Intramolecular Cyclization of *ω*-Haloalkylsubstituted Thiophosphorylacetonitriles: Synthesis and Stereochemistry of 3-Cyano-2 oxo-1,2-thiaphosphacyclanes

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ABSTRACT: *Intramolecular S-alkylation in a series of* -*-haloalkyl-substituted thiophosphorylacetonitriles* **5–7** *presents an effective synthetic route to the hitherto unknown 3-cyano-2-oxo-1,2-thiaphospholanes* **14** *and thiaphosphinanes* **15***. The compounds were obtained as a mixture of* cis*- and* trans*-isomers that were resolved to individual stereoisomers in most cases. For some of them, X-ray diffraction analysis has been performed. It was shown that 31P NMR spectroscopy can be used to assign the stereochemistry of* 3-cyano-2-oxo-1,2-thiaphosphacyclanes. © 2002 John Wiley & Sons, Inc. Heteroatom Chem 13:1–21, 2002; DOI 10.1002/hc.1101

INTRODUCTION

By now, many phosphorus-containing heterocycles have been described in the literature, among them phosphacyclanes with a single phosphorus heteroatom [1], and polyheteraphosphacyclanes and unsaturated derivatives [2–6] have been investigated most extensively. As regards saturated 1,2 heteraphosphacyclanes with oxygen, nitrogen, or sulfur as heteroatoms, 1,2-oxaphosphacyclanes have been studied sufficiently thoroughly, and some of them have found practical application in the industry, specifically as nonflammable hydraulic liquids and plasticizers [7,8], additives to oils, lubricants [9], flame retardants [10,11], thermostabilizing additives to fibers [12–14], and surfactants [15]. Moreover, despite the fact that 1,2-thiaphosphacyclanes should be of no less practical interest, at the beginning of the present investigation, only a few of the simplest, nonfunctionalized representatives of such types of compounds have been described. It was shown recently in our laboratory [16–19] that thiophosphoryl compounds **1**, with the phosphorus atom bearing the *ω*-haloalkyl moiety, rather easily undergo intramolecular S-alkylation reactions yielding fiveand six-membered 1,2-thiaphosphacyclanium salts **2**. In the case of the presence of at least one alkoxy group at the phosphorus atom, a subsequent dealkylation takes place leading to formation of the 2-oxo-1,2-thiaphosphacyclanes **3** (Scheme 1).

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SCHEME 1

The subsequent functionalization of such compounds, with the introduction of substituents to the ring, represents, however, a rather complicated task. Nevertheless, such an S-alkylation must obviously be a general method of 1,2-thiaphosphacyclane structure formation. Therefore, one should strive to obtain the functional compounds of the said type starting from the functionalized thiophosphoryl compounds having an *ω*-haloalkyl group.

Earlier [20,21], we confirmed this assumption by the synthesis of a few (3-halopropyl)-substituted thiophosphonylacetonitriles and thiophosphinylacetonitriles. In this case, the intramolecular cyclizations proceeding under elevated temperatures, yielded the 2-ethoxy- and 2-methyl-3-cyano-2-oxo-1,2-thiaphosphinanes. To elucidate the limits of the application of the aforementioned synthetic scheme and to estimate the influence of the initial compound structures on the result and stereochemistry of the process, we have synthesized a variety of *ω*-haloalkylsubstituted compounds **5–8** and investigated their intramolecular transformations.

RESULTS AND DISCUSSION

Synthesis of ω-Halo-2 thiophosphorylalkanonitriles

Compounds **5–8**, having a terminal chlorine atom, were obtained as the result of an alkylation of thiophosphorylacetonitriles **4** under phase-transfer catalysis conditions based on the conditions of these reactions investigated by us earlier [22,23] (Scheme 2). Namely, it was established that Cmonoalkylation of compounds **4** with monohaloalkanes and 1,3-bromochloropropane can be carried out in a 50% aq. NaOH/CH₂Cl₂ medium with high a degree of selectivity. The secondary alkylation of C-alkylated thiophosphorylacetonitriles with monohaloalkanes and unsymmetrical *α, ω*dihaloalkanes, in turn, proceeded easily in the heterophase medium s. KOH/CH₃CN. It should be noted that the synthesis of disubstituted thiophosphorylacetonitriles **6–8** is most readily affected without

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isolation of the intermediate alkylation products. The optimal catalyst for both biphasic systems is triethylbenzylammonium chloride (TEBA).

Use of the *α, ω*-dibromoalkanes for an alkylation of monoalkylsubstituted thiophosphorylacetonitriles, under the aforementioned conditions, could afford alkylation compounds having a terminal bromine atom. In this case, however, a side reaction proceeds to give symmetrical bis(thiophosphoryl) alkanedinitriles. For example, the reaction of diethylthiophosphorylpropionitrile with 1,3-dibromopropane to give **7a** is accompanied to a great extent by the formation of the heptanecarbodinitrile **9**, even with substantial dilution of the reaction mixture and with use of an excess of the alkylating agent (Scheme 3). When the said reaction was carried out under the standard dilution conditions suggested previously [23] for unsymmetric *α, ω*-bromochloroalkanes, **9** was found to be the main reaction product (yield 77%).

Compound **9** was obtained as a mixture with the statistical ratio of meso and racemic d,l-isomers and was almost insoluble in hexane. This allowed us to separate it readily from **7a**. Stereoisomers **9** were, in turn, separated by column chromatography. The structure of the meso-form $9(R_c, S_c)$ was confirmed by single crystal X-ray diffraction analysis (Figure 1). The bond lengths and angles in **9** are characterized by the typical values for analogous

FIGURE 1 Molecular structure of meso-**9**. The disordering of OEt groups is omitted for clarity. Selected bond lengths (A): ˚ P(1)-O(2) 1.561(3), P(1)-O(1) 1.571(3), P(1)-C(1) 1.849(3), P(1)-S(1) 1.917(2), P(2)-O(4) 1.562(3), P(2)-O(3) 1.565(3), P(2)- C(5) 1.838(3), P(2)-S(2) 1.915(2); Selected angles (deg.): O(2)-P(1)-O(1) 103.8(2), O(2)-P(1)-C(1) 101.0(2), O(1)-P(1)-C(1) 100.4(2), O(2)-P(1)-S(1) 116.7(2), O(1)-P(1)-S(1) 117.1(1), C(1)-P(1)-S(1) 115.3(1), O(4)-P(2)-O(3) 104.6(2), O(4)-P(2)-C(5) 101.9(1), O(3)-P(2)-C(5) 100.2(1), O(4)-P(2)-S(2) 116.8(1), O(3)-P(2)-S(2) 116.8(1), C(5)-P(2)-S(2) 114.3(1).

compounds [24]. In both of the $S = P(OEt)$ fragments, the $P = S$ group is characterized by the antiperiplanar orientation with respect to the CN group. The slight shortening of $P = S$ bonds, to 1.915(2)A in θ , is probably caused by the electronwithdrawing effect of the cyano group [24].

Yields, elemental analysis data, physicochemical constants, and spectroscopy data of the compounds **5–8** are summarized in Tables 1 and 2. It should be mentioned that, in order to accelerate the intramolecular S-alkylation, compounds with a terminal chlorine atom were transformed, in some cases, to the corresponding iodo-derivatives by the exchange reaction with $NaI/CH₃CN$.

