

(Iminomethylidene)phosphines (RP=C=NR)

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There has been much recent interest in compounds incorporating $P=C^{2,3}$ and $P=P^4$ double bonds. Furthermore, the stable (2,2dimethylpropylidyne)phosphine, (CH₃)₃CC=P, featuring a PC triple bond has been prepared.⁵ However, previous attempts to isolate cumulenes possessing a two-coordinate phosphorus atom, RP=C=X, have failed, 2,3,6 except in a single case, the preparation of the stable and sterically protected $(CH_3)_3CP = C = NC(CH_3)_3$ We now wish to report a general method of preparation of unstable (iminomethylidene)phosphines (monophosphorus analogues of carbodiimides), their direct observation by IR and mass spectrometry, and their initial chemical reactions.

The required starting materials, (iminomethyl)phosphines 1, exist in equilibrium with carbamoylphosphines [(CH₃)₃SiP(R)- $CON(Ph)Si(CH_3)_3$ and smaller amounts of alkylidenephosphines $RP=C(OSi(CH_3)_3)N(Ph)Si(CH_3)_3$ ⁸ In analogy with the reaction used in the preparation of $(CH_3)_3CC \equiv P$,⁵ the compounds 1 could be expected to eliminate hexamethyldisiloxane and thus give rise to the desired (iminomethylidene)phosphines, RP= C=NPh (Scheme I).

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In the event, the treatment of 1 with a catalytic amount of solid NaOH led only to the isolation of the dimers $3.^9$ 3b and 3e were also obtained by spontaneous decomposition of the adducts 2b and

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^{(9) (}a) Satisfactory elemental analyses (C, H, N, P) were obtained for these compounds. (b) X-ray crystallographic determinations of 3c and 3d confirm the structure assignments. The two exocyclic phosphorus ligands are trans oriented. Full crystallographic data will be published: Becker, G.; Härer, J.; Riffel, H.; Uhl, G.; Wessely, H.-J., in preparation.



Figure 2. CID mass spectrum of m/z 211 from the pyrolysis of 3a at 430

2e, the latter formed by the addition of phenyl isothiocyanate to the alkyl- or aryl(trimethylsilyl)phosphine.¹⁰

If the diphosphetanes 3 are, in fact, dimers of (iminomethylidene)phosphines, the monomers might be regenerated via a cycloreversion under suitable reaction conditions. This, indeed, is readily achieved by flash vacuum pyrolysis of 3. Product formation was monitored by low-temperature IR spectroscopy, and a pyrolysis unit has also been attached to the ion source of a reversed-geometry Varian MAT 311A mass spectrometer¹¹ equipped with a collision chamber for obtaining CID (collisionally induced dissociation) spectra of initial pyrolysis products.

2,4-Bis(phenylimino)-1,3-diphenyl-1,3-diphosphetane (3a) (mp 139 °C) was vaporized at 139-145 °C (10^{-5} torr). When the pyrolysis temperature was increased to 400 °C, an IR band at 1853 cm⁻¹ appeared in the pyrolyzate condensed at -196 °C. The maximum intensity of this band was observed at a pyrolysis temperature at 480 °C (Figure 1). On warming the pyrolyzate to -55 °C, the intensity of the 1853-cm⁻¹ band started decreasing; at the same time, a band at 1560 cm⁻¹ due to the starting material 3a started increasing. This process was complete at -30 °C, and after warming to room temperature 3a was recovered in better than 90% yield.12

These observations indicate that 3a dissociates into two molecules of the (iminomethylidene)phosphine 4a on gas-phase pyrolysis. The monomeric 4a reverts to 3a above -55 °C in the solid state. The strong band at 1853 cm⁻¹ in the IR of 4a is assigned to the asymmetric stretching vibration of the P=C=N moiety. For comparison, carbodiimides (RN=C=NR) absorb near 2100 cm⁻¹.

The analogous pyrolysis of 3b (mp 220 °C) at 480 °C (sublimation temperature 110 °C, increasing to 200 °C in 35 min) gave rise to a strong IR band at 1839 cm⁻¹ ascribed to 4b. This material started redimerizing to 3b at -25 °C, a process that was complete at 0 °C. Comparable results were obtained with the precursors 3c-e.

The IR assignments were corroborated by using the mass spectrometry reactor. With increasing pyrolysis temperature, the

M⁺ peak due to 3a decreased, while that of 4a increased. The CID mass spectrum of 4a obtained at 430 °C is shown in Figure 2. The base peak at m/z 183 may be ascribed to the dibenzophospholyl cation, $C_{12}\dot{H}_8P^+$, which is typical of diphenylated phosphorus compounds.¹³ In the pyrolysis of **3b** the parent peak of the precursor (m/z 382) disappeared at 490 °C, while that of **4b** $(m/z \ 191)$ reached maximum intensity. Here, the spectrum is dominated by a loss of isobutene to give m/z 135, formally corresponding to PhN=C=Ph+ or PhNH-C=P+

We are continuing the studies of (iminomethylidene)phosphines, in particular cycloaddition reactions and attempts to obtain other phosphorus-containing cumulenes, e.g., RP=C=O.

Registry No. 1a, 24103-42-2; 1b, 87218-80-2; 1c, 87729-47-3; 1d, 87729-48-4; 1e, 87729-49-5; 2b, 87729-50-8; 2e, 87729-51-9; 3a, 87729-52-0; 3b, 87218-81-3; 3c, 87729-53-1; 3d, 87729-54-2; 3e, 87729-55-3; 4a, 87729-56-4; 4b, 87218-77-7; 4c, 87729-57-5; 4d, 87729-58-6; **4e**, 87729-59-7.

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Perlaux, J. Bull. Soc. Chim. Belg. 1976, 85, 69. (12) The band at 2120 cm⁻¹ in Figure 1 is due to phenyl isocyanide, formed in a competing thermal fragmentation of 3a. This material evaporates during warm-up and thus does not contaminate the final product. A yield of ca. 10% of phenyl isocyanide was obtained by distilling it into a cold trap and subsequently identifying it by comparison with an authentic sample.