Attenuated Organo-magnesium Activation of White Phosphorus

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Sequential reactions between a 2,6-di-iso-propylphenyl-substituted \( \beta \)-diketiminato magnesium n-butyl derivative and \( \text{P}_4 \) allow the highly discriminating synthesis of unusual \([\text{nBu}_2\text{P}_4]^{2-}\) and \([\text{nBu}_2\text{P}_8]^{2-}\) cluster dianions.

Keywords: cluster compounds • magnesium • \( \text{P}_4 \) activation • phosphorus • structure elucidation

White phosphorus is the starting point in the synthesis of many organophosphanes and phosphorus-containing molecules. As a consequence, the controlled activation of \( \text{P}_4 \) has attracted considerable attention.\(^{[1]}\) While notable metal-free examples of \( \text{P}_4 \) activation are provided by Bertrand’s use of nucleophilic \( \text{N} \)-heterocyclic carbenes for sequential P-P activation of the tetrahedral molecule,\(^{[2]}\) a majority of approaches have utilized either reactive transition metal or main group reagents. Typical transition metal-based activation can induce not only the cleavage of bonds within \( \text{P}_4 \) but also the aggregation of smaller fragments to form \( \text{P}_n \) (\( n>4 \)) cluster anions. Cummins, for example, has shown that disproportionation of a niobium(IV) tris-enolate in the presence of \( \text{P}_4 \) provides the \( \{\text{P}_8\} \) cluster compound \( 1^{,[3]} \) while in a very recent report, Bergman and Arnold have described a trinuclear \( \beta \)-diketiminato \( \mu-\text{P}_{12} \) niobium complex, which was isolated as one of multiple minor by-products during the reaction of the Nb(V) complex \( (\text{DippBDI})\text{Nb}(\text{N}^\text{Bu})\text{Me}_2 \) \( (\text{DippBDI} = \text{HC}\{\text{C}(\text{Me})_2\text{N}(2,6-i-\text{Pr}_2\text{C}_6\text{H}_3)\}_2) \) with \( \text{H}_2 \) in the presence of \( \text{P}_4 \).\(^{[4]}\) Notably, this latter reaction was reported to yield a compound containing a cyclo-\( \text{P}_4 \) dianion as its predominant reaction product and, in some respects, the irrationality of the formation both of these compounds is typical of the synthesis of high nuclearity \( \{\text{P}_n\} \) clusters.\(^{[1f]}\)
Although reductive chemistry to yield cyclo-P₄ anions has also been described through the formal oxidative addition of P₄ to low oxidation state Group 13 and 14 derivatives (for example, compounds 2 and 3)\[1c,5,6]\ and there is precedent for the activation of P₄ through its interaction with transition metal hydrides and alkyls,\[7\] well defined reactions with anionic Group 14 derivatives of elements from Groups 1 and 2 are uncommon and dominated by the use of highly sterically demanding carbon- or silicon-reagents. Wright and co-workers have described the synthesis of the [R₂P₈]₂⁻ cluster dianion (4) derived from a {P₇}³⁻ Zintl ion core through the reaction of the very bulky hypersilyl potassium reagent with P₄.\[8\] The disubstituted butterfly compound 5 (Mes⁺ = 2,4,6-tBu₃C₆H₂) and the diphosphene Mes⁺P=PMes⁺ were similarly isolated from the reaction of Mes⁺Li and P₄ in the presence of Mes⁺Br.\[9\] Production of compound 5 presumably occurs through initial addition of Mes⁺Li across a P-P bond within the P₄ unit and the intermedicy of the consequent organopolyporphosphorus anion. The existence of species of this latter type has very recently been confirmed by Lammertsma and co-workers who reported that the B(C₆F₅)₃-stabilized lithium bicyclo[1.1.0]tetraphosphabutane species 6a and 6b could be isolated from reactions of P₄ and sterically demanding lithium aryls in the presence of the borane Lewis acid.\[10\]

Although similar but indiscriminate reactivity was posited some 50 years ago during reactions with simple Grignard reagents,\[11\] the activation of white phosphorus by nucleophilic Group 2 reagents is even less widespread as that by the alkali metals. Our own research interests lie in the development of a defined bond activation and homogeneous catalytic chemistry for the Earth-abundant and environmentally benign alkaline earth
As a first step toward the incorporation of white phosphorus in Group 2-derived catalysis, we present in this contribution the results of our initial studies which indicate that remarkable kinetic control during P₄ activation is achievable with organomagnesium reagents (Scheme 1). Key to the success of this chemistry is the application of the sterically demanding DippBDI supporting ligand employed in the syntheses of Arnold and Bergman’s aforementioned niobium systems, compounds 2 and 3 and very recently described binuclear cobalt(I) and copper(I) species containing a P₄-derived tetraphosphacyclobutadiene and an unactivated P₄ ligand respectively.[13]