Intramolecular S-Alkylation of ω-Halo-2-thiophosphorylalkanonitriles

Investigation of the intramolecular S-alkylation in a series of *ω*-halo-2-thiophosphorylalkanonitriles **5– 8** (performed by the example of the compounds bearing the diphenylthiophosphinyl group **5f, 7g**) has demonstrated that the presence of the electronwithdrawing *α*-cyano group results in a regular decrease in the thione sulfur atom nucleophilicity, and, consequently, in an increase in *ω*-halo-2 thiophosphorylalkanonitriles stability.

Thus, unlike (3-iodobutyl)diphenylphosphine sulfide, which readily undergoes spontaneous intramolecular S-alkylation yielding diphenyl 1,2*λ*5 thiaphosphinanium iodide [16], during the preparation from (3-chlorobutyl)diphenylphosphine sulfide by the exchange reaction with sodium iodide, its cyanosubstituted analogs **10a** and **11a** are stable and may be isolated. The corresponding cyclic forms **12** and **13** were formed slowly in acetonitrile solutions of **10a** and **11a**, the equilibrium between linear and cyclic forms (ring-chain halotropic tautomerism) being established in about six months (Scheme 4). After the equilibrium had been reached, the amounts of cyclic forms were found to be 14% (**12**) and 9% (**13**), according to the 31P NMR data.

However, due to the subsequent irreversible dealkylation in the intermediate thiaphosphacyclanium salts, the cyclization of the initial compounds **5–7** proceeds rather readily, even for the compounds having a terminal chlorine atom, when there is at least one alkoxy group bonded to the phosphorus atom. Thus, **5–7** are only partly transformed to the five- and six-membered 3-cyano-2-oxo-1,2 thiaphosphacyclanes **14** and **15**, even under distillation *in vacuo* (Scheme 5, method I). Hence, although the introduction of the cyano group decreases the thione sulfur atom nucleophilicity, it promotes, at the same time, the dealkylation process in the intermediate thiaphosphacyclanium salt. It should be noted that the substituted thiophosphorylacetonitrile **8a**, bearing the 4-chlorobutyl moiety

$$
\begin{array}{ccc}\n\text{Ph}_{2}\text{P(S)C(R)CN} & \xrightarrow{\text{NaLCH}_{3}CN} & \text{Ph}_{2}\text{P(S)C(R)CN} \\
\downarrow & & \downarrow \\
\text{CH}_{2}\text{B} & & \downarrow \\
\text{St, 7g} & & 10a, 11a & & 12, 13 \\
& R = H (10a, 12); R = \text{Me (11a, 13)}\n\end{array}
$$

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TABLE 1 Physical and Elemental Analyses Data for Compounds **5–9, 14** and **15**

Compound	Yield ^a	$m.p./°C$ or $b.p./°C$ (mm Hg)		Found (%) Calculated (%)				
	(%)		Formula	$\cal C$	H	Ν	\boldsymbol{P}	
$5a^b$	83 (46)	Oil	C_9H_{17} CINO ₂ PS	40.11	6.28	5.35 5.19	11.58	
5b ^b	70 (41)	Oil	$C_{11}H_{21}CINO2PS$	40.08 43.67	6.35 7.13	—	11.48 10.12	
5c	82 (60)	$170 - 175(1)$	C_9H_{17} CINOPS	43.37 43.88	7.11 6.84	$\overline{}$ 5.45	10.40 12.23	
$5d^b$	65 (40)	Oil	$C_{10}H_{19}$ CINOPS	43.60 44.31	6.75 7.20	5.52 5.35	12.20 11.23	
$5e^b$	87(61)	Oil	$C_{13}H_{17}$ CINOPS	44.86 51.94	7.15 5.87	5.23 5.06	11.57 10.16	
5f ^b	75 (55)	88-89	$C_{17}H_{17}CINPS$	51.74 62.68	5.64 4.98	4.64 4.29	10.28 9.26	
6a ^b	93 (70)	Oil	C_9H_{17} CINO ₂ PS	62.17 40.55 40.07	5.10 5.87	4.20 5.38 5.19	9.29	
$6b^b$	80 (69)	Oil	$C_{13}H_{17}$ CINOPS	51.88 51.74	6.31 5.42 5.64	4.83 4.64	$\qquad \qquad -$ $\qquad \qquad -$ $\qquad \qquad -$	
$\mathbf{6c}^b$	71 (60)	Oil	$C_{14}H_{19}$ CINOPS	53.18 53.25	5.93 6.02	4.68 4.44	$\overline{}$ $\qquad \qquad -$	
$7a^c$	65 $(57)^d$	$165 - 175(1)$		$\overline{}$			$\qquad \qquad -$	
7b	75 (63)	$155 - 160(1)$	$C_{12}H_{23}CINO2PS$	47.10 47.22	7.93 7.43	\equiv	\equiv	
$7c^c$	85 (50)	$150 - 165(1)$	$C_{12}H_{23}CINO2PS$	46.16 46.23	7.39 7.38	4.96 4.49	9.85 9.95	
7d	80(64)	$170 - 175(1)$	$C_{10}H_{19}$ CINOPS	47.19 46.89	7.69 7.51	—	11.31 10.99	
$7e^b$	90(68)	Oil	$C_{11}H_{21}$ CINOPS	47.10 46.89	7.69 7.51		11.31 10.99	
7f ^b	100 (83)	Oil	$C_{14}H_{19}$ CINOPS	53.22 53.24	6.03 6.06	4.36 4.44	9.55 9.81	
$7g^b$	83 (65)	$65 - 66$	$C_{18}H_{19}CINPS$	62.30 62.15	5.50 5.50	4.30 4.03	8.80 8.90	
$8a^e$	92(73)	$150 - 195(0.1)$						
8b	71 (59)	87-89 (Petrol. ether)	$C_{19}H_{21}$ CINPS	62.96 63.06	5.77 5.85	3.68 3.87	$\qquad \qquad$	
9		Oil (meso: $D, L = 30:70$) 94-95 (meso: $D, L = 70:30$) 93 (hexane, meso)	$C_{17}H_{32}N_2O_4P_2S_2$	44.93 44.93	6.91 7.05	6.05 6.17	$\overline{}$	
14b		166-167 (A:B = 94:6)	$C_{11}H_{12}NOPS$	55.58	5.08	5.81		
14 _c	61 (30) 95 (54) Quant. (57)	$108 - 109$ (B) 101-102 (A) $71 - 72$ (B)	$C_{12}H_{14}NOPS$	55.70 57.31 57.37	5.06 5.69 5.58	5.91 5.62 5.58		
15a	75 (51)	65-70 ($\mathbf{A}:\mathbf{B} = 50:50$)	$C_7H_{12}NO_2PS$	40.60	5.93	6.70	15.12	
15 _b	55 (21) 32(17)	116–118 (B) 73-74 (A:B = 50:50) ^{$\frac{1}{2}$}		40.96	5.89	6.83	15.09	
15 _c	30(18)	136–137 (A)	$C_6H_{10}NOPS$	41.73	5.72	7.78		
15d	53 (25) 44 (21) 79 (34)	$132 - 134$ (B) $111 - 112(A)$	$C_{11}H_{12}NOPS$	41.14 55.56 55.70	5.71 5.06 5.06	8.00 5.70 5.91		
15e	29 (10)	$80 - 81$ (B)	$C_9H_{16}NO_2PS$	46.36	7.08	5.99	13.05	
15f	19(6) Quant. (24)	$134-136$ (A:B = 84:16)	$C_{12}H_{14}$ NOPS	46.35 56.44 56.37	6.87 5.82 5.58	6.01 5.44 5.58	13.30 11.77 12.35	
15g 15h	15(6) 64 (22)	Oil (A:B = 64:36) ^g $151 - 152$ (B)	$C_7H_{12}NOPS$	44.35	6.62	$\qquad \qquad \qquad$ 7.31	16.09	
15i	75 (45)	$94 - 95$ (A:B = 70:30) $107-108$ (A:B = 5:95)	$C_8H_{14}NO_2PS$	44.44 43.62 43.83	6.35 6.36 6.50	7.41 6.43 6.39	16.40	

