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$[P_4H]^+[Al(OTeF_5)_4]^-$: protonation of white phosphorus with the Brønsted superacid $H[Al(OTeF_5)_4]_{(solv)}^{\ddagger\ddagger}$

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A sustainable transformation of white phosphorus (P_4) into chemicals of higher value is one of the key aspects in modern phosphorus research. Even though the chemistry of P_4 has been investigated for many decades, its chemical reactivity towards the simplest electrophile, the proton, is still virtually unknown. Based on quantum-chemical predictions, we report for the first time the successful protonation of P_4 by the Brønsted acid $H[Al(OTeF_5)_4]_{(solv)}$. Our spectroscopic results are in agreement with acid-mediated activation of P_4 under protonation of an edge of the P_4 -tetrahedron and formation of a three-center two-electron P–H–P bond. These investigations are of fundamental interest as they permit the activation of P_4 with the simplest electrophile as a new prototype reaction for this molecule.

Introduction

White phosphorus (P_4), discovered by Henning Brand in 1669 while searching for the philosopher's stone, is the thermodynamically least stable and most reactive form of phosphorus at room temperature and consists of tetrahedral P_4 molecules. Despite its spontaneous flammability and severe toxicity, P_4 is the easiest form to produce on an industrial scale and is therefore the commercially most important allotrope.¹ Especially its conversion to PCl_3 is of high interest, as it is a base chemical for the production of many organophosphorus compounds.

From a historical point of view, two important chemical reactions of P_4 are described in every good textbook of inorganic chemistry:² (a) the slow oxidation of P_4 vapor to P_4O_{10} under emission of light. This chemoluminescence has coined the name phosphorus, which is derived from the greek mythology ("light-bearer"). (b) the activation and disproportionation of P_4 by aqueous solutions of alkali metal hydroxides. In this way, the industrially relevant phosphine gas (PH_3) is obtained in high purity next to the alkali metal salt of hypophosphorous acid (NaH_2PO_2). More recent studies deal with the degradation of white phosphorus in the presence of other strong nucleophiles, such as organolithium and organomagnesium compounds, carbenes or silylenes, under the topic " P_4 -activation and functionalization".^{3–6} From a mechanistic point of view, a charged

nucleophile (Nu^-) interacts with one of the three energetically degenerate LUMOs of the P_4 molecule (Fig. 1a) under opening of the P_4 tetrahedron to yield a substituted butterfly-like bicyclo [1,1,0]tetraphospha-butane anion (Fig. 1b, I).⁷

Electrophiles, on the other hand, should react at an edge of the tetrahedron, as the two energetically degenerate highest occupied molecular orbitals (HOMO and HOMO–1) have large coefficients at two adjacent phosphorus atoms (Fig. 1a). The situation is, however, much more complicated and the formation of various products is usually observed in this seemingly simple reaction. In the case of Ph_2P^+ and NO^+ , the insertion of these small molecules into one of the P–P bonds is indeed observed, as also theoretically predicted for NO^+ (Fig. 1b, II).^{9–11} In the case of Ag^+ as an example of an electrophilic transition metal center, a weak coordination of Ag^+ to the edge of the P_4 tetrahedron occurs (Fig. 1b, III).¹² However, depending on the steric demand of the metal fragment, the coordination of P_4 via the apex can be enforced, even though the interaction of the energetically low-lying HOMO–5 with the electrophile is necessary to achieve this coordination mode (Fig. 1b, IV).¹³

In contrast to the experimental observations made for the reaction of P_4 with nucleophiles as well as coordinatively and electronically unsaturated transition metal complexes, experimental proof for the structure of the elusive $[P_4H]^+$ cation in solution is still missing in the literature. In fact, weak acids do not react with P_4 due to the rather poor nucleophilicity and weak basicity of white phosphorus. Common strong acids, such as sulfuric acid (H_2SO_4) and nitric acid (HNO_3) cannot be used for the generation of $[P_4H]^+$ as they directly oxidize P_4 to either phosphorous acid (H_3PO_3) and sulfur dioxide (SO_2), or to phosphoric acid (H_3PO_4), nitrogen oxide (NO_2) and water (H_2O), respectively. Hydrogen chloride (HCl) can react with P_4 to form

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† Dedicated to Prof. Dr Peter Jutzi on the occasion of his 80th birthday.