![Scheme 1: Synthesis of the complexes described herein.](image)

An initial reaction between an equimolar quantity of P₄ and the β-diketiminato n-butylmagnesium complex, [(DippBDI)Mg-nBu] (7), resulted in the smooth consumption of the magnesium starting material to provide a single new species (8) characterized by two mutually coupled doublet resonances at δ –25.6 and –195.5 ppm (J_P-P = 99.6 Hz) in the resultant ³¹P{¹H} NMR spectrum. Notably, although compound 8 formed with only 50% consumption of the P₄, some evidence of further onward reaction was apparent on continued monitoring (vide infra). Repetition of this reaction with the reagent quantities adjusted to reflect the apparent 2:1 stoichiometry resulted in the complete consumption of both P₄ and 7 and the sole production of compound 8, which was also observed to comprise a single set of temperature invariant (from –90 to +90 °C) n-butyl and (DippBDI) ligand environments in the corresponding ¹H and ¹³C NMR spectra. The origin of these observations was resolved through the isolation of colorless single crystals of compound 8 suitable for X-ray diffraction analysis from n-hexane solution at low temperature, the results of which are displayed in
Figure 1 with selected bond length and angle data presented in the figure caption. The molecular structure of 8 comprises a dinuclear magnesium complex of an unprecedented non-planar \([n\text{Bu}_2\text{P}_4]^{2-}\) dianion which may be considered as arising from the formal addition of two magnesium \(n\)-butyl fragments to, and the consequent activation of two P-P single bonds within, the P₄ tetrahedron. The P-C bonds to the \(n\)-butyl chains are located on the adjacent P(1) and P(2) atoms while the magnesium centers each coordinate to one of these alkylated phosphorus atoms and the phosphorus atom located at the opposite diagonal corners of the tetraphosphorus unit. The adoption of a square array is prevented by a distinct pyramidalization at each phosphorus center such that the P-P-P angles are compressed to ca. 80°. The P(3)-P(4) bond length \([2.2647(7) \text{ Å}]\) is significantly elongated in comparison to the interatomic separation between the alkylated phosphorus atoms \([P(1)-P(2) 2.1762(7) \text{ Å}]\) but is comparable to the P-P distances observed in molecules containing the \([P_4]^{4-}\) anion such as compound 2 \([\text{mean } 2.29 \text{ Å}]\).\(^5\) The Mg-P distances to P(3) and P(4) \([\text{Mg}(1)-P(3) 2.5625(7), \text{Mg}(2)-P(4) 2.5592(7) \text{ Å}]\) are also shorter than those to P(1) and P(2) \([\text{Mg}(1)-P(1) 2.6336(7), \text{Mg}(2)-P(2) 2.6652(7) \text{ Å}]\) and we interpret both of these structural features to indicate that the distribution of negative charge is polarized toward the non-alkylated P(3) and P(4) centers and that the \([n\text{Bu}_2\text{P}_4]^{2-}\) unit as a whole may be considered as derived from the formal addition of two \(n\)-butyl carbenium ions to the \([P_4]^{4-}\) constituent anion of compound 2.
Figure 1: ORTEP representation (30% probability ellipsoids) of compound 8. Hydrogen atoms and iso-propyl groups are removed for clarity. Selected bond lengths (Å) and angles (°): Mg(1)-N(1) 2.0462(15), Mg(1)-N(2) 2.0465(15), Mg(1)-P(3) 2.5625(7), Mg(1)-P(1) 2.6336(7), Mg(1)-P(4) 3.1607(7), Mg(2)-N(4) 2.0424(15), Mg(2)-N(3) 2.0466(15), Mg(2)-P(4) 2.5592(7), Mg(2)-P(2) 2.6652(7), P(1)-C(59) 1.8584(19), P(1)-P(2) 2.1762(7), P(1)-P(4) 2.2127(7), P(2)-C(63) 1.8575(19), P(2)-P(3) 2.2108(7), P(3)-P(4) 2.2647(7), N(1)-Mg(1)-P(3) 126.57(5), N(2)-Mg(1)-P(3) 122.72(5), N(1)-Mg(1)-P(1) 127.76(5), N(2)-Mg(1)-P(1) 122.16(5), P(3)-Mg(1)-P(4) 57.14(2), N(1)-Mg(1)-P(4) 167.48(5), N(2)-Mg(1)-P(4) 99.06(5), P(3)-Mg(1)-P(4) 45.131(17), P(1)-Mg(1)-P(4) 43.731(16), N(4)-Mg(2)-N(3) 93.52(6), N(4)-Mg(2)-P(4) 126.28(5) N(3)-Mg(2)-P(4) 121.83(5), N(4)-Mg(2)-P(2) 121.40(5), N(3)-Mg(2)-P(2) 130.79(5), P(4)-Mg(2)-P(2) 66.130(19).

