Ca(THP)]^+$ fragments. A similar product, $[(BDI)Ca]_3(P_7)$, has been obtained by Roesky and co-workers by reduction of P_4 with $[(BDI)Ca(\mu_2 - H)]_2$. However, in contrast to this previous report which describes a major side-product with a ^{31}P -resonance at -241.3 ppm, the reduction of P_4 with the Ca^I synthon is highly selective.

Although the structure of 3 is close to being C_3 -symmetric, the complex does not show crystallographic symmetry. The Ca metal centres have shortest interactions to the three twocoordinate P atoms (P4-P6) in P₇³⁻ which formally carry a negative charge (Ca-P: 3.0557(7)-3.1163(6) Å). These bonds are considerably longer than the corresponding Ca-P bonds in $[(BDI)Ca]_3(P_7)$ which vary from 2.8667(9) to 2.9346(9) Å.¹⁹ The long Ca-P contacts in 3 are mainly due to the bulkier BDI* ligand and the additional THP coordination. While in 3 there are no Ca-P contacts to the apical P atom P7, the formally neutral three-coordinate P atoms in the P₃-triangle (P1-P3) show long contacts (Ca-P: 3.3253(7)-3.4611(5) Å). The P-P bond lengths in the P_7 -cage (2.1601(6)–2.2610(4) Å) are in the range of previous reported Zintl ions.37 Reports of group 2 metal based Zintl ions are particularly rare and especially their selective formation from P4 remains difficult. Hill and co-workers isolated a Mg flanked P7 Zintl cluster that is structurally very similar to [(BDI)Ca]₃(P₇). However, this product could only be



Scheme 3 Activation of P₄ by Ca¹ synthons.

obtained in poor yields as a minor side product by fractional crystallization of the raw product.³⁸

Whilst the ³¹P{¹H} NMR spectrum of [(BDI*)Ca(THP)]₃(P₇) at room temperature showed one very broad singlet at -85.28 ppm due to fast exchange between the three different P-positions in the P_7^{3-} anion, at $-90\,^{\circ}\text{C}$ this resonance was split into seven broad but distinct signals (Fig. S22†), partially with visible magnetic coupling (ppm: -147.6, -130.4, -111.6, -75.7, -69.2, -45.1, and -38.2). This behaviour differs from other reports on the dynamics of P_7^{3-} which generally describe splitting in only three signals in a ratio of 1:3:3 upon cooling. ^{19,37} These can be assigned to the apical position, the P_3 basal triangle and the three connecting P atoms. Our observed splitting into seven signals at low temperature can only be rationalized by a loss of trigonal symmetry by different coordination geometries at the Ca centres, making each P atom magnetically inequivalent.

The central 2e-donor X in (BDI*)Ca-X-Ca(BDI*) influences the synthon's stability and reactivity.22-24 In order to evaluate the effect of the central X²⁻ anion on P₄ reduction, the most reducing Ca^I synthon, [(BDI*)Ca]₂(N₂) was converted to the corresponding arene complexes by reaction with benzene or pxylene which led to N₂ release. These binuclear arene complexes were further reacted with equimolar quantities of P4 in cyclohexane- d_{12} and conversion was monitored with $^{31}P\{^{1}H\}$ NMR. Similarly to P_4 reduction with N_2^{2-} , two signals were observed: a sharp low-field resonance at 458.4 ppm and a rather broad high-field singlet at -85.3 ppm, corresponding to the P_7^{3-} Zintl anion. Over time, the sharp low-field resonance at 458.4 ppm disappeared and clean formation of the P7 complex was observed. This suggests that the species with a ³¹P resonance at 458.4 ppm is an intermediate on the way to formation of the P₇ complex. The only difference is the rate at which this happens

which seems to be related to the reducing power of the central X^{2-} anion: $N_2^{2-} > p$ -xylene $^{2-} >$ benzene $^{2-}$. While in the case of the N_2 complex, the intermediate species was nearly fully converted to the P_7 complex after an hour, the less reducing benzene complex required stirring overnight (Fig. S45†). As we were not able to isolate the intermediate, we sought a weaker Ca^I synthon to slow the reaction further.