‡ Electronic supplementary information (ESI) available. See DOI: 10.1039/c8sc03023e



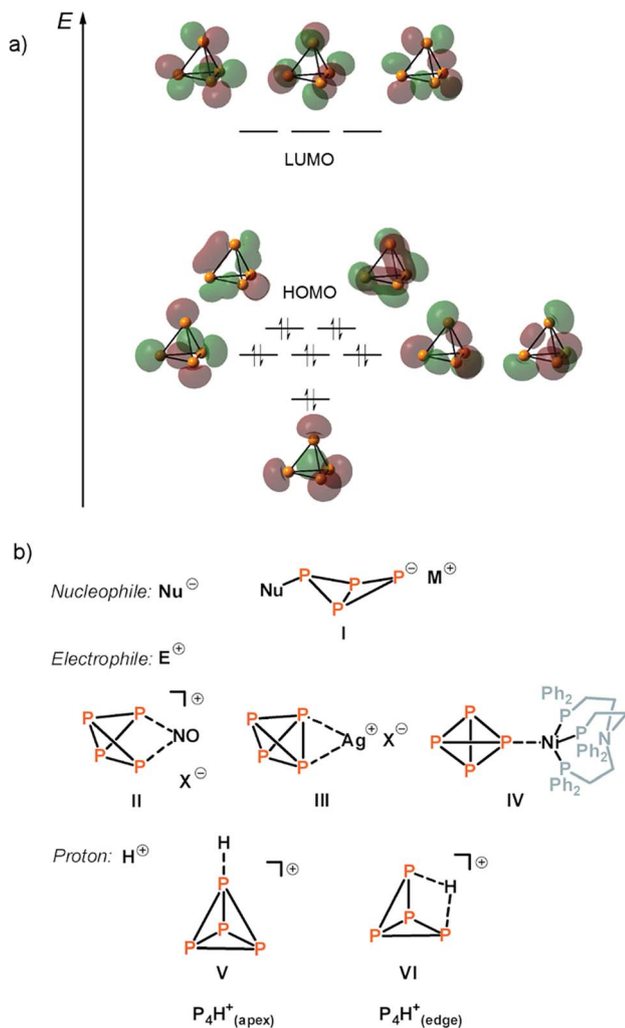


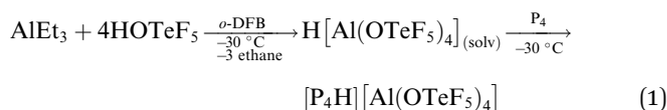
Fig. 1 Molecular orbital scheme of P₄ (a)⁸ and examples of nucleophilic and electrophilic attacks at P₄ and the assumed structures for protonated P₄ (b).

phosphine gas (PH₃) and phosphorus trichloride (PCl₃). Based on *ab initio* calculations, Fluck *et al.*¹⁴ predicted in 1979 that the weakly bound proton in [P₄H]⁺ is located at the apex of the tetrahedron (Fig. 1b, V), while protonation at the edge was predicted to be energetically less favored (Fig. 1b, VI). The authors exclude protonation at the P₃-face. More recent *ab initio* molecular orbital calculations at the MP2/6-31G(d,p) level of theory in 1996 by Abboud, Yáñez and co-workers reveal, however, that the thermodynamically most favourable process is the protonation at the edge under formation of a three-center two-electron (3c-2e) P–H–P bond (Fig. 1b, VI).¹⁵ The same group determined the gas-phase basicity of P₄ by means of Fourier transform ion cyclotron resonance mass spectrometry. In 2000, Ponc and co-workers provided an additional theoretical support for the existence of a non-classical 3c-2e P–H–P bond in [P₄H]⁺ using the generalized population analysis.¹⁶ More recently, Lobayan and Boichichio used a topological analysis of the electron density to describe the 3c-2e P–H–P bond in [P₄H]⁺.¹⁷

Results

Taking the above-mentioned considerations into account, we anticipated that strong acids of conjugated weakly coordinating and non-reactive anions should be excellent reagents for the protonation of P₄. Reed and Nixon, for instance, have shown that phosphabenzene can be protonated by the *in situ* generated Brønsted superacid H(CHB₁₁Me₅Br₆).¹⁸ Also these phosphorus heterocycles are known for their extremely weak basicity. As one of us¹⁹ has recently reported a novel aluminum-based superacidic system containing the weakly coordinating anion [Al(OTeF₅)₄]⁻, we report here now the synthesis and the first spectroscopic proof on the structure of [P₄H]⁺ in solution.