aAccording to ³¹P NMR data (shown in parenthesis is the product yield after purification); for compounds **14** and, **15** the yield is based on the both diastereomers; shown first is the yield for chloroderivatives cyclization, the second is that for iododerivatives cyclization; for purification conditions, see Experimental section. bPurified by column chromatography (silica gel 40-100 μ m, hexane—acetone 100:4).

^cPurified by distillation and column chromatography.

 ${}^{d}86%$ Purity according to 1 H NMR data.

^eA 85:15 mixture of compounds having terminal chlorine and bromine atoms according to ¹H NMR data.

 f 87% purity according to ¹H NMR data.

 986% purity according to 1 H NMR data.

TABLE 2 Selected 31P, 1H, and 13C NMR and IR Data for Compounds **5–8**

¹³C NM_h

⁶L

⁶L

R¹R²P(S)-C-CN

CH₂(CH₂)_{n-2}CH₂Cl

<u>2</u>

<u>2</u>

<u>1</u>

5

TABLE 2 (continued) Selected ³¹P, ¹H, and ¹³C NMR and IR Data for Compounds 5–8
 $-\frac{6!}{-C}$

^aSolvent CDCl3; 1H NMR (**5a–e; 6a–c; 7a–d,f,g; 11a**); NMR 13C (**5a–c,e; 6a–c; 7a–d**).

^bSolvent C6D6; 1H NMR (**5f, 7e**); NMR 13C (**5d,f , 7e,f,g**).

^cTwo diastereomers **A:B** ⁼ 1:1 (**5c,d**), 60:40 (**5e**), 57:43 (**6b**), 64:36 (**6c**), 1:1 (**7d**), 65:35 (**7e**), 64:36 (**7f**).

 d In CH₃CN the diastereomers signals are interchanged (δ _P(**A**) 84.71, δ _P (**B**) 84.82).

^eThe compound having bromine terminal atom.

 f The proton P–C–CH₂ (Pr) and CH₂CH₂CH₂CI signals are overlapped.

gSolvent CD₃CN for ¹H, ³¹P, ¹³C NMR, spectrum is shown for the corresponding chlorine derivative. h Solvent CH₂Cl₂.</sup>

SCHEME 5

(the cyclization of which could afford only a sevenmembered ring), does not cyclize under the aforementioned conditions. This can apparently be explained by the lesser thermodynamic stability of the said ring compared to the corresponding five- and six-membered analog.

The yield of 3-cyano-2-oxo-1,2-thiaphosphacyclanes **15** under distillation *in vacuo* depends mainly on the easiness of dealkylation for the RO group (Table 3). In other words, it decreases with an increase of the volume of the radical (compare **15a** and **15b**) and also when passing from a primary radical to a secondary one (compare **15c**, obtained from **5c** and **5d**). Furthermore, the yields of the six-membered heterocycles are decreased drastically (up to 5–7%) with the insertion of the alkyl radical into the *α*-position to the phosphorus (compare **15b** and **15e**). Even **7a**, bearing a terminal bromine atom, transforms under distillation to the cyclic product with a yield of 7% only. Compounds **6**, when distilled, afforded the 3-alkyl-3-cyano-2-oxo-1,2 thiaphospholanes **14** far more readily and in considerably higher yields than observed for six-membered compounds, **15**, even in spite of the presence of an alkyl radical in the α -position to the phosphorus in the initial compounds **6a,c**. This fact corresponds to the general rule that five-membered rings are formed more readily than six-membered ones [25– 27]. It should be mentioned that addition of catalytic amounts of TEBA to the initial compounds **5–7** increases the yields of the target compound **14** and **15**.

As one could expect, the intramolecular cyclization of the *ω*-iodoalkyl-substituted thiophosphinates and thiophosphonates **10** and **11** obtained from the corresponding chloroderivatives **5–7** in situ (with use of 2 equiv. NaI/CH_3CN and with 2 hours reflux, method II) proceeds under milder conditions. Intramolecular S-alkylation proceeds slowly even at room temperature. Thus, cyclic derivative **15c** is formed in 62% yield from the iodo derivative **5d** $(R^1 = Me, R^2 = Bu - i, R^3 = H, n = 3)$ during one year at 20◦ C. Upon refluxing **5–7** in acetonitrile for 2–20 hours, 2-alkyl(phenyl)-3-cyano-2-oxo-1,2-thiaphosphacyclanes **14** and **15** are

TABLE 3 Dependence of Yields and Diastereomer Ratio of 3-Cyano-2-oxo-1,2-thiaphosphacyclanes **14** and **15** on the Initial Substrate Structure and Cyclization Method

Substrate 6a		Yield 14 and 15, $%$ (³¹ P NMR), Diastereomer Ratio						
	Thiaphosphacyclane		Method I Distillation in vacuo	Method II Nal/MeCN, Reflux				
	14a	19 40 ^c	A > B(60:40)	quant ^a	b			
6b	14b	b	b	61	A < B(40:60)			
6c	14с	95	$A > B$ (70:30)	quant.	$A < B$ (30:70)			
5a	15a	75	$A \approx B$	quant. ^d				
5b	15b	55	$A \approx B$	32	$A \approx B$			
5c	15c		A > B(60:40)	83	A < B(40:60)			
5d	15c	30	A > B(60:40)	53	A < B(40:60)			
5e	15d	44 ^c	$A > B$ (55:45)	79	$A \approx B$			
7a	15i			75^e	$A < B$ (40:60)			
7b	15e	5.6 17 ^c	A > B(60:40)	29	A < B(40:60)			
7c	15g	15	A > B(60:40)	30	$A < B$ (30:70)			
7e	15h			64	$A \approx B$			
7f	15f	19	$A > B$ (55:45)	quant.	A < B(40:60)			

^aCyclic sodium salt **16**.

 b The experiment was not carried out.

c Triethylbenzylammonium chloride addition (cat.) upon distillation.

^dCyclic sodium salt **17**.

e Cyclization of the compound having terminal bromine atom upon refluxion in the acetonitrile solution.

SCHEME 6

formed in yields up to quantitative (Scheme 6, Table 3).

It should be noted that cyclization of the thiophosphonates **5a** and **6a** occurred under such conditions (with an excess of NaI), followed by dealkylation of the second alkoxy group at the phosphorus atom in the desired 1,2-thiaphosphocyclanes **14a** and **15a**. As a result, the sodium salts of the corresponding cyclic acids **16** and **17** were obtained in practically quantitative yields. In the case of **8a**, this cyclization method resulted in a polymer product of nonidentified structure schemes.