The previously noted onward reaction of compound 8 and additional P₄ (vide supra) prompted us to carry out a further reaction of an equimolar quantity of P₄ and compound 8. Reaction in benzene solution at room temperature provided the slow and simultaneous consumption of the starting materials and direct conversion to a single new compound (9) characterized by the emergence of eight mutually coupled signals in the ³¹P NMR spectrum. The corresponding ¹H and ¹³C NMR spectra indicated that this new compound comprised two differentiated groups of resonances indicative of two separate n-butyl and (DippBDI) ligand environments. Compound 9 could also be synthesized directly through the slow (7 days) reaction of compound 7 with an equimolar amount of P₄ at room temperature or by heating of
a similar reaction at 60°C for 2 days. In this latter case compound 9 crystallized as colorless single crystals suitable for a further X-ray diffraction analysis, allowing the resolution of the solution NMR spectroscopic observations. The result of this analysis is shown in Figure 2 and highlights that the disposition of the phosphorus and magnesium centers within compound 9 is consistent with the coupling pattern observed in the solution-state $^{31}$P NMR spectrum and the multiplicity of signals arising in the $^1$H and $^{13}$C NMR data. The structure comprises two ($^\text{Dipp}$BDI) magnesium units bonded to a [n-Bu$_2$P$_8$]$^{2-}$ cluster dianion derived from the formal reductive coupling of two neutral P$_4$ units. Although the rationality of this \{P$_8$\} cluster synthesis from the combination of two \{P$_4$\} fragments appears to be is unique, its structure may be considered as a heptaphosphanorbornane fused to an exo-oriented cyclotriphosphane ring and a constitutional isomer of the \{P$_8$\} cage within the niobium(V) species [Ph$_2$CP$_8$Nb(OC$^\text{2-Ad}$Mes)$_3$], which was derived through treatment of compound 1 with benzophenone.$^{[14]}$ The single tris-enolato niobium dication and the \{Ph$_2$C\} unit of this previously described species is replaced in the present case by two ($^\text{Dipp}$BDI) magnesium monocations and two n-butyl substituents. The magnesium centers contact the \{P$_8$\} cage structure in endo [Mg(1)] and exo [Mg(2)] configurations with respect to the P(4) bridgehead and provide Mg-P distances of 2.575(3) Å [P(4)-Mg(2)], 2.625(3) Å [P(8)-Mg(2)], 2.594(3) Å [P(2)-Mg(1)] and 2.831(3) Å [P(7)-Mg(1)]. Although these latter values indicate some variability in the charge donation across the cage structure, it should be noted that the Mg-P contacts are all within the range expected [2.45 to 2.985 Å] from a search of the Cambridge structural database.$^{[15]}$ All of the P-P distances within this new cluster are also consistent with P-P single bonds and range between 2.096(3) Å [P(3)-P(4)] and 2.262(4) Å [P(5)-P(6)].
Figure 2: ORTEP representation (30% probability ellipsoids) of compound 9. Hydrogen atoms, iso-propyl methyl groups and \( n \)-butyl groups aside from the P-bonded C(59) and C(63) atoms are removed for clarity. Selected bond lengths (Å) and angles (°): P(1)-P(5) 2.195(3), P(2)-P(3) 2.160(2), P(2)-Mg(1) 2.594(3), P(3)-P(4) 2.096(3), P(3)-P(7) 2.186(3), P(4)-P(5) 2.222(3), P(4)-Mg(2) 2.575(3), P(5)-P(6) 2.262(4), P(6)-P(7) 2.170(4), P(6)-P(8) 2.185(3), P(7)-P(8) 2.153(3), P(7)-Mg(1) 2.831(3), P(8)-Mg(2) 2.625(3), Mg(1)-N(2) 2.046(5), P(2)-P(1)-P(5) 105.23(12), P(1)-P(2)-P(3) 96.79(11), P(1)-P(2)-Mg(1) 90.77(11), P(3)-P(2)-Mg(1) 97.47(9), P(4)-P(3)-P(7) 113.74(12), P(2)-P(3)-P(7) 97.95(10), P(3)-P(4)-P(5) 88.12(10), P(3)-P(4)-Mg(2) 101.38(10), P(5)-P(4)-Mg(2) 91.12(11), P(1)-P(5)-P(4) 102.66(14), P(1)-P(5)-P(6) 92.00(12), P(4)-P(5)-P(6) 105.80(12), P(7)-P(6)-P(8) 59.27(11), P(7)-P(6)-P(5) 106.69(12).