According to quantum-chemical calculations at the B3LYP/def2-TZVPP level, the protonation of P₄ can be achieved by a medium consisting of the Brønsted superacid H[Al(OTeF₅)₄]_(solv) and *ortho*-difluorobenzene (*o*-DFB), see eqn (1).¹⁹ This is due to a slightly lower proton affinity of *o*-DFB (741.6 kJ mol⁻¹) compared to P₄ (748.4 kJ mol⁻¹), as computed at CCSD(T)/aug-cc-pVTZ level of theory.



The reaction product of P₄ and the Brønsted superacid was obtained as a temperature-, moisture- and oxygen-sensitive salt. It shows a clean low-temperature proton-coupled ³¹P NMR spectrum with two equally intense signals at δ = -481.7 and δ = -405.8 ppm with a weak roof effect (Fig. 2a). No other signals were observed in the ³¹P NMR spectrum in the region between δ = 400 ppm and δ = -800 ppm.

This spectrum, which also reveals an additional splitting owing to the higher order of the system, is in accordance with an AX₂Y₂ spin system (¹J(³¹P_X, ³¹P_Y) = 233.95 Hz, ¹J(¹H_A, ³¹P_Y) = 36.70 Hz, ²J(³¹P_X, ¹H_A) = 4.91 Hz). This can only result from the protonation of the P₄ molecule at the P–P-edge. Furthermore, a triplet of triplets at δ = -5.35 ppm appears in the ¹H NMR spectrum (Fig. 2c) showing the corresponding couplings of the proton to P_X and P_Y, respectively. Interestingly, both the chemical shifts and coupling constants are in excellent agreement with the simulated spectra of [P₄H]⁺_(edge), obtained by quantum-chemical calculations (Fig. 2b and c and S3, Table S1†). For comparison reasons, Fig. 2d shows the simulated ³¹P NMR spectrum of the species [P₄H]⁺_(apex), which clearly differs from the experimental results. The NMR studies clearly prove the presence of [P₄H][Al(OTeF₅)₄] and that P₄ is protonated at an edge of the tetrahedron as predicted by quantum-chemical calculations.^{15–17}

We further started to investigate the dynamics of the cation in solution. Interestingly, variable temperature NMR spectroscopy indicates a coalescence of the signals at T = -10 °C (Fig. 3).

The triplet of triplets observed at T = -40 °C in the ¹H NMR spectrum broadens with increasing temperature resulting in a broad singlet (approx. FWHM = 75 Hz) at the coalescence temperature. The chemical shift slightly changes from δ =