Reaction of P₄ with (BDI*)Ca-(anthracene)-Ca(BDI*)²³ gave a myriad of products of which one could be recognized as the P₇ complex (3) by ³¹P NMR analysis (Fig. S48†). This prompted us to look further for a suitable bridging ligand. Due to aromaticity in its dianionic state, the boron-doped 9,10-dimethyldiboraanthracene (DBA) dianion is less reducing and has markedly different electronic properties.39-41 Recently, first lanthanide triple-decker complexes featuring DBA²⁻ ligands were reported. 42 Reduction of DBA with the Ca^I synthon [(BDI*) Ca](N₂) in cyclohexane resulted in gas evolution and immediate precipitation of a microcrystalline orange solid (Scheme 3). ¹H NMR showed selective formation of the target complex [(BDI*) Ca]₂(DBA) (4) which could be isolated in 88% yield (Fig. S49†). Unlike (BDI*)Ca-(C₆H₆)-Ca(BDI*), which in benzene shows reductive benzene coupling $(C_6H_6^{2-} + C_6H_6 \rightarrow biphenyl^{2-})^{43}$ [(BDI*)Ca]₂(DBA) is even at 60 °C remarkably stable in aromatic solvents. Its 11B NMR spectrum did not show a clear signal, even when quartz NMR tubes were used. Bright orange crystals suitable for single crystal X-ray diffraction were obtained by recrystallization from a cyclohexane/n-pentane mixture at room temperature.

The complex crystallized in the $P\bar{1}$ space group with two independent [(BDI*)Ca]₂(DBA) (4) inverse sandwich complexes in the asymmetric unit (Fig. 2b). The structure shows a planar $(\mu^2-\eta^6:\eta^6-DBA)^{2-}$ ligand of which the central B₂C₄-ring is sandwiched between the two (BDI*)Ca⁺ fragments with Ca-

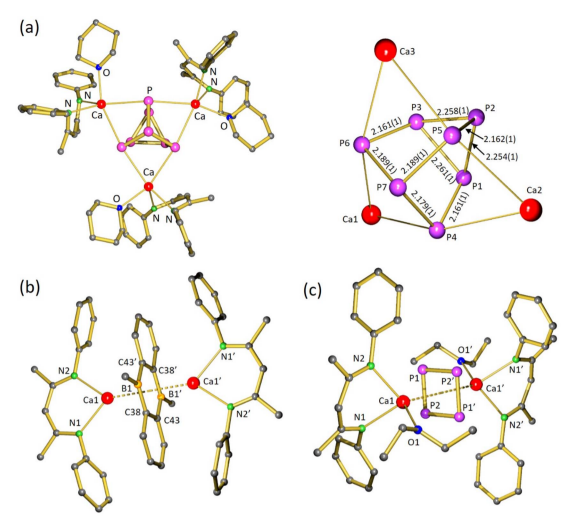


Fig. 2 (a) Crystal structure of $[(BDI^*)Ca(THP)]_3(P_7)$ (3) and a view of the Ca_3P_7 core. (b) Centrosymmetric crystal structure of $[(BDI^*)Ca]_2(DBA)$ (4). (c) Centrosymmetric crystal structure of $[(BDI^*)Ca(OEt_2)]_2(cyclo-P_4)$ (5). In all figures the H atoms and Et_2CH -groups have been omitted for clarity.

ring_{centre} distances ranging from 2.3552(4) to 2.3786(4) Å. Preference for metal coordination to the central B_2C_4 -ring was also observed in alkali metal complexes: $M(THF)_n$ –(DBA)– $M(THF)_n$ complexes (M = Li, Na, K).^{40,44} Like in these alkali metal inverse sandwiches, the planar DBA²⁻ anion in 4 is isoelectronic to anthracene and shows in its central ring C–C bond distances of 1.466(3)–1.468(2) Å and B–C bond distances of 1.530(3)–1.541(3) Å, specifying the extended aromatic nature of this dianion. Computational investigation of 4 shows that Ca ligand bonding is highly ionic. NPA charges: (BDI*) –0.88, Ca +1.79, DBA –1.87 (Fig. S67†).

A C_6D_6 solution of [(BDI*)Ca]₂(DBA) (4) and one equivalent of P_4 was stirred overnight at room temperature resulting in a colour change from orange to bright yellow. The $^{31}P\{^1H\}$ NMR spectrum showed a sharp singlet at 453.9 ppm indicative for exclusive formation of the hitherto unidentified intermediate (Fig. S51†) whereas the high field signal at -85 ppm for P_7^{3-} formation is missing. In addition, 1H , ^{11}B , and ^{13}C NMR confirmed the release of neutral DBA (Fig. S52–S54†). The sharp low-field singlet of this intermediate could be assigned to the