To obtain 3-cyano-2-ethoxy-3-methyl-2-oxo-1,2 thiaphosphinane, **15i**, which is formed under thermal cyclization of the initial chloride in low yield (7%) and apparently could not be obtained from the corresponding iodo derivative (see Scheme 7), we have used S-alkylation of the corresponding *ω*-bromo-2-thiophosphorylalkanonitrile proceeding under reflux in the acetonitrile solution schemes.

$$
\underbrace{(EtO)_2P(S)C(R^3)CN}_{(CH_2)_\Pi} \xrightarrow[\text{NaL/CH}_3\text{CN}]{} \xrightarrow[\text{NaCl}]{\text{NaL/CH}_3\text{CN}]{} \xrightarrow[\text{CH}_2)_\Pi} \xrightarrow[\text{CH}_2)_\Pi} \xrightarrow[\text{CH}_2)_\Pi
$$

SCHEME 7

SCHEME 8

The presence of two asymmetric centers phosphorus and carbon C(3) atoms—in the 3-cyano-2-oxo-1,2-thiaphosphacyclanes determines their formation as a mixture of two diastereomers (**A** and **B**), each being a racemic mixture of the enantiomers. In the $31P$ NMR spectrum (Table 4), two singlets correspond to these diastereomers **A** and **B** (with those denoted by diastereomer **A** showing a lowfield chemical shift). The ratio of the diastereomers formed during the cyclization depends on the initial compound structure (mainly by the presence of the alkyl substituent in the α -position), the method of cyclization and the ring size. The influence of these factors on the diastereomeric composition of 3-cyano-2-oxo-1,2-thiaphosphacyclanes **14** and **15** obtained is shown in Table 3.

For 2-alkoxy-3-cyano-2-oxo-1,2-thiaphosphinanes, **15a,b**, without an alkyl substituent in the 3 position, the diastereomeric ratio is 1:1, while for their 3-alkylsubstituted analog, **15e–i**, the proportion of diastereomer **A** is generally higher when the cyclization is carried out by distillation *in vacuo* (Hlg = Cl), and diastereomer **B** is formed in a greater amount under milder conditions (when cyclizing 5 iodo(bromo)-2-thiophosphorylpentanonitriles). The same regularities may be observed for 3-cyano-2 oxo-1,2-thiaphospholanes, **14**. The preferential formation of one or another diastereomer obviously depends on steric and electronic factors determining the structure of the intermediate phosphonium salt and conditions of the cyclization as well.

The compounds **14** and **15** were isolated from the reaction mixtures as diastereomer mixtures (e.g., by precipitation using ether). These mixtures were resolved into the individual diastereomers in most cases by column chromatography and fractional crystallization. Satisfactory elemental analysis data were obtained both for the diastereomeric mixtures isolated and for the individual isomers (Table 1). The main spectral parameters of the heterocycles synthesized (1 H, 13 C, 31 P NMR and IR) are listed in Tables 4–6.

In the IR spectra, the absorption bands characteristic of $P = 0$ and CN groups are observed in the expected areas, and the difference for diastereomers **A** and **B** does not exceed 10 cm−¹ (Table 5).

In the 31P NMR spectra (Table 4), 1,2-thiaphospholanes 14 show signals in CH_2Cl_2 in the $\delta = 61-73$ ppm region. For compounds 15 the range of the chemical shift values in CH_2Cl_2 is 35–44 ppm for 2-alkoxyderivatives **15a,b,e,g**, 41–52 ppm for 2-methylsubstituted compounds **15c,h**, and 32–44 ppm for 2-phenylsubstituted analogs **15d,f**. Note that, as for the initial linear compounds **5– 7**, the introduction of an alkyl substituent into the

Solvent Compound	CH ₃ CN			CH ₂ Cl ₂			CDC ₃		C_6H_6			
	δ (A)	δ (B)	$\Delta\delta$	δ (A)	δ (B)	$\Delta\delta$	δ (A)	δ (B)	$\Delta\delta$	δ (A)	δ (B)	$\Delta\delta$
14a				69.62	68.12	1.50						
14b	74.20	68.61	5.59	71.79	65.56	6.23						
14 _c	73.83	66.14	7.69	72.21	61.79	10.42	72.71	65.11	7.60	70.62	62.87	7.75
15a	36.49	36.19	0.30	35.89	35.58	0.31	36.65	36.21	0.44	34.92	34.05	0.87
15 _b	37.06	36.58	0.48	36.00	35.55	0.45	36.54	36.16	0.38			
15c	44.98	44.61	0.37	42.64	41.91	0.73	41.99	39.38	2.61	39.36	37.42	1.94
15d	35.44	34.56	0.88	35.05	34.29	0.76	35.15	34.57	0.58	$\overline{}$		
15e	44.04	41.43	2.61	43.56	41.00	2.56	43.88	41.38	2.50			
15f				43.98	40.05	3.93	44.51	40.54	3.97			
15g				43.28	40.93	2.35	43.69	41.43	2.26			
15h	52.48	47.64	4.84	51.67	47.45	4.22				51.43	45.50	5.93
15i	43.70	41.04	2.66				43.93	41.51	2.42			

TABLE 4 Chemical Shift Values (*δ* ppm) in 31P NMR Spectra for 3-Cyano-2-oxo-1,2-thiaphosphacyclanes **14** and **15** (**A** and **B** diastereomers) in Different Solvents

six membered heterocycles results in a considerable downfield shift (*δ* 7–9 ppm).

The difference between the chemical shifts of the 1,2-thiaphosphacyclanes **A** and **B** diastereomers $(\Delta \delta)$ in the ³¹P NMR spectra depends on the ring size, the presence of an alkyl substituent in the 3-position, and the solvent (Table 4). All other factors being the same, $\Delta \delta$ is generally higher for the compounds with a methyl or phenyl group bonded to the phosphorus atom. Furthermore, for the initial compounds **5–7** having two asymmetric centers, the change of the solvent often results in the change of the mutual signal positions of diastereomers **A** and **B** in 31P NMR spectra, while for **14** and **15**, the mutual position of stereomers **A** and **B** remains unchangeable independently of the solvent in use, which is easy to follow in comparing their integral intensities.

In its 1H NMR spectra, 3-cyano-2-oxo-1,2 thiaphosphacyclanes show the characteristic $SCH₂$ signal at the $\delta = 2.80-3.60$ region, which in some cases appears as an overall complex multiplet for both protons, and, for **14b,c**, and **15c-e,h**, two separate multiplets are observed corresponding to the protons in the *trans*- and *cis*-positions to the phosphoryl group oxygen atom. For 3-cyano-2 oxo-1,2-thiaphosphinanes, **15a–d**, the signal of the methine proton at the *α*-position to the phosphorus atom appears, as a rule, as a doublet of doublets of doublets, as it does for the initial compounds, **5**, but in lower field compared with the latter ones (Table 5).

