A Reinvestigation of the Reactions of Enolate Anions with Cyclotriphosphazenes

CHRISTOPHER W. ALLEN, KOLIKKARA RAMACHAN-DRAN, RANDALL P. BRIGHT and JONATHAN C. SHAW

Department of Chemistry, University of Vermont, Burlington, Vt. 05405, U.S.A.

Received October 30, 1981

Introduction

Enolate anions of ketones are ambident nucleophiles which have attracted considerable attention [I] . We have previously reported the reactions of the lithium enolate anions of acetophenone and cyclohexanone with hexafluorocyclotriphosphazene, $N_3P_3F_6$, and based on limited nmr data suggested that attack occurred at the carbon end of the nucleophile leading to ketones with the phosphazene moiety bonded to the α -carbon atom [2]. Recently, Tate et al. reported the reactions of polydichlorophosphazene with various enolate anions and proposed that attack occurred at the oxygen end of the nucleophile [3], thus leading to vinyl alcohol derivatives. This investigation is an attempt to resolve these conflicting interpretations concering the nature of these materials.

Experimental

Hexachlorocyclotriphosphazene, $N_3P_3Cl_6$, (Ethyl Corp.) was recrystallized from petroleum ether to a constant m.p. 113 °C. n-Butyl lithium (1.6 M solution in hexane) was obtained from Aldrich. Tetrahydrofuran (THF) (Aldrich) was distilled from sodium-benzophenone ketyl. Petroleum ether (b.p. $35-55$ °C), benzene and ethyl acetate (Fisher) were distilled by standard procedures. NMR spectra $(in$ CDCl₃) were recorded on a Bruker WM250 spectrometer operating at 250.1 MHz (^1H) , 62.9 MHz (^{13}C) , and 101.2 MHz (^{31}P) . Tetramethyl silane (TMS) was used as an internal reference for ¹H and ¹³C NMR measurements. For ³¹P NMR, 85% H3P04 was used as an external standard. Chemical shifts upfield to the reference are assigned a negative sign. 13 C and 31 P NMR spectra were recorded under conditions of broad band decoupling. Infrared (IR) spectra were obtained as their thin films (NaCl discs) on a Beckman IR 20A spectrometer. Mass spectra were recorded on a Perkin-Elmer RMU-6D spectrometer operating at 80 eV. Elemental analyses were performed by Integral Microanalytical Laboratories. Hexafluorocyclotriphosphazene [4] and the derivatives derived from the enolate anions of acetophenone $(I, N_3P_3F_5OC_8H_7)$ and cyclohexanone (II, $N_3P_3F_5OC_6H_9O$ [2] were prepared by previously reported procedures.

Preparation of N₃P₃ $Cl₅ OCH = CH₂ (III)$

A solution of n-butyl lithium (45 ml, 0.07 mol) was added to tetrahydrofuran (THF) (75 ml) at room temperature, using the apparatus described elsewhere [5]. The resulting reaction mixture was stirred for 16 hr then slowly added to a solution of 10.5 g (0.03 mol) of $N_3P_3Cl_6$ in tetrahydrofuran at 0 °C under a nitrogen atmosphere. The resulting mixture was allowed to come to room temperature and was kept stirred for four days. The solvent was removed and the residue extracted with hexane. Following filtration and removal of hexane, 10.2 g of a pale yellow liquid was obtained. A 2 gm sample of this material was purified using the flash chromatography technique [6] using petroleum ether as the eluant. A 0.92 g (44.3% of theory) sample of a colorless liquid, b.p. 75 "C at 0.05 mm Hg, was obtained. *Anal.* Calcd. for $N_3P_3Cl_5OC_2H_3$ (III); C, 6.75; H, 0.84; mol. wt. 353. Found: C, 6.74; H, 0.75; mol. wt. 353 (mass spectrum).

IR $(cm⁻¹)$: 1650 (s, C=C str), 1220 (s, PN str), 1110 (s, PO str), 1035(s), 93O(w, PCl), 875(m, PCl) 750(m, PCl).

Preparation of N₃P₃F₅ OCH=CH₂ (IV)

The reaction of 40 ml (0.064) of n-butyl lithium solution in excess THF was added to 16.0 g (0.064 $m \sim 100$ F₆ as previously described. After removal of most of the solvent, the remaining product/solvent mixture was carefully fractionated at room temperature. The remaining solvent distilled at 5 mm Hg and the product at 1 mm Hg. A sample of 0.56 g (3.2% of theory) of a colorless liquid was obtained. Anal. Calcd. for $N_3P_3F_5OC_2H_3$ (IV); mol. wt. 273. Found: mol. wt. 273 (mass spectrum).

IR $(cm⁻¹)$: 1645 $(m, C=C)$, 1290 $(s, PN str)$, 1125 (s, PO str), 1060(m), 1010(m), 95O(s, PF asym), 880(s), 845(s, PF), 800(m).

A ttemp ted Derivatization of I and II

In separate experiments, the acetophenone (I) and cyclohexanone (II) derivatives of $N_3P_3F_6$ were treated with bromine (Br_2/CCl_4) and hydrogen (Hz/lo% Pt on activated carbon). In both the bromination and hydrogenation experiments, a large number of products formed which resisted separation.

^a All coupling constants are given in Hz. ^{b 1}H and ¹³C chemical shifts are given in ppm relative to TMS. ^{e 31}P chemical shifts are given in ppm relative to 85% H₂PO₄.