four chemically equivalent phosphorus atoms of the *cyclo-*P₄ dianion, in the form of an inverse sandwich complex. Comparing to few reports in literature, the ^{31}P NMR chemical shift of the $cyclo-P_4{}^{2-}$ ring is sensitive to its environment: $[(^{DIPP}Form)_2Sm]_2(cyclo-P_4)$ with $\delta=+453$ ppm 45 ($^{DIPP}Form=HC(N-DIPP)_2), Cs_2P_4\cdot 2NH_3$ with $\delta=+348$ ppm, 46 [(NON) Sm(THF)_2]_2(cyclo-P_4) with $\delta=+480$ ppm (NON = 4,5-bis(2,6-diisopropylphenyl-anilido)-2,7-di-*tert*-butyl-9,9-dimethyl-xanthene), 47 [(NON)Yb(THF)_2]_2(cyclo-P_4) with $\delta=+382$ ppm, 47 and a U complex with η^2,η^2 -bridging $cyclo-P_4{}^{2-}$ with $\delta=+718$ ppm. 48 Despite this considerable range in chemical shifts, all values are considerably downfield shifted, substantiating the 6π -electron aromatic character of this dianion.

Crystallization from a saturated pentane solution, layered with drops of Et₂O, allowed for isolation and structural characterization of [(BDI*)Ca(OEt₂)]₂(cyclo-P₄) (5) by X-ray diffraction. The crystal structure of 5 confirms the formation of an inverse sandwich complex with a bridging cyclo-P₄ dianion (Fig. 2c). The 6π -electron aromatic cyclo-P₄²⁻ ring bridges in η^4 , η^4 -fashion between two cationic (BDI*)Ca⁺ fragments with

Ca-P₄(centroid) bond lengths of 2.6460(5) Å and Ca-P bond lengths between 3.0446(9) and 3.0682(8) Å. The P₄ ring is disordered over two positions which are rotated in respect to each other around the Ca···Ca' axis by circa 45°. The P-P bond lengths range from 2.156(1) Å to 2.158(1) Å and internal P-P-P angles are close to 90° (P2-P1-P2' 89.40(4)°, P1'-P2-P1 90.60(4)°). Hence, the aromatic P_4^{2-} ring is almost perfectly square planar. The P-P bond lengths of the phosphorus ring are in the range of reported examples and seems independent of the metal atoms which sandwiches this ring (Sm: 2.144(1)-2.162(1) Å, 45 U: 2.149(2)-2.152(2) Å, 48 Cs: 2.146(1)-2.148(1) Å). 46 It is of interest to note that the geometry of P₄²⁻ depends strongly on the metals that sandwich this dianion. The ring structure of 6π -electron aromatic cyclo- $P_4^{\ 2-}$ in the Ca inverse sandwich 5 contrasts strongly with the butterfly structure of P₄²⁻ in the Mg inverse sandwich 2. A detailed discussion on these differences follows below (vide infra).

It is noteworthy, that the reduction of P_4 with [(BDI*) $Ca]_2(DBA)$ (4) exclusively led to formation of the aromatic (*cyclo*- P_4)²⁻ dianion. Even with a large excess of P_4 and using forcing reaction conditions (60 °C), only the P_4 complex 5 was formed. This represents the first example of a quantitative reduction of P_4 to (*cyclo*- P_4)²⁻ mediated by s-block metals. Treatment of P_4 with Cs in THF followed by solvation in liquid ammonia led to the formation of $Cs_3P_7 \cdot 3NH_3$ as major product and the desired cyclotetraphosphide $Cs_2P_4 \cdot 2NH_3$ was only a by-product.⁴⁶

In order to evaluate its role as an intermediate in polyphosphide formation, crystalline $[(BDI^*)Ca(OEt_2)]_2(cyclo-P_4)$ (5) and one equivalent of P_4 were suspended in cyclohexane- d_{12} (Scheme 3). Monitoring the reaction with $^{31}P\{^1H\}$ NMR showed after 2 hours at room temperature slow conversion of the $(cyclo-P_4)^{2-}$ complex into the $(P_7)^{3-}$ complex $(P_4:P_7=0.63:0.37)$. Complete, selective conversion to the P_7 product was achieved after two days at room temperature (Fig. S55†).

Two observations need further attention. (1) The DBA complex 4 cannot be converted to a P_7 complex and reacts with excess white phosphorous only to the P_4 product. (2) Once crystallized in presence of ether and isolated, 5 selectively reacts with P_4 to the P_7 complex. Although at first sight contradicting, these combined observations can only lead to one conclusion. The neutral DBA that is released in reaction of 4 with P_4 must be an inhibitor for the P_4 -to- P_7 conversion.