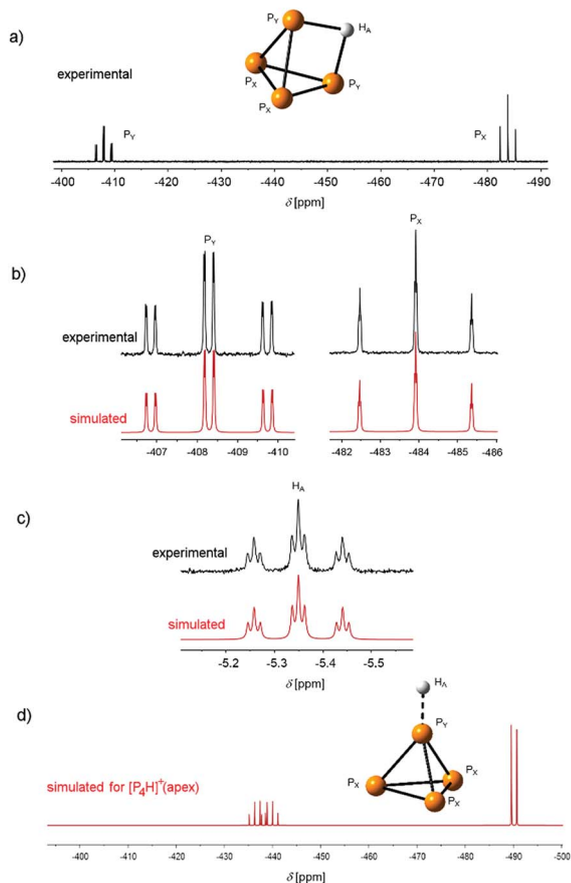


Fig. 2 (a) Low-temperature ($T = -40\text{ }^{\circ}\text{C}$) experimental ^{31}P NMR spectra of $[\text{P}_4\text{H}][\text{Al}(\text{OTeF}_5)_4]$ in *o*-DFB (external lock: $[\text{D}_6]$ acetone). The atom labelling is indicated in accordance with the AX_2Y_2 spin system. (b) ^{31}P NMR spectrum (162 MHz, top) and simulated P_X and P_Y signals (bottom) of the species $[\text{P}_4\text{H}]_{(\text{edge})}^+$. (c) Experimental and simulated ^1H NMR spectrum (401 MHz). The full spectra are provided in Fig. S1 and S2.† (d) Simulated ^{31}P NMR P_X and P_Y signals of the species $[\text{P}_4\text{H}]_{(\text{apex})}^+$.

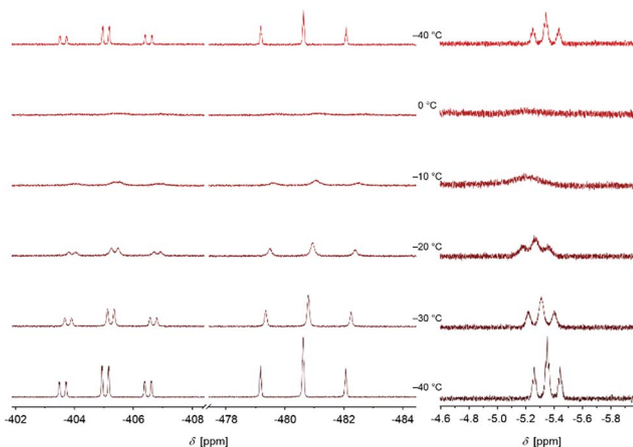


Fig. 3 Excerpt of the ^{31}P and ^1H NMR spectra of $[\text{P}_4\text{H}][\text{Al}(\text{OTeF}_5)_4]$ in *o*-DFB (external lock: $[\text{D}_6]$ acetone) at various temperatures. The sample was first measured at $T = -40\text{ }^{\circ}\text{C}$, annealed stepwise to $T = 0\text{ }^{\circ}\text{C}$ and cooled down again to $T = -40\text{ }^{\circ}\text{C}$ afterwards.

-5.35 ppm at $T = -40\text{ }^{\circ}\text{C}$ to $\delta = -5.19\text{ ppm}$ at $T = 0\text{ }^{\circ}\text{C}$. In the ^{31}P NMR spectrum, a similar process is observed. The two signals are broadened at the coalescence temperature and shifted to a higher field by 0.5 ppm for P_X and P_Y at $T = 0\text{ }^{\circ}\text{C}$. This process is reversible by re-cooling the sample to $T = -40\text{ }^{\circ}\text{C}$ again. No signal for P_4 is detected during this process. Based on these observations, we anticipate a dynamic intramolecular migration of the proton on the P_4 surface. From the experimental dynamic-NMR data, the corresponding barrier can be estimated to $\Delta G^\ddagger = 54.2\text{ kJ mol}^{-1}$.

We noticed, that if an excess of P_4 is present in the reaction mixture, the previously described $[\text{P}_9]^+$ cation²⁰ is formed next to $[\text{P}_4\text{H}]^+$. Upon warming a sample containing a mixture of $[\text{P}_4\text{H}]^+$ and P_4 from $T = -40\text{ }^{\circ}\text{C}$ to $T = -10\text{ }^{\circ}\text{C}$, a fast and full conversion to $[\text{P}_9]^+$ is observed, as detected by NMR spectroscopy. This observation indicates that activation of the P_4 molecule by protonation already occurs at low temperature, while broad band UV/Vis irradiation is necessary to form $[\text{P}_9]^+$ from a $\text{P}_4/[\text{P}_4\text{NO}]^+$ mixture, as reported in the literature before.¹⁰