In the 13C NMR spectra of **14** and **15** (Table 6) the signals of the carbon atoms C(4) [28] bonded to the cyano-group are placed at $\delta = 31.49 - 34.09$ for 3-cyano-2-oxo-1,2-thiaphosphinanes **15a–d** and are shifted downfield (36.87–43.99) for their 3-alkylsubstituted analogs **15e–i**. In the case of the fivemembered compounds **14**, the signals of the C(4) atom are placed at $\delta = 43.31 - 50.11$. It should be noted that, in six-membered ring compounds, the above C(4) signals are shifted upfield about 2–5 ppm compared to the P–C(CN) signal in the initial linear *ω*-haloalkyl-substituted thiophosphorylacetonitriles **5** and **7**. On the contrary, in the five-membered rings **14**, the signals of the C(4) atom are shifted about 3– 10 ppm downfield with respect to the P–C(CN) signal in the initial **6**. Furthermore, the decrease of ${}^{1}J_{PC}$ (4– 17 Hz) is noted in the cyclic compounds compared to the initial compounds **5–7**. This fact is explained logically by the change in the surroundings of the phosphorus atom attached to the aforementioned carbon atom (from $R^1(RO)P(S)$ - to $R^1(O)PSR$ -). It should be mentioned that ${}^{1}J_{\text{PC}}$ is generally less in case of the **A**-isomer both for five- and six-membered rings. In 1,2-thiaphosphinanes, the signal of the cyclic $C(1)$ atom attached to sulfur appears as a doublet with ² J_{PC} 3.7–5.9 Hz at δ = 24.85–29.78 for **15a–d** and at δ = 33.45–37.98 for their alkyl-substituted analogues **15e–i**. For 1,2-thiaphospholanes **14b,c**, the doublet of the C(1) atom is placed in the same region as for their 6-membered analogues, but with the $^{2}J_{\text{PC}}$ coupling constant increasing to 7.6–8.5 Hz. This constant is maximal for the cyclic salt **16** (9.2 Hz). Thus, alkyl substituent insertion into the *α*-position to the phosphorus atom results in a downfield shift of C(4), $C(1)$, and $C(6)$ signals. The signal of the CN-group carbon atom is located in the characteristic region of δ = 115–120, appearing usually as a doublet for the **A**-isomer and as a singlet for the **B**-isomer. It should be noted that the signal of this group for the initial compounds **5–7** usually is situated in the same *δ* region as the doublet with ${}^2J_{\text{PC}}$ coupling constant equal to 4.4–8.0 Hz.

TABLE 5 Selected 1H NMR and IR spectra parameters for 3-cyano-2-oxo-1,2-thiaphosphacyclanes **¹⁴** and **¹⁵**

^aTwo conformers.

 b The signals of OCH₂ protones of **A** and **B** diastereomers are overlapped.

^cThe signals of C1H2 protones of **A** and **B** diastereomers are overlapped. ^d**A** and **B** diastereomers mixture in 63:37 (**15b**), 29:71 (**15d**), 64:36 (**15g**), 5:95 (**15i**) ratio.

 ${}^e\mathsf{T}$ he signals of C²H₂ and C³H₂ protones of **A** and **B** diastereomers are overlapped. f Solvent CD ${}_3$ CN.

^gThe signals of minor diastereomer **B** are overlapped by signal of major isomer **A**.
"Solvent DMSO-d₆.

 $(\tilde{CH}_2)_{n-2}$

^aSolvent CD₃CN.
^bSolvent DMSO-d₆.

The structures of the series of the individual diastereomers **14** and **15** have been confirmed by the Xray diffraction analysis. Important bond lengths and bond angles are shown in Table 7. According to the X-ray data, five-membered rings in **14** adopt the twist (deviation of the $C(2)$, $C(4)$ atoms) and the envelope conformation (deviation of the C(2) atom) (Figures 2 and 3), while all the six-membered rings, both reported herein and described by us previously [20,21], are characterized by chair conformations (Figures 4–6), with the axial position for the CN group in all molecules **14** and **15**. The phosphorus atoms are characterized by a slightly distorted tetrahedral configuration with the endocyclic $C(4)P(1)S(1)$ angles in the ranges of $95.84(4) \div 97.87(3)$ [°] in five-membered rings **14** and $103.43(7) \div 105.79(5)°$ in six-membered rings **15**. The torsion angles $C(5)C(4)P(1)O(1)$ are in the interval −35.0 to 55.2◦ for molecules having the equatorial position of the $P = O$ group (synclinal conformation, *cis*-isomer), and from 147.6 to 170.6◦ for the axial $P = O$ group (antiperiplanar conformation, *trans*-isomer).

The bond lengths and bond angles in fiveand six-membered rings exhibit the expected values (Table 7). The P–S bond lengths in 1,2-thiaphosphinanes **15** (2.044(2) \div 2.065(2)Å) are slightly shortened in comparison with the typical single P–S bond $(2.08 \div 2.10\text{\AA})$ [24] and are close to those observed earlier in the series of $1,2\lambda^5$ -thiaphosphinanium salts (2.05\AA) [18,19]. Such similarity of thiaphosphinanium salts and cyanosubstituted cycles **14** and **15** means that the phosphorus atom in the latter compounds is characterized by a significant fractional positive charge, mainly due to the electronwithdrawing effect of both the CN group and the phosphoryl oxygen. The same conclusion can be made on the base of comparison of the P–S bond lengths in the nonfunctionalized 2-oxo-2-phenyl-1,2 thiaphosphinane [29] and **15d(A)** (Figure 5), where the P–S bond length is shortened by $0.02A$, while the $P(1)$ –C(1) bond is elongated by 0.04A due to the presence of the CN group (Table 7). Thus, one can conclude that, for the P–S bond, the presence of electronwithdrawing substituents leads not to an elongation

	14c(A)	14c(B)	15a(B) a	$15c(A)^a$	15c(B)	15d(A)	15i(B) a	15h(B) b	$\mathbf{G}^{a,c}$
$P(1) - S(1)$	2.0778(9)	2.0751 $(4)^d$ 2.0832(4)	2.044(2)	2.062(2)	2.065(2)	2.059 $(1)^d$ 2.061(1)	2.0558(7)	2.054(1)	2.082(1)
$P(1) - O(1)$	1.478(2)	1.4805(8) 1.4817(8)	1.456(3)	1.488(2)	1.484(2)	1.479(2) 1.477(2)	1.469(1)	1.470(4)	1.474(2)
$P(1) - R_1$	1.793(2)	1.800(1) 1.801(1)	1.573(3)	1.785(3)	1.788(2)	1.797(2) 1.801(2)	1.572(1)	1.785(4)	1.801(2)
$P(1)-C(4)$	1.857(2)	1.869(1) 1.873(1)	1.817(4)	1.838(4)	1.833(2)	1.839(2) 1.832(2)	1.836(1)	1.852(2)	1.790(3)
$S(1) - C(1)$	1.837(2)	1.840(1) 1.834(1)	1.807(6)	1.835(3)	1.845(3)	1.838(3) 1.827(3)	1.832(2)	1.837(3)	1.828(3)
C(4)P(1)S(1)	97.29(8)	95.84(4) 97.87(3)	105.5(1)	103.9(1)	103.43(7)	103.45(8) 103.56(8)	105.79(5)	104.62(9)	103.4(1)
O(1)P(1)R ¹	112.2(1)	117.73(5) 110.84(5)	116.7(2)	114.6(1)	114.1(1)	113.8(1) 113.6(1)	116.56(8)	113.6(4)	113.0(1)
C(1)S(1)P(1)	95.7(1)	94.93(4) 95.20(4)	105.5(1)	97.9(2)	100.13(7)	99.1(1) 97.96 (9)	99.54(8)	100.2(1)	96.8(1)
C(5)C(4)P(1)O(1)	-35.0	165.4 147.6	170.6	170.6	50.0	167.9 170.00	55.2	50.5	n/a
$Lp_S S(1)P(1)X^e$	118	-148 -124	175	176	177.1	173 175	175	176	176
Conformation ^T	Twist $C(2)$, $C(4)$ $(0.52, -0.11E)$	Twist $C(2)$, $C(4)$ $(0.69, -0.13E)$ Envelope C(2) (0.64E)	chair	chair	chair	chair	chair	chair	chair

TABLE 7 Important Bond Lengths (A) and Bond Angles (deg.) for Some Representatives of ˚ **14** and **15**

ªThe numbering of atoms in **15a(B), 15c(A), 15i(B),** and **G** is the same as for all other six-membered cycles in Figures 2–6.
^bFor **15h(B)** the bond lengths and angles are presented only for the molecule with the occupa

c**G** presents 2-oxo-2-phenyl-1,2-thiaphosphinane [29].