Indeed, whereas the reaction of crystalline [(BDI*) $Ca(OEt_2)]_2(cyclo-P_4)$ (5) with P_4 in cyclohexane- d_{12} showed slow but selective conversion to the P_7 complex, addition of DBA to 5 inhibited P_4 -to- P_7 conversion. With catalytic quantities of DBA as low as 5 mol% no conversion was observed, even after one day at room temperature. However, after 30 hours at 60 °C a small amount of the P_7 complex was observed (Fig. S58†). The mechanism of this inhibitor effect are still unclear but leave room for speculation (*vide infra*).

P_4^{2-} : butterfly or aromatic ring?

The geometry of P_4^{2-} depends strongly on the metals that sandwich this dianion. Captured between two (BDI*)Mg⁺ cations it takes the form of a butterfly with η^2, η^2 -bridging (2)

but between (BDI*)Ca⁺ a η^4 , η^4 -bridging 6π -electron aromatic *cyclo*-P₄²⁻ dianion (5) is favoured. These intriguing differences in geometry and coordination modes may be understood by Density Functional Theory (DFT) calculations.

The structures of 2 and ether-free 5 were optimized at the PBE0/def2-SVP level of theory. The calculated structure of 2 fits reasonably well with that from the crystal structure (Fig. S66†), indicating a sufficient level of theory. The aromaticity of 5 has been investigated before, and differs from that in classical aromatic hydrocarbons like in benzene.49 It can be noticed that the P-P bonds in *cyclo*- P_4^{2-} in 5 (calculated: 2.173–2.184 Å) are not much different from the single P-P bonds in P₄ (calculated: 2.201 Å). Electron Localization Function (ELF) analyses showed that the P-P bonds in cyclo-P₄²⁻ have P-P single bond character and show no sign of the high electron delocalization as found in benzene.49 In contrast to the high population in C-C bonds of benzene, ELF analysis does not show a high population in the P-P bonding orbitals. Instead, a high population was found in the P lone-pairs in cyclo-P₄²⁻. Hence, the concept of "lone-pair aromaticity" was defined. In contrast to benzene, for which a circle within the C₆-ring depicts aromaticity, for cyclo-P₄²⁻ a circle around the P₄-ring has been proposed.⁴⁹

The question remains why the P₄²⁻ dianion can have two different appearances, butterfly or cyclic structure, depending on the metal that sandwich this entity. The answer may lie in the character of metal-P₄ bonding. Based on electronegativity differences (Mg: 1.31, Ca: 1.00, P: 2.19),50 Mg-P4 bonding in 2 should be slightly more covalent than Ca-P₄ bonding in 5. The rather low Wiberg Bond Indices (WBI's) for Mg-P4 bonding (0.11/0.17) and Ca-P₄ bonding (0.05/0.06) show that both bonds have ionic character but the Mg-P₄ bonds are somewhat more covalent (Fig. 3a). Natural Population Analysis (NPA) indeed shows a significantly higher negative charge on the cyclo-P₄ unit in Ca complex 5 (-1.74) than that on butterfly P₄ unit in Mg complex 2 (-1.56); see Fig. 3a. This corresponds to higher positive charges on the Ca cations (+1.74) compared to the Mg cations (average: +1.67). The butterfly form of P₄²⁻ shows two different P atoms. The two-coordinate P atoms that carry most of the negative charge (-0.55) shows strong bonding to Mg and feature WBI's of 0.17. The three-coordinate P atoms with charges of -0.23 have weaker bonds to Mg (WBI: 0.11). This confirms the view that 2 can be seen as a magnesate anion (BDI*)Mg(P₄)[−] with a total NPA charge of −0.82 that interacts with a (BDI*)Mg⁺ cation carrying a charge of +0.82. The WBI's for the P-P bonds (0.94/0.98) are close to those for a covalent single bond.