The $[\text{P}_4\text{H}]^+$ cation was further analyzed by means of mass spectrometry. In the mass spectrum (positive mode), a signal allocated to $[\text{P}_4\text{H}]^+$ appears at $m/z = 124.9$. In addition, signals due to $[\text{P}_4]^+$ ($m/z = 123.8$) as well as of $[\text{Te}]^+$ and $[\text{TeH}]^+$ in a natural isotope distribution ($n = 122, 124\text{--}126$) arise with less intensity. Furthermore, the cations $[\text{P}_3]^+$, $[\text{o-DFB}]^+$, $[\text{o-DFB-H}]^+$ and $[\text{P}_5]^+$ were found. The mass spectrum recorded in the negative mode shows only signals of the four anions $[\text{AlF}_3(\text{OTeF}_5)]^-$, $[\text{AlF}_2(\text{OTeF}_5)_2]^-$, $[\text{AlF}(\text{OTeF}_5)_3]^-$ and $[\text{Al}(\text{OTeF}_5)_4]^-$, see Fig. S7–S9.†

Finally, we investigated $[\text{P}_4\text{H}][\text{Al}(\text{OTeF}_5)_4]$ by means of Raman spectroscopy both in an *o*-DFB solution at $T = -30\text{ }^{\circ}\text{C}$ and as a neat powder at $T = -78\text{ }^{\circ}\text{C}$ in the solid state (Fig. 4). In both spectra, two prominent bands can be observed. The band around $\tilde{\nu} = 1615\text{ cm}^{-1}$ corresponds to the symmetrical P–H–P stretching mode and the band at $\tilde{\nu} = 598\text{ cm}^{-1}$ occurs slightly

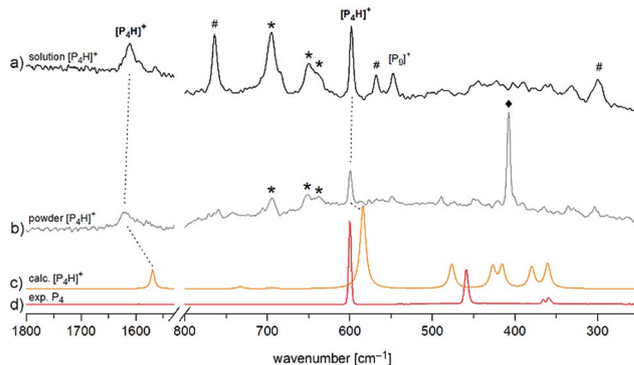


Fig. 4 Enlarged Raman spectrum of (a) $[\text{P}_4\text{H}][\text{Al}(\text{OTeF}_5)_4]$ in *o*-DFB at $T = -30\text{ }^{\circ}\text{C}$ and (b) $[\text{P}_4\text{H}][\text{Al}(\text{OTeF}_5)_4]$ washed with *n*-pentane at $T = -78\text{ }^{\circ}\text{C}$, (c) calculated spectrum of $[\text{P}_4\text{H}]^+$ at the B3LYP/def2-TZVPP level of theory and (d) experimental spectrum of solid P_4 at $T = -196\text{ }^{\circ}\text{C}$. Bands of the anion $[\text{Al}(\text{OTeF}_5)_4]^-$ at $\tilde{\nu} = 695, 650$ and 637 cm^{-1} are marked by an asterisk (*). Bands of the solvent (*o*-DFB: #, *n*-pentane: ♦) are indicated as well. Full spectra are provided in Fig. S5 and S6.†



shifted with respect to the breathing mode of neat P_4 and is assigned to the corresponding mode of $[P_4H]^+$.

Both the experimental band positions agree well with the computed wavenumbers at the B3LYP/def2-TZVPP level of theory at $\tilde{\nu} = 1569\text{ cm}^{-1}$ (A_1) and $\tilde{\nu} = 584\text{ cm}^{-1}$ (A_1), respectively. Further Raman bands are predicted between $\tilde{\nu} = 360\text{ cm}^{-1}$ and $\tilde{\nu} = 476\text{ cm}^{-1}$ but they are difficult to assign in the experimental spectrum due to their rather low intensities and their partial interference with the bands of $[P_9]^+$ impurities. It should be pointed out that special care must be taken by isolating $[P_4H][Al(OTeF_5)_4]$ as a solid, as one sample exploded during the Raman measurement at dry ice temperature after approx. 300 scans at 75 mW, Fig. 4b. Attempts to record low-temperature IR spectra of the $[P_4H]^+$ cation were unsuccessful, as the strongest IR band of $[P_4H]^+$ is hidden by very prominent bands of the anion at $\tilde{\nu} = 713\text{ cm}^{-1}$ and $\tilde{\nu} = 695\text{ cm}^{-1}$. Also, the strongest band of *o*-DFB occurs at $\tilde{\nu} = 750\text{ cm}^{-1}$. Nevertheless, low temperature ($T = -30\text{ }^\circ\text{C}$) IR spectra of the liquid phase of $[P_4H][Al(OTeF_5)_4]$ have been recorded using a glass fiber ATR head, which are provided in Fig. S10 and S11.†