Results and Discussion

The two possible reaction pathways for the ambi- $\frac{d}{dx}$ and $\frac{d}{dx}$ is the sphere with a health and $\frac{d}{dx}$ ϵ ne cholate amons with a hexanalocyclotifulosphazene are shown below. Our first attempt at resolving
this question involved derivatization with bromine on

hydrogen. If the vinyloxy (route b) derivative y the simple and the simple and the simple address would be a simple would be a simple would be a simple would formed, then the simple addition compounds would be readily identifiable. Unfortunately, these reactions gave rise to a large number of products so we turned
our attention to nmr spectroscopic techniques. In order to gain more useful information from the nmr studies, we prepared derivatives containing hydrogen tuties, we prepared derivatives containing hydrogen tonis on both carbon atoms. The enotate amon or acetaldehyde is quantitatively generated from the metalation of THF with n-butyl lithium [7]

$$
C_4H_8O + n \cdot C_4H_9Li \rightarrow LiOC_2H_3 + C_2H_4 + C_4H_{10}
$$

and gives derivatives of both the hexachloro- and $\frac{1}{2}$ and $\frac{1}{2}$ is the mnr definition of $\frac{1}{2}$.

$$
LiOC2H3 + N3P3X6 \rightarrow N3P3X5OC2H3 III: X = ClIV: X = F
$$

 $13C$, $31P$) for compounds I-IV are reported in Table I. The compounds it incrependem rubber.

The at min spectrum of the closely resembles that of vinyl acetate except for the additional phosphorus coupling for each proton and can be analyzed as such. The observed spectrum is not consistent with a carbon bonded derivative since H_x exhibits more extensive coupling than it would as an aldehyde proton. The chemical shifts are more appropriate for olefinic rather than for alkyl and aldehyde protons. The same arguments apply to IV. The previous H nmr data on compounds I and II were obtained at 60 MHz and were poorly resolved $[2]$. In the 250 MHz spectrum, the additional coupling appropriate to an olefin but not to an α -substituted ketone, is observed. The separation between H_a and H_b in I is too large to be $\frac{1}{2}$ cparation between $\frac{1}{2}$ and $\frac{1}{2}$. In 1 is too large to be F_{eff} for F_{eff} in an any phosphaze in F_{eff} . Furthermore, only H_a is observed in II, thus the two peaks (H_a, H_b) observed in I are not reasonably ascribed to phosphorus-proton coupling. The s cribed to phosphorus proton coupling. The from the the theorem the 13C nmr data. The on comes non the ϵ min that filter are no resonances in the alkyl or carbonyl ranges [9] while those in the olefin range match the number predicted

from a vinyloxy derivative. The assignments of C_1 and C_2 shifts in III were confirmed by off resonance decoupling in which C_1 becomes a doublet and C_2 a triplet. The large shielding of C_1 is due to mesomeric interactions of C_1 with oxygen lone pair electrons [9]. The magnitude of the phosphoruscarbon coupling constants is variable and does not provide a useful structural tool in this case. The ^{31}P nmr spectra of I-IV show little variation with or without broad band proton decoupling. If the phosphorus atom were bonded to a methylene group, s_{S} is a steam were bonded to a metry fine group, E^r
Resolution in CH PXX resonance.
All of the nmr data which we have presented sup-

port the proposal of Tate [3] that attack of the enolate is at the oxygen end, rather than at the carbon end, of the nucleophile. The IR data are also consistent with this conclusion. The bands around 1650 cm^{-1} can be ascribed to olefinic stretching modes rather than carbonyl modes and bands around 1130 cm^{-1} can be assigned to a POC vibration rather than unusual PN vibrations. These materials, especially the newly reported III and IV, represent any the hewry reported in and \mathbf{r} , represent nomer example or organorunctional prospilatores [10] which may be expected to exhibit interesting
monomer and polymer chemistry. Work along these lines is currently in progress in our laboratory.

Acknowledgements

This work was supported in part by the Office of Naval Research. We also wish to thank Dr. D. P. Tate of Firestone Corp. for useful discussion.

References

- 1 H. 0 House and V. Kromer, *J. Org. Chem., 28, 3362 2* J. G. DuPont and C. W. Allen, Inorg. *Chem., 16, 2964* (1, U H
- *3* D. P. Tate, W. L. Hergenrother, 3. W. Kang and D. 1:. . v. DI
.
- Graves, *Pol_vmer Preprints, 20, 177* (1979). Graves, *Polymer Preprints*, 20, 177 (1979).
- *(London)*, 34 */* (1961). *(London), 347* (1961).
- and K. RamachandranfJ. *Chem. Educ., 57, 564* (1980). ng K. Kamachangran, *J. Chem. Eguc.*, *J*., 304 (1980).
W. O. Orill. M. K. L. A. Mitra. *J. Orn. Chem., 43*,
- *7* R. B. Bates, L. M. Kroposki and D. E. Potter, *J. Org.* 7. L. SIIII, N
2004/1078).
- *8 P. T. A. S. M. Rangardan, S. M. Rangardan, S. M. Rangardan, Indian I. L. Paddock, ¹ Chem., 37, 560 (1972).*
- **6221 (1979).**
 $\overline{a} = \overline{b}$ *Chem., 12, 316 Chem., 21, 316 2020*. hem., 12, 310 (1973);
i. F. *i. H. J. Harris, <i>J. A. Chem. Sot.,* 101, 1. К. АЦСОСК
.001 (1979).
- \overline{a} μ B. Strothers, Carbon-Academic Press, N.Y. (1972).
- *Res. Dev., 18,* 81 (1979).