In comparison, the NPA charges on the P atoms in *cyclo*-P₄ are more similar (-0.38/-0.49) and intermediate to those in butterfly P₄²⁻ (-0.23/-0.55). Charge differences are likely dictated by their different environments. Atoms-In-Molecules (AIM) analyses show that the P₄²⁻ dianions in 2 and 5 are also involved in weak P···H–C bonding with organic fragments of the BDI* ligands (Fig. S72 and S74†). In agreement with more ionic character of the Ca complex, the WBI's for Ca–P₄ bonding (0.05/0.06) are smaller than for Mg–P₄ bonding (0.11/0.17). However, the WBI's for the P–P bonds in *cyclo*-P₄ (1.23/1.25)

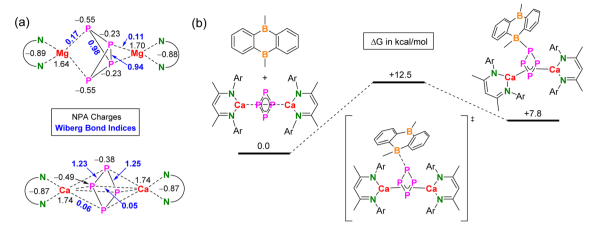
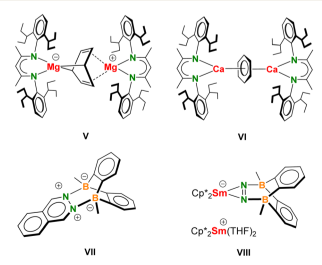


Fig. 3 Computational studies at the PBE0-D3BJ(PCM = cyclohexane)/def2-TZVP level of theory. (a) NPA charges and WBI's for $[(BDI*)Mg]_2(P_4)$ and $[(BDI*)Ca]_2(cyclo-P_4)$. (b) Energy profile for the reaction of $[(BDI*)Ca]_2(cyclo-P_4)$ with DBA.

are slightly larger than that expected for a single bond. This is in agreement with some extent of aromaticity (*vide supra*).

The differences observed in bonding of the P₄²⁻ dianion in Mg and Ca complexes are comparable to differences observed in bonding of the $C_6H_6^{2-}$ dianion (V and VI, Scheme 4). Bonding in the Mg complex (BDI*)Mg-(C₆H₆)-Mg(BDI*) is more covalent than that in the corresponding Ca complex. The $C_6H_6^{2-}$ in the Mg complex shows the typical boat form with strong localized Mg-C bonding to bow and stern and much weaker Mg-C interactions to the C=C bonds.33 Similarly as for 2, the complex can be seen as a magnesate anion (BDI*)Mg(C₆H₆)⁻ interacting with a (BDI*)Mg+ cation through unusual Mg-alkene coordination for which we recently found ample proof. 51-53 In contrast, (BDI*)Ca-(C₆H₆)-Ca(BDI*) shows a nearly flat C₆H₆²⁻ dianion and η^6, η^6 -bridging between the Ca²⁺ ions.²² These parallels between $C_6H_6^{\,2-}$ and $P_4^{\,2-}$ bonding in Mg or Ca sandwich complexes find their origins in the more covalent nature of the Mg-ligand bond but could also be related to the considerably larger size of the Ca²⁺ cation compared to the Mg²⁺ cation. This explanation is in agreement with the occurrence of P₄²-



Scheme 4 Formulas V-VIII.

butterfly structures in Al, Ga, Si or Ni complexes which are even more covalent in character than the Mg complex. 12,14,34-36

How does diboraanthracene (DBA) inhibit P_4 to P_7 conversion?

Whilst [(BDI*)Ca(OEt₂)]₂(cyclo-P₄) (5) reacts selectively with P₄ to the P₇ complex, addition of DBA inhibited this conversion. Initially, we sought an explanation in possible interaction of DBA with the P₄ reactant. It is known that the electrophilic diborane DBA can interact with bidentate electron-rich ligands like 1,2-diazines (e.g. in VII)⁵⁴ and even assists in N₂ fixation with Cp*₂Sm (VIII) (Cp* = 1,2,3,4,5-penta-methyl-cyclopentadienyl).⁵⁵ However, ¹H and ³¹P NMR spectra of a mixture of DBA and P₄ did not show any changes in the chemical shifts when compared to the pure species. This excludes significant DBA···P₄ interaction. Attempts to optimize the structure of potential DBA···P₄ complexes by DFT calculation also only led to separation of these molecules.

Alternatively, DBA can interact with the P_4 complex [(BDI*) $Ca]_2(cyclo-P_4)$, thus inhibiting further reactivity with P_4 . DFT calculations indeed show that this complex is able to interact with DBA by formation of a P–B bond (Fig. 3b; for structures see Fig. S78†). Although the activation free energy for this process is only $\Delta G_{298}^{\dagger} = 12.5$ kcal mol^{-1} , the adduct is $\Delta G_{298} = 7.8$ kcal mol^{-1} higher in energy than the unbound molecules. This implies a fast equilibrium that lies mainly on the side of free P_4 and [(BDI*)Ca] $_2(cyclo-P_4)$. This means that, although complexation of DBA with $cyclo-P_4^{2-}$ is possible, this cannot be a reason to inhibit further reactivity towards the formation of P_7 complex. Especially, when one considers that also small, catalytic quantities of DBA already work as inhibitor.