Our quantum-chemical calculations at the coupled-cluster CCSD(T)/aug-cc-pVTZ level agree very well with the experimental results of the protonation of a P_4 edge. This position is also expected from the MO diagram, where the HOMO orbital is located along the P_4 edge (Fig. 1a), leading to a three-center two-electron P–H–P bond. This gives rise to a C_{2v} symmetric structure with an elongation of the $P_Y \cdots P_Y$ bond of 20.4 pm compared to the bond length of 221.8 pm in the P_4 tetrahedral structure. The bond distances P_Y-P_X and P_X-P_X are less affected by 1.4 and 6.3 pm, respectively (Fig. 5). The computed minimum structure for protonation at the apex of the P_4 molecule is 61.4 kJ mol^{-1} higher in energy than the global minimum structure. An apex protonation would also lead to a computed P–H stretching mode at $\tilde{\nu} = 2502\text{ cm}^{-1}$, which is more than $\tilde{\nu} = 900\text{ cm}^{-1}$ above the experimentally observed band at $\tilde{\nu} = 620\text{ cm}^{-1}$. Surprisingly, the protonation and simultaneous opening of the tetrahedral structure lead to a P_4 -butterfly type minimum structure (Fig. 5), while protonation of the triangle surface shows a higher order saddle point. Both structures will be higher in energy by 74.3 and 88.5 kJ mol^{-1} compared to the

global minimum structure of $[P_4H]^+$, see Fig. 5. For details of the computed structural parameters see Tables S2 and S3.†

Conclusions

Based on these results, we could reveal for the first time the structure of protonated white phosphorus in solution. Both experimental results and quantum-chemical calculations provide evidence for a protonation at the edge of the P_4 molecule. The opening of the P_4 -tetrahedron *via* the simplest electrophile (H^+) under formation of a three-center two-electron P–H–P bond is of fundamental interest for understanding the reactivity of this intriguing phosphorus allotrope. It is expected that this groundbreaking result is important for the development of chemical processes related to the activation and further functionalization of elemental phosphorus by electrophiles.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- 1 F. Krafft, *Angew. Chem., Int. Ed. Engl.*, 1969, **8**, 660.
- 2 R. Steudel, *Chemie der Nichtmetalle, Synthesen - Strukturen - Bindung - Verwendung de Gruyter*, Berlin, edn 4, 2014.
- 3 M. Scheer, G. Balázs and A. Seitz, *Chem. Rev.*, 2010, **110**, 4236.
- 4 (a) J. E. Borger, A. W. Ehlers, M. Lutz, J. Chris Slootweg and K. Lammertsma, *Angew. Chem., Int. Ed.*, 2014, **53**, 12836; (b) L. Xu, Y. Chi, S. Du, W.-X. Zhang and Z. Xi, *Angew. Chem., Int. Ed.*, 2016, **55**, 9187; (c) M. Arrowsmith, M. S. Hill, A. L. Johnson, G. Kociok-Köhn and M. F. Mahon, *Angew. Chem., Int. Ed.*, 2015, **54**, 7882.
- 5 (a) J. D. Masuda, W. W. Schoeller, B. Donnadiou and G. Bertrand, *Angew. Chem., Int. Ed.*, 2007, **46**, 7052; (b) J. D. Masuda, W. W. Schoeller, B. Donnadiou and G. Bertrand, *J. Am. Chem. Soc.*, 2007, **129**, 14180; (c) O. Back, G. Kuchenbeiser, B. Donnadiou and G. Bertrand, *Angew. Chem., Int. Ed.*, 2009, **48**, 5530; (d) D. Martin, M. Soleilhavoup and G. Bertrand, *Chem. Sci.*, 2011, **2**, 389; (e) C. L. Dorsey, B. M. Squires and T. W. Hudnell, *Angew. Chem., Int. Ed.*, 2013, **52**, 4462; (f) C. D. Martin, C. M. Weinstein, C. E. Moore, A. L. Rheingold and G. Bertrand, *Chem. Commun.*, 2013, **49**, 4486; (g) M. Cicač-Hudi, J. Bendi, S. H. Schlindwein, M. Bispringhoff, M. Nieger, H. Grützmacher and D. Gudat, *Eur. J. Inorg. Chem.*, 2016, **5**, 649.