^dFor molecules **14c(B)** and **15d(A)** bond lengths and angles are indicated for two independent molecules.

^eThe torsion angle between the hypothetical sulfur electron lone pair (Lp_s) and the phosphorus axial subsistent (X is O(1) in diasteromer A and R¹ in **B**). The positions of Lp_s were calculated in the assumption of the tetrahedral arrangement of Lp_S 's and $C(1)$, $P(1)$ atoms.

f For five-membered cycles the value in brackets corresponds to the deviation of the atom from the mean plane of remaining atoms of the cycle.

FIGURE 4 Molecular structure of **15c(B)**.

FIGURE 2 Molecular structure of **14c(A)**.

FIGURE 3 Molecular structure of **14c(B)**. Only one of two independent molecules is shown. The differences in their conformations are described in Table 7.

but to a shortening of the bond that is usual for the polar or ionic bonds.

Besides the inductive effects of the substituents, variation of the P–S bond can also be affected by the interaction between the lone pair (Lp_s) of the sulfur atom and the antibonding *σ*-orbital of the axial P–X bond (n- σ^* interaction), which differs for P=O, OAlk, and Alk groups [30]. The increase of a n-*σ*[∗] interaction must be reflected in the shortening of the P–S endocyclic bond and elongation of the axial exocyclic one.

Analysis of the important bond lengths and bond angles in the six-membered rings indicates clearly

FIGURE 5 Molecular structure of **15d(A)**. Only one of two independent molecules is shown.

FIGURE 6 Molecular structure of **15h(B)**. The disordering is omitted for clarity.

some influence of the n-*σ*[∗] interaction in the case of the OEt axial group in **15a(B)**, where not only the P– S bond attains the minimum value $(2.044(2)$ A), but also the $S(1)$ -C(1) bond is the shortest one in the series of compounds studied $(1.807(6)\text{\AA})$. In spite of pronounced shortening of the P–S bond in **15a(B)**, introduction of the Me group in the 3-position of the ring (compound **15i(B)**) causes an increase of the P–S bond length by 0.01\AA , annulling, in fact, the influence of the n-*σ*[∗] interaction. It's noteworthy that the formal strength of the n- σ^* interaction, in accordance with the values of the pseudotorsion angle L_p _sS(1)P(1)X and the length of the P(1)–O(2)(Et) bond in **15a(B)** and **15i(B)**, were found to be the same.

Opposite to the axial alkoxy group at the phosphorus atom, the influence of the $n-\sigma^*$ interaction on the P–S bond is insignificant in the case of an axial $P = 0$ bond. For example, as it may be seen from Table 7, the $P(1)$ –O(1) and $P(1)$ –S(1) bonds in **15c(A)** and **15c(B)** are similar. The same situation is observed in the case of the 3-cyano-3-ethyl substituted 5-membered rings where the $P(1)$ –S(1) and P(1)–O(1) bonds in **14c(B)**, with the axial position of the $P = O$ group are close to the corresponding values in $14c(A)$ with the equatorial $P = 0$ bond.

For the $1,2\lambda^5$ -thiaphosphacyclanium salts, the elongation of the P–S bond when passing from the five-membered rings to the six-membered ones $(2.0751(4) \div 2.0832(4)$ Å) was not observed [18]. Therefore, taking into account the aforementioned influence of the alkyl substituent in the 3-position on the P–S bond length in **15a(B)** and **15i(B)**, one can suppose that the difference $(0.019A)$ between the P-S bond lengths in **14c(B)** and **15d(A)** is also caused mainly, not by the ring size change, but by introduction of the 3-ethyl substituent decreasing the fractional positive charge on the phosphorus atom.

Summarizing, the P–S bond length in 1,2 thiaphosphacyclanes is more affected by the inductive effects of the substituents at the phosphorus atom and in the 3-position of the ring than by the n-*σ*[∗] interaction.

Comparison of the $31P$ NMR spectra and X-ray diffraction data allows one to conclude that for the 3-cyano-2-oxo-1,2-thiaphospholanes **14**, the isomer **A** (having a downfield chemical shift) is characterized by the synclinal conformation of the cyano group and phosphoryl oxygen atom (*cis*-isomer) with the identical configuration of the chiral centers (2*R*[∗] ,3*R*[∗]), while diastereomer **B** is characterized by an antiperiplanar disposition of the said groups (*trans*-isomer) with the opposite configuration of the chiral centers (2*R*[∗] ,3*S*[∗]). For the 2-

SCHEME 9

oxo-1,2-thiaphosphinanes, **15**, the situation changes to the opposite one: diastereomer **A** with a downfield shift in its 31P NMR spectrum corresponds to the *trans*-disposition of the cyano group and phosphoryl oxygen atom with the opposite configuration of the chiral centers (2*R*[∗] ,2*S*[∗]), while the signal of the *cis*-isomer **B** with the identical configuration of the asymmetric centers is shifted upfield. The only exception is the 3-cyano-2-oxo-2-phenyl-1,2-thiaphosphinane **15f**, wherein the alteration of substituent precedence leads to the identical configuration of the chiral centers in the *trans*-**A** isomer and to the opposite one in the *cis*-**B** isomer. Scheme 9 illustrates stereoisomer **14** and **15** structures in comparison with δ^{31} P.

Thus, 31P NMR spectroscopy allows one to assign unambiguously the type of geometric isomer and the configuration of the asymmetric centers in 3-cyano-2-oxo-1,2-thiaphosphinanes **14** and **15**. To determine the stereochemical structures of phosphacyclanes and 1,2-oxaphosphacyclanes having a tetracoordinated phosphorus atom 1H NMR spectra are commonly used [1,31]. This method of determination is based on the deshielding effect of the P = 0 bond on all kinds of protons in the *cis* position to the $P = O$ group. Undoubtedly, this anisotropic effect allows the assignement of the *cis* and *trans* arrangements of the $P = O$ group and CH-CN cyclic proton for the compounds **15a–d** or methyl [P– $C(CH₃)CN-$] or methylene $[P-C(CH₂-R)CN-]$ group in 3-alkylsubstituted compounds, **14a–c** and **15e–i:** $\delta_{\rm H}$ *trans*-isomer > $\delta_{\rm H}$ *cis*-isomer [32]. Nevertheless, ³¹P NMR spectroscopy is more preferable to determine the geometric structure of **14** and **15** as

having no need for chemically pure samples and allowing one to monitor the crude reaction mixtures. We believe that regularities described previously for the disposition of signals of *cis* and *trans* 2 oxo-1,2-thiaphosphacyclanes in ³¹P NMR spectra have to be similar in passing to other 3-substituted 1,2-thiaphophacyclanes, provided that the 3 substituent will represent an electron-withdrawing group.