Subsequent preliminary investigations indicated that the kinetic control exerted during the syntheses of compounds 8 and 9 is an apparent consequence of the steric demands of both the supporting \(^{\text{Dipp}}\)BDI platform and the reactive \( n \)-butyl co-ligand. Analysis by \(^{31}\)P NMR spectroscopy of a reaction between \([^{\text{Dipp}}\text{BDI}]_{2}\)MgH and half a molar equivalent of \( \text{P}_4 \) provided a mixture of several phosphorus-containing species. Although a minor component comprised broadened resonances at \( \delta \approx -3.5 \) and \(-159 \) ppm reminiscent of those observed for compound 8, the major product of this reaction was characterized by a triplet signal observed at \( \delta \approx -284.8 \) ppm (\( J_{P-H} = 45 \) Hz). This latter species was identified as the trimeric primary magnesium phosphide, compound 10, through a further X-ray analysis performed on single crystals produced by fractional crystallization of the reaction mixture (Figure 3a). Analytically pure samples of compound 10, notable as the first primary phosphide of a Group 2 element to have been structurally characterized, were found to be unstable once redissolved
in toluene-d$_8$. Monitoring by $^1$H and $^{31}$P NMR spectroscopy at room temperature evidenced the onset of a redistributive process to as yet unidentified magnesium-containing species and the production of PH$_3$, which was observed as a binomial quartet at $\delta$ -245.0 ppm in the $^{31}$P NMR spectrum.

In a similar manner, monitoring by NMR spectroscopy of a reaction analogous to that employed in the synthesis of compound 8 but with replacement of the $N$-{2,6-}iPr$_2$C$_6$H$_3$ groups of the (DippBDI) ligand of 7 by less sterically demanding $N$-{2,4,6-}Me$_3$C$_6$H$_2$ substituents (Me$_3$BDI), indicated less discriminating behavior and reactivity which again resulted in the formation of multiple products. A further reaction performed between [(Me$_3$BDI)Mg$_n$-Bu] and P$_4$ in a 2:3 molar ratio again provided a complex mixture containing at least three new phosphorus-containing compounds. Fractional crystallization from this solution and a resultant single crystal X-ray diffraction analysis of compound 11 (Figure 3b) provided some insight into the nature of these processes. Compound 11 may be viewed as a [P$_3$]$^3^-$ Zintl ion cage reminiscent of compound 4 but decorated with three $\beta$-diketiminato magnesium units. We suggest, therefore, that the formation of this phosphorus cage constitutes a thermodynamic sink during the solution comproportionation of magnesium polyphosphorus species of varying nuclearity.[16]

**Figure 3:** ORTEP representations (30% probability ellipsoids) of (a) compound 10 with hydrogen atoms except those attached to P(1), P(2) and P(3) and iso-propyl methyl groups removed for clarity; (b) compound 11 with hydrogen atoms and mesityl methyl groups removed for clarity.
In conclusion we have shown that the controlled, sequential organomagnesium-mediated activation of white phosphorus may be achieved through selection of an appropriate combination of supporting and reactive co-ligands. We are continuing to explore the parameters of this reactivity and will report our findings in future publications.

**Methods**

Details of the synthesis, characterization data and the crystallographic protocols employed in this study are given in the Supporting Information. CCDC 1057657 - 1057660 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Center via [www.ccdc.cam.ac.uk/data_request/cif](http://www.ccdc.cam.ac.uk/data_request/cif).

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**References**


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