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Versatile Synthesis of Phospholides from Open-Chain Precursors. Application to Annelated Pyrrole− and Silole−Phosphole Rings

Youzhi Xu,[†] Zhihua Wang,[†] Zhenjie Gan,[†] Qiuzhen Xi,[†] Zheng Duan,^{*,†} and François Mathey*^{,†,‡}

† College of Chemistry and Molecular Engineering, International Phosphorus Laboratory, [Jo](#page-2-0)int Research Laboratory for [Fu](#page-2-0)nctional Organophosphorus Materials of Henan Province, Zhengzhou University, Zhengzhou 450001, P. R. China ‡ Division of Chemistry & Biological Chemistry, Nanyang Technological University, 21 Nanyang Link, Singapore 637371

S Supporting Information

[AB](#page-2-0)STRACT: [Phospholides](#page-2-0) are easily obtained by treatment of the open-chain acetylenic phosphines shown by an excess of lithium at room temperature in THF (12 examples).

Phospholides are one of the two fundamental aromatic rings of phosphorus−carbon heterocyclic chemistry.¹ However, the number of synthetic routes to these species is very limited in spite of their key role in organophosphorus sy[n](#page-2-0)thesis and coordination chemistry. In fact, they are practically always synthesized by reductive cleavage of the P−R exocyclic bond of phospholes, 2 the driving force being the conversion of the nonaromatic phosphole into the aromatic phospholide. Alternativel[y,](#page-2-0) it is possible to couple the easy $\begin{bmatrix} 1,5 \end{bmatrix}$ shift of the sp²-carbon substituent of phospholes from P to C α with a deprotonation of the resulting $2H$ -phosphole.³ To the best of our knowledge, the only case where a noncyclic precursor has been used concerns the formation of a calciu[m](#page-2-0) phospholide by reaction of a calcium bis(trimethylsilyl) phosphide with a 1,3 diyne.⁴ We wish to report hereafter on a versatile access to phospholides starting from easily made noncyclic phosphines and [all](#page-2-0)owing to synthesis of hitherto unknown annelated species.

Our work began with the synthesis of two families of openchain phosphines incorporating a carbon−carbon triple bond at the γ position from the phosphorus center. The first family (2) derives from triphenylphosphine. Its synthesis relies on a classical Sonogashira coupling (Scheme 1). Phosphine 2a has

already been described in the literature.⁵ The second family (4) , incorporating a Z-enyne functionality, was prepared from the appropriate bromo enynes (3) wh[os](#page-2-0)e synthesis has been recently described 6 (Scheme 2). These two families of phosphines have a backbone similar to those of secondary phosphine oxides t[ha](#page-2-0)t have been previously used in a synthesis of phosphindoles.

Scheme 2. Synth[es](#page-2-0)is of Diphenylphosphinoenynes

This led us to investigate the reaction of lithium with our two families of phosphines. A statistical cleavage of the P−C bonds would lead to a 66% yield of the anion 5 whose cyclization would give a phosphole or a phosphindole (Scheme 3).

The actual results of our experiments with a 5- to 6-fold excess of lithium were more satisfactory than expected. In fact, the formation of Ph₂PLi, as detected by its ${}^{31}P$ resonance at

Scheme 3. Proposed Synthetic Scheme

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Figure 1. ³¹P chemical shifts of phospholides and phosphindolides (in THF with Li⁺ as counterion).

Scheme 4. Synthesis of Phosphindoles

Scheme 5. Synthesis of Phospholes

Due to the use of an excess of lithium, the crude solutions containing the anions 6 or 8 also contain phenyllithium which is formed by reduction of the initial phenyl radical by the excess of metal. Treating the solution of 6a by iodine and sulfur led to 7i $(R = R¹ = Ph, X = S)$. In order to investigate why lithium almost selectively cleaves the P−Ph bonds of 2 and 4, we decided to study the electronic structure of radical 10 by DFT

at the UB3LYP/6-311G (d,p) level. Radicals such as 10 are known to be involved in the reduction of triarylphosphines by alkali metals.¹⁰ The geometrical structure of 10 is shown in Figure (2). The most striking characteristic of this structure is

Figure 2. Computed structure of radical anion 10. Main distances (Å) and angles (deg): P12−C13 1.875, P12−C3 1.835, P12−C24 1.835, C24−C25 1.456, C25−C34 1.411, C34−C35 1.216; C3−P12−C13 101.38, C3−P12−C24 105.06; C13−P12−C24 102.92 C24−C25− C34 122.39.

that the P−C13 (P−Ph) bond at 1.875 Å is significantly longer than the two other P−C bonds at 1.835 Å. The SOMO (Figure 3) is antibonding on this P−C13 bond, whereas it is bonding

and highly localized on the two other P−C bonds. These data indicate that the radical evolves preferentially by cleavage of this P−Ph bond.

A recent theoretical paper has stressed the interesting properties of ladder-type heterotetracenes with dual bridging atoms for optoelectronic applications.¹¹ A mixed phosphorussilicon species 11b was included among the studied molecules. It is possible to use this new r[out](#page-2-0)e to phospholides to synthesize 11b and its derivatives as shown in Scheme 6.

Scheme 6. Synthesis of P,Si-Tetracenes

The initially formed β -lithiated phosphindole spontaneously cyclizes by demethylation of the trimethylsilyl substituent to build the silole ring. Somewhat similar demethylations leading to benzosiloles have been described in the literature.^{12,13} The X-ray crystal structure of 11c is shown in Figure 4. The core of the molecule is slightly bent by 3.2° around the C7−C8 junction between the phosphole and the silole planes.

Figure 4. X-ray crystal structure of 11c. Main distances (Å) and angles (deg): P1−O1 1.481(3), P1−C1 1.811(4), P1−C8 1.821(4), P1−C15 1.813(5), Si1−C7 1.887(4), Si1−C10 1.886(3), C1−C6 1.405(6), C6−C7 1.474(5), C7−C8 1.355(5), C8−C9 1.468(5), C9−C10 1.415(5); C1−P1−C8 91.55(18), C7−Si1−C10 90.72(16).

Another original species 14b incorporating fused pyrrole and phosphole rings has also been obtained as shown in Scheme 7.

In conclusion, we have developed the first general synthesis of phospholides and annelated phospholides starting from open-chain precursors. This synthesis has served to prepare a new phosphorus−silicon heterotetracene of interest for optoelectronic studies and a new phosphole−pyrrole annelated system. Many more applications of this new scheme can be envisaged.

■ ASSOCIATED CONTENT

S Supporting Information

Experimental section, NMR data for 2−14 and X-ray data for 11c This material is available free of charge via the Internet at http://pubs.acs.org.

■ AUTHOR INFORMATION

Corresponding Authors

*E-mail: duanzheng@zzu.edu.cn.

*E-mail: fmathey@ntu.edu.sg.

Notes

The authors declare no competing financial interest.

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■ **DEDICATION**

This work is dedicated to Prof. Tamotsu Takahashi on the occasion of his 60th birthday.

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