Reaction mechanisms that give rise to higher nuclearity clusters are in general poorly understood.³⁷ However, it seems reasonable to assume that reduction of P_4 with s-block metal reducing agents starts with electron-transfer. As we could show that the P_4^{2-} dianion is a likely intermediate en route to P_7^{3-} , the first step of this transformation could be a single electron transfer (SET) process. As DBA is a molecule with a low-lying

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LUMO and can be easily reduced, its mode of inhibiting the P₄to-P₇ conversion may simply be a reversible electron-capture process. The high affinity of DBA for electrons inhibits SET to P₄. Using DFT, we found that reduction of DBA is indeed much more facile than reduction of P_4 . The reaction, DBA⁻ + $P_4 \rightarrow P_4$ + DBA, was calculated to be highly endothermic (in cyclohexane: $\Delta H = +23.5 \text{ kcal mol}^{-1}$). This is also in agreement with the observation that catalytic quantities of DBA function as a stabilizer for the P₄²⁻ dianion and would explain the highly selective formation of the P_4 complex 5.

Conclusion

We achieved first P₄ reductions with low-valent β-diketiminate Mg^I complexes and found the selectivity to be greatly dependent on the bulk of the ligand. Using a superbulky BDI ligand with DIPeP substituents (BDI*) led to highly selective formation of a butterfly-shaped P₄²⁻ dianion bridging two (BDI*)Mg⁺ fragments in a unique η^2 , η^2 -fashion. The same complex with Ca instead of Mg features a cyclo-P₄²⁻ dianion bridging (BDI*)Ca⁺ fragments in an η^4 , η^4 mode. This difference finds its origin in the more covalent nature of the Mg-(P₄) bond but could also be related to the considerably larger size of the Ca²⁺ cation compared to the Mg²⁺ cation, favouring a delocalized cyclo-P₄²⁻ dianion.

Reaction of low-valent Ca^I synthons selectively gave products with the P_7^{3-} Zintl anion: $[(BDI^*)Ca]_3(P_7)$. Monitoring these reactions by NMR shows first unambiguous proof that P₇³⁻ formation proceeds through a cyclo-P42- intermediate. The kinetics of the $P_4 \rightarrow cyclo$ - $P_4^{2-} \rightarrow P_7^{3-}$ conversion depends strongly on the reducing power of the Ca^I synthon: [(BDI*) $Ca]_2(X)$ in which X is dianionic N_2^{2-} or arene²⁻. Conversion is faster along the row $X = benzene^{2-} < p$ -xylene²⁻ $< N_2^{2-}$.

case of Ca^I synthons, using 9,10-dimethyldiboraanthracene (DBA) as a bridging dianion led to exclusive formation of the cyclo-P₄²⁻ product. Even with excess P₄ and forcing reaction conditions no further $P_4 \rightarrow P_7$ conversion was observed in the time frame conducted. As addition of trace quantities of DBA already inhibited and considerably retarded further reactivity of the cyclo-P₄²⁻ dianion, it is proposed that DBA prevents radical reactivity by functioning as a reversible electron trap.

These first investigations on P₄ reduction with low-valent Ae metal complexes show that selectivities depend on the bulk of the BDI ligand, the metal and the presence of inhibitors for radical reactivity. We continue our research with investigations on P4 reduction with heavier low-valent Ae metal synthons.

Data availability

Crystallographic data has been deposited with the Cambridge Structural Database.