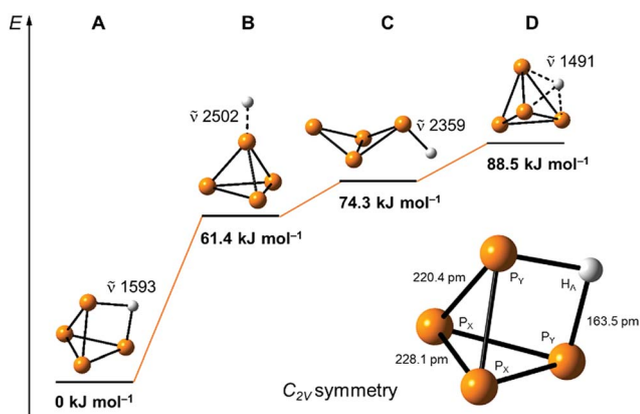


Fig. 5 Computed relative energies and P–H vibrations of optimized $[P_4H]^+$ structures at the CCSD(T)/aug-cc-pVTZ level of theory.



- 6 (a) Y. Xiong, S. Yao, M. Brym and M. Driess, *Angew. Chem.*, 2007, **119**, 4595; (b) S. Khan, R. Michel, S. S. Sen, H. W. Roesky and D. Stalke, *Angew. Chem., Int. Ed.*, 2011, **50**, 11786.
- 7 R. Riedel, R. H.-D. Hausen and E. Fluck, *Angew. Chem.*, 1985, **97**, 1050.
- 8 B. M. Cossairt and C. C. Cummins, *J. Am. Chem. Soc.*, 2009, **131**, 15501.
- 9 J. J. Weigand, M. Holthausen and R. Fröhlich, *Angew. Chem.*, 2009, **121**, 301.
- 10 T. Köchner, S. Riedel, A. J. Lehner, H. Scherer, I. Raabe, T. A. Engesser, F. W. Scholz, U. Gellrich, P. Eiden, R. A. Paz Schmidt, D. A. Plattner and I. Krossing, *Angew. Chem., Int. Ed.*, 2010, **49**, 8139.
- 11 K. F. Hoffmann, A. Wiesner, N. Subat, S. Steinhauer, S. Riedel and Z. Anorg, *Z. Anorg. Allg. Chem.*, 2018, DOI: 10.1002/zaac.201800174.
- 12 I. Krossing and L. van Wüllen, *Chem. - Eur. J.*, 2002, **8**, 700.
- 13 (a) P. Dapporto, S. Midollini and L. Sacconi, *Angew. Chem., Int. Ed.*, 1979, **18**, 469; (b) M. Peruzzini, S. Mañas, A. Romerosa and A. Vacca, *Mendeleev Commun.*, 2000, **10**, 134; (c) T. Gröer, G. Baum and M. Scheer, *Organometallics*, 1998, **17**, 5916.
- 14 E. Fluck, C. M. E. Pavlidou and R. Janoschek, *Phosphorus Sulfur Relat. Elem.*, 1979, **6**, 469.
- 15 J.-L. M. Abboud, M. Herreros, R. Notario, M. Esseffar, O. Mó and M. Yáñez, *J. Am. Chem. Soc.*, 1996, **118**, 1126.
- 16 R. Bochicchio, L. Lain, A. Torre and R. Ponc, *Croat. Chem. Acta*, 2000, **73**, 1039.
- 17 R. M. Lobayan and R. Bochicchio, *J. Phys. Chem. A*, 2015, **119**, 7000.
- 18 Y. Zhang, F. S. Tham, J. F. Nixon, C. Taylor, J. C. Green and C. A. Reed, *Angew. Chem., Int. Ed.*, 2008, **47**, 3801.
- 19 A. Wiesner, T. W. Gries, S. Steinhauer, H. Beckers and S. Riedel, *Angew. Chem., Int. Ed.*, 2017, **56**, 8263.
- 20 T. Köchner, T. A. Engesser, H. Scherer, D. A. Plattner, A. Steffani and I. Krossing, *Angew. Chem., Int. Ed.*, 2012, **51**, 6529.