CONCLUSION

Thus, using as an example the *ω*-haloalkylsubstituted thiophosphorylalkanonitriles, the intramolecular Salkylation has proven to be the general principle of the 1,2-thiaphosphacyclane ring construction which opens wide opportunities to synthesize different hitherto elusive 1,2-thiaphosphacyclanes, including functionalized ones.

EXPERIMENTAL

General

NMR spectra were recorded on Bruker WP-200SY and AMX-400 spectrometers in CH_2Cl_2 , CDCl₃, C₆D₆, $CD₃CN$ solutions using a deutero solvent as an internal standard (1 H, 13 C) and 85% H₃PO₄ as an external standard. IR spectra were recorded on a UR-20 spectrophotometer in a thin layer, in KBr pellets, or in vaseline oil. The starting thiophosphorylalkanonitriles **4** were obtained by the reaction of phosphorylacetonitriles with the Lawesson reagent according to the literature [33].

X-Ray Crystal Structure Determination

The crystallographic data for **9, 14c(A** and **B), 15c(B), 15d(A)** and **15h(B)** are presented in Table 8. All structures were solved by the direct method and refined by full-matrix least squares against F^2 in the anisotropic (H-atoms isotropic) approximation using the SHELXTL-97 package. Analysis of the electron density synthesis in **9** and **15h(B)** has revealed disordering of molecules in crystals. In the stucture of **9,** OEt groups were disordered by two positions, while in **15h(B)** the disordering was caused by the superposition of two molecules with the occupancies 0.8 and 0.2, respectively. The disorder in **15h(B)** with the same ratio of occupancies was observed for two independent single crystals. All hydrogen atoms in **9** and **14–15** were located from the electron density difference synthesis and were included in the refinement in isotropic approximation.

Supplementary Material

Crystallographic data, namely atomic coordinates, anisotropic parameters, a full list of the bond lengths, bond and torsion angles, and F_0/F_c values (excluding structure factors), reported in this article are given in the supplementary materials and have been deposited as supplementary nos. CCDC-154959 (**9**), CCDC-154958 (**14c(A)**), CCDC-154955 (**14c(B)**), CCDC-154956 (**15c(B)**), CCDC-154957 (**15d(A)**), and CCDC-154954 (**15h(B)**) at the Cambridge Crystallographic Data Centre. Copies of the data may be obtained free of charge on application to the CCDC,12 Union Road, Cambridge CB2 1EZ UK (Fax: [internat.] +44-1223/336-033; E-mail: deposit @ccdc.cam.ac.uk).

5-Chloro-2-thiophosphorylpentanonitriles (5): General Procedure

To a stirred solution of thiophosphorylacetonitrile **4** (10 mmol) and TEBA (200 mg) in CH_2Cl_2 (based on 15 mL of solvent for the solid **4f** and 5 mL for the liquid **4a–e**) NaOH (1.6 g, 40 mmol) was added as a 50% aqueous solution. To the reaction mixture, 1,3-bromochloropropane was added dropwise over 10 minutes. The reaction mixture was stirred for 3 hours at 20◦ C and diluted with water (15 mL); the organic layer was separated, and the aqueous layer was extracted with $CH₂Cl₂$. The organic layer was dried $(Na₂SO₄)$ and concentrated *in vacuo*, and the residue was subjected to column chromatography (silica gel $40-100 \mu m$; hexane–acetone elution) or distillation *in vacuo* (for **5c**).

ω-Chloro-2-thiophosphorylalkanonitriles 6–8: General Procedure

Step a: To a stirred solution of thiophosphorylacetonitrile $4(12 \text{ mmol})$ and TEBA (200 mg) in CH₂Cl₂ (20 mL), NaOH as a 50% aqueous solution (48 mmol) was added followed by the dropwise addition of the alkyl halide R^3 Hlg (14.4 mmol). The reaction mixture was stirred for 3 hours at 20◦ C and diluted with water (15 mL); the organic layer was separated, and the aqueous layer was extracted with CH_2Cl_2 . The combined organic fractions were dried $(Na₂SO₄)$ and concentrated *in vacuo*. The oil thus obtained, representing the intermediate thiophosphorylalkanonitrile of 87–96% purity (according to ${}^{1}H$ NMR spectrum), was used in the subsequent step without further purification.

Step b: To a stirred solution of the aforementioned oil and TEBA (200 mg) in CH_3CN (15 mL) Br(CH_2)_nCl ($n = 2 - 4$) (15 mmol) was added, followed by powdered KOH (15 mmol) portionwise

Compound	9	14c(A)	14c(B)	15c(B)	15d(A)	15h(B)
Formula	$C_{17}H_{32}N_2O_4P_2S_2$	$C_{12}H_{14}$ NOPS	$C_{12}H_{14}$ NOPS	$C_6H_{10}NOPS$	$C_{11}H_{12}NOPS$	$C_7H_{12}NOPS$
Molecular weight	454.51	251.27	251.27	175.18	237.25	189.21
Crystal system, Space group	Monoclinic, $P2_1/c$	Monoclinic, C ₂ /c	Triclinic, P1	Monoclinic, $P2_1/c$	Triclinic, P1	Orthorhombic, Pbca
a. A	11.478 (4)	19.072 (4)	10.6650(4)	9.113(6)	10.312(5)	12.4500 (9)
b, A	17.654(4)	12.890(3)	10.9258(4)	6.073(5)	10.996(5)	11.3047(8)
c, A	12.634(4)	10.796 (2)	11.5450(5)	15.077 (9)	11.523(5)	13.1782 (10)
α (°)			72.467 (1)		65.13(3)	
β (°)	95.78(3)	98.99(3)	71.922 (1)	98.62(5)	89.34 (4)	
			86.670 (1)		79.30 (4)	
V, \mathring{A}^3	2547(1)	2621.4 (9)	1218.52 (8)	825(1)	1161(1)	1854.7 (2)
		8				8
d (calc.), g cm ^{-3}	1.185	1.273	1.370	1.410	1.357	1.355
μ (Mo K α), cm ⁻¹	3.56	3.48	3.75	5.19	3.88	4.67
Diffractometer	Siemens P3/C	Siemens P3/C	Smart 1000 CCD	Siemens P3/C	Siemens P3/C	Smart 1000 CCD
Temperature, (K)	298(2)	153(2)	110(2)	298(2)	298(2)	100(2)
$2\theta_{\text{max}}(°)$	50	50	60	64	50	60
No. collected refl., no. unique refl.	4684, 4454	2068, 1996	12751, 6760	3135, 2977	4371.4123	13466, 2666
(Rint)	(0.0318)	(0.0275)	(0.0128)	(0.0403)	(0.0582)	(0.0333)
R_1 (on F for refl. with $1 > 2\sigma(1)$	0.0714	0.037	0.0321	0.0429	0.0425	0.0496
	(for 3537 ref.)	$(for 1363$ refl.)	$(for 6173$ refl.)	(for 1756 refl.)	(for 3754 refl.)	(for 2184 refl.)
wR_2 (on F^2 for all refl.)	0.1849	0.0920	0.0962	0.0950	0.1124	0.1252
	(for 4454 ref.)	(for 1996 refl.)	(for 6760 refl.)	(for 2977 refl.)	(for 4123 ref.)	(for 2666 refl.)
Goodness of fit on F^2	1.048	0.931	1.031	0.850	1.003	1.147

TABLE 8 Crystal Data, Data Collection, and Structure Refinement Parameters for **9** and Some Representatives of **14** and **15**

addition. The reaction mixture was stirred for 2 hours at 20◦ C, a further 15 mmol of the corresponding dihaloalkane was added, and stirring was continued for 1 hour. The reaction mixture was concentrated *in vacuo*, taken up in benzene (15 mL), and diluted with water (10 mL). The organic layer was separated, and the aqueous one was extracted with benzene. The combined organic fractions were dried (Na_2SO_4) and concentrated *in vacuo*. The residue was purified by distillation or column chromatography.