Author contributions

S. Thum: conceptualization, investigation, validation, formal analysis, writing - original draft, visualization. O. P. E. Townrow: investigation, validation, formal Analysis. J. Langer: formal analysis, validation. Sjoerd Harder: conceptualization, writing - original draft - review and editing, visualization, validation, supervision, project administration.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 U. Lennert, P. B. Arockiam, V. Streitferdt, D. J. Scott, C. Rödl, R. M. Gschwind and R. Wolf, Nat. Catal., 2019, 2, 1101-1106.
- 2 D. J. Scott, Angew. Chem., Int. Ed., 2022, 61, e202205019.
- 3 C. M. Hoidn, D. J. Scott and R. Wolf, Chem.-Eur. J., 2021, 27, 1886-1902.
- 4 B. M. Cossairt, N. A. Piro and C. C. Cummins, Chem. Rev., 2010, 110, 4164-4177.
- 5 M. Caporali, L. Gonsalvi, A. Rossin and M. Peruzzini, Chem. Rev., 2010, 110, 4178-4235.
- 6 L. Giusti, V. R. Landaeta, M. Vanni, J. A. Kelly, R. Wolf and M. Caporali, Coord. Chem. Rev., 2021, 441, 213927.
- 7 M. Scheer, G. Balázs and A. Seitz, Chem. Rev., 2010, 110, 4236-4256.
- 8 S. Khan, S. S. Sen and H. W. Roesky, Chem. Commun., 2012, 48, 2169.
- 9 N. A. Giffin and J. D. Masuda, Coord. Chem. Rev., 2011, 255, 1342-1359.
- 10 W. Uhl and M. Benter, Chem. Commun., 1999, 771-772.
- 11 C. Dohmeier, H. Schnöckel, C. Robl, U. Schneider and R. Ahlrichs, Angew. Chem., Int. Ed., 1994, 33, 199-200.
- 12 G. Prabusankar, A. Doddi, C. Gemel, M. Winter and R. A. Fischer, Inorg. Chem., 2010, 49, 7976-7980.
- 13 J. D. Masuda, W. W. Schoeller, B. Donnadieu and G. Bertrand, Angew. Chem., Int. Ed., 2007, 46, 7052-7055.
- 14 Y. Xiong, S. Yao, M. Brym and M. Driess, Angew. Chem., Int. Ed., 2007, 46, 4511-4513.
- 15 C. Jones, Nat. Rev. Chem, 2017, 1, 0059.
- 16 B. Rösch and S. Harder, Chem. Commun., 2021, 57, 9354-9365.
- 17 L. A. Freeman, J. E. Walley and R. J. Gilliard, Nat. Synth., 2022, 1, 439-448.
- 18 S. P. Green, C. Jones and A. Stasch, Science, 2007, 318, 1754-1757.
- 19 R. Yadav, M. Weber, A. K. Singh, L. Münzfeld, J. Gramüller, R. M. Gschwind, M. Scheer and P. W. Roesky, Chem.-Eur. J., 2021, 27, 14128-14137.
- 20 M. M. Rauhut and A. M. Semsel, J. Org. Chem., 1963, 28, 471-473.

- 21 A. S. S. Wilson, C. Dinoi, M. S. Hill, M. F. Mahon, L. Maron and E. Richards, *Angew. Chem.*, *Int. Ed.*, 2020, **59**, 1232–1237.
- 22 B. Rösch, T. X. Gentner, J. Langer, C. Färber, J. Eyselein, L. Zhao, C. Ding, G. Frenking and S. Harder, *Science*, 2021, 371, 1125–1128.
- 23 J. Mai, B. Rösch, N. Patel, J. Langer and S. Harder, *Chem. Sci.*, 2023, **14**, 4724–4734.
- 24 J. Mai, B. Rösch, J. Langer, S. Grams, M. Morasch and S. Harder, *Eur. J. Inorg. Chem.*, 2023, **26**, e202300421.
- 25 A. M. Griffin, P. C. Minshall and G. M. Sheldrick, J. Chem. Soc., Chem. Commun., 1976, 809–810.
- 26 W. Huang and P. L. Diaconescu, Chem. Commun., 2012, 48, 2216–2218.
- 27 M. E. Barr, B. R. Adams, R. R. Weller and L. F. Dahl, *J. Am. Chem. Soc.*, 1991, 113, 3052–3060.
- 28 F. Spitzer, C. Graßl, G. Balázs, E. M. Zolnhofer, K. Meyer and M. Scheer, *Angew. Chem., Int. Ed.*, 2016, 55, 4340–4344.
- 29 S. N. Konchenko, N. A. Pushkarevsky, M. T. Gamer, R. Köppe, H. Schnöckel and P. W. Roesky, *J. Am. Chem. Soc.*, 2009, **131**, 5740–5741.
- 30 F. Hennersdorf, J. Frötschel and J. J. Weigand, *J. Am. Chem. Soc.*, 2017, **139**, 14592–14604.
- 31 C. Ganesamoorthy, C. Wölper, A. S. Nizovtsev and S. Schulz, *Angew. Chem., Int. Ed.*, 2016, 55, 4204–4209.
- 32 T. X. Gentner, B. Rösch, K. Thum, J. Langer, G. Ballmann, J. Pahl, W. A. Donaubauer, F. Hampel and S. Harder, *Organometallics*, 2019, **38**, 2485–2493.
- 33 T. X. Gentner, B. Rösch, G. Ballmann, J. Langer, H. Elsen and S. Harder, *Angew. Chem., Int. Ed.*, 2019, **58**, 607–611.
- 34 S. Pelties, A. W. Ehlers and R. Wolf, *Chem. Commun.*, 2016, 52, 6601–6604.
- 35 M. M. D. Roy, A. Heilmann, M. A. Ellwanger and S. Aldridge, *Angew. Chem., Int. Ed.*, 2021, **60**, 26550–26554.
- 36 M. S. Hill, M. F. Mahon, C. L. McMullin, S. E. Neale, K. G. Pearce and R. J. Schwamm, *Z. Anorg. Allg. Chem.*, 2022, 648, e202200224.
- 37 R. S. P. Turbervill and J. M. Goicoechea, *Chem. Rev.*, 2014, 114, 10807–10828.
- 38 M. Arrowsmith, M. S. Hill, A. L. Johnson, G. Kociok-Köhn and M. F. Mahon, *Angew. Chem., Int. Ed.*, 2015, **54**, 7882–7885.