Interaction of the Methylated 4a with Br(CH²)3Br

The methylation of **4a** (14 mmol) was performed according to the aforementioned step a to give the intermediate product in quantitative yield with 96% purity (according to ${}^{1}H$ NMR spectroscopy). For the physicochemical and spectral parameters of the methylated **4a** see Ref. [22]. The substance was used for the subsequent stage without further purification.

The reaction of the latter compound with $Br(CH₂)₃Br$ (18.8 mmol) was carried out according to step b, using the standard workup of the reaction mixture. After concentration, hexane was added to the residue, and this was allowed to stand for 3 days at room temperature. The solid **9** (2.08 g, 64%; meso:d, $l = 1:1$) that precipitated was filtered off.

On attempted isolation of the isomers by column chromatography (100:1 to 100:13 hexane-acetone elution), two main fractions were obtained, the first (100:11 hexane-acetone) representing an oil (meso: $d, l = 3:7$ and the second (100:12 hexane-acetone) being a solid with m.p. $94-95°C$ (meso:d,l = 7:3). After recrystallization of the second fraction from hexane, the meso form was obtained in the pure state, (m.p. 93◦ C).

Anal. found (%): C, 44.93; H, 6.91; N, 6.05; Calcd. for $C_{17}H_{32}N_2O_4P_2S_2$ (%): C, 44.93; H, 7.05; N, 6.17.

Compound 9 (meso): ³¹P NMR (CDCl₃) δ 92.3.¹H NMR (CDCl₃) *δ* 1.35 (t, ³ *J*_{HH} 7.2 Hz, C<u>H</u>₃CH₂O, 12H), 1.55 (d, ² *J*_{PH} 17.6 Hz, C(CN)CH₃, 6H), 1.62-1.90 $(m, C(CN)CH₂, 4H), 2.0–2.13 (m, CH₂CH₂CH₂, 2H),$ 4.14–4.31 (m, OCH2, 8H). 13C NMR (CDCl3) *δ* 15.94 and 16.05 (CH_3CH_2O), 19.54 (C(CN) CH_3), 20.44 (t, $CH_2CH_2CH_2$, ${}^3J_{PC}$ 9.0 Hz), 34.0 ($CH_2CH_2CH_2$), 41.18 (d, ${}^{1}J_{PC}$ 114.6 Hz, P–C), 64.41–64.30 (two d, ${}^{2}J_{PC}$ 7.5 and 7.8 Hz, OCH₂), 119.24 (d, ${}^{2}J_{PC}$ 7.1 Hz, CN).

Compound **9** (**d,l**): ³¹P NMR (CDCl₃) δ 92.1. ¹H NMR (CDCl₃) *δ* 1.36 (t, ³ *J*_{HH} 7.2 Hz, C<u>H</u>₃CH₂O, 12H), 1.57 (d, ² J_{PH} 17.6 Hz, C(CN)CH₃, 6H), 1.62–1.90 (m, $C(CN)CH₂$, 4H), 2.0–2.13 (m, $CH₂CH₂CH₂$, 2H), 4.14– 4.31 (m, OCH₂, 8H).

The reaction of methylated $4a$ with $Br(CH_2)_3Br$ was carried out in the same manner, except that 25 mL of acetonitrile was used in the step of the reaction with $Br(CH_2)$ ₃Br; the residue obtained after the workup of the reaction mixture comprised (according to 31P NMR spectroscopy) the target **7a** (65%); **9** (20%), and the C-methylation product **4a** (15%). The residue was distilled *in vacuo,* the fraction with b.p. 150– 165◦ C (1 mm Hg) being collected and comprising the target **7a** (86%); **15i** (8%) and the C- methylation product **4a** (6%). The fraction was used to produce 2-oxo-1,2-thiaphosphinane **15i** without further purification.

2-Diphenylthiophosphinyl-5-iodopentanonitrile 10a

The mixture of 5-chloro-2-diphenylthiophosphinylpentanonitrile **5f** (0.15 g, 0.45 mmol) and sodium iodide (81.36 mg, 0.54 mmol) in absolute acetonitrile was heated under reflux for 6 hours. After having been allowed to cool, the reaction mixture was concentrated, the residue was taken up in benzene, and sodium chloride was filtered off. The filtrate was concentrated to give the target compound **10a** (0.18 g, 94% purity according to ¹H NMR spectroscopy). ^{31}P NMR (CDCl₃) *δ* 45.69. ¹H NMR (CDCl₃) *δ* 1.85–1.87, 1.94–1.96 (two m, $1H + 1H$, CHCH₂), 2.11–2.17 (m, $CH₂CH₂I$, 2H), 3.11–3.17 (m, ³ J_{HHC} = ³ J_{HH_D} 8.0, ² $J_{H_AH_B}$ 14 Hz,CH2I, 2H, AB-system), 3.54–3.63 (m, CH, 1H), 7.52–7.88 (m, C_6H_5 , 10H).

In the same manner from 5-chloro-2-diphenylthiophosphinyl-2-methylpentanonitrile **7g** (0.15 g, 0.43 mmol) and sodium iodide (77.5 mg, 0.52 mmol), **11a** (0.17 g) was obtained (the reaction mixture comprised about 20% **7g** according to ${}^{31}P$ NMR spectroscopy). ³¹P NMR (CDCl₃) δ 55.09.¹H NMR (CDCl₃) *δ* 1.58 (d, ³*J*_{PH} 16.4 Hz, CH₃C(CN), 3H), 1.82–1.90, 1.98–2.04, 2.09–2.14 (three m, CH₂CH₂CH₂I, 1H + 1H + 2H), 3.10 (t, ${}^{3}J_{\text{HH}}$ 8.0 Hz, 2H, CH₂I), 7.52–7.58, 8.19–8.27 (two m, C_6H_5 , 10H).

³¹P NMR Study of the Intramolecular 10a and 11a S-Alkylation

A solution of 15 mg of the compound **10a, 11a** in absolute acetonitrile was placed in a 5 mm NMR ampule, which was sealed and stored at 20℃ with the ³¹P NMR spectra being recorded weekly.

					Yield $(%)^a$
Compound	Cyclization Method	$b.p.$ (1 mm Hg) (I) or the reflux duration (h) (II)	Methods of Purification and Isolation of Diastereomers	According to the Reaction Mixture NMR ³¹ P	After Purification
14 a^b		150-170	Column chromatography $(C_6H_{14}$:Me ₂ CO = 10:4), A:B (60:40)	40	
14 _b	\mathbf{