- 39 M. Dietz, M. Arrowsmith, K. Drepper, A. Gärtner, I. Krummenacher, R. Bertermann, M. Finze and H. Braunschweig, J. Am. Chem. Soc., 2023, 145, 15001–15015.
- 40 E. Von Grotthuss, S. E. Prey, M. Bolte, H. W. Lerner and M. Wagner, *J. Am. Chem. Soc.*, 2019, 141, 6082–6091.
- 41 S. E. Prey and M. Wagner, *Adv. Synth. Catal.*, 2021, **363**, 2290–2309.
- 42 C. Uhlmann, L. Münzfeld, A. Hauser, T. T. Ruan, S. Kumar Kuppusamy, C. Jin, M. Ruben, K. Fink, E. Moreno-Pineda and P. W. Roesky, *Angew. Chem., Int. Ed.*, 2024, **63**, e202401372.
- 43 J. Mai, M. Morasch, D. Jędrzkiewicz, J. Langer, B. Rösch and S. Harder, *Angew. Chem., Int. Ed.*, 2023, **62**, e202212463.
- 44 E. von Grotthuss, S. E. Prey, M. Bolte, H. W. Lerner and M. Wagner, *Angew. Chem., Int. Ed.*, 2018, 57, 16491–16495.
- 45 C. Schoo, S. Bestgen, R. Köppe, S. N. Konchenko and P. W. Roesky, *Chem. Commun.*, 2018, **54**, 4770–4773.
- 46 F. Kraus, J. C. Aschenbrenner and N. Korber, *Angew. Chem.*, Int. Ed., 2003, 42, 4030–4033.
- 47 A. Hauser, L. Münzfeld, S. Schlittenhardt, R. Köppe, C. Uhlmann, U.-C. Rauska, M. Ruben and P. W. Roesky, *Chem. Sci.*, 2023, 14, 2149–2158.
- 48 A. S. P. Frey, F. G. N. Cloke, P. B. Hitchcock and J. C. Green, *New J. Chem.*, 2011, 35, 2022–2026.
- 49 F. Kraus and N. Korber, Chem.-Eur. J., 2005, 11, 5945-5959.
- 50 L. R. Murphy, T. L. Meek, A. Louis Allred and L. C. Alien, J. Phys. Chem. A, 2000, 104, 5867–5871.
- 51 J. Martin, J. Langer, M. Wiesinger, H. Elsen and S. Harder, Eur. J. Inorg. Chem., 2020, 2582–2595.
- 52 K. Thum, A. Friedrich, J. Pahl, H. Elsen, J. Langer and S. Harder, *Chem.-Eur. J.*, 2021, 27, 2513–2522.
- 53 K. Thum, J. Pahl, J. Eyselein, H. Elsen, J. Langer and S. Harder, *Chem. Commun.*, 2021, 57, 5278–5281.
- 54 S. N. Kessler and H. A. Wegner, *Org. Lett.*, 2010, **12**, 4062–4065.
- 55 S. Xu, L. A. Essex, J. Q. Nguyen, P. Farias, J. W. Ziller, W. H. Harman and W. J. Evans, *Dalton Trans.*, 2021, 50, 15000-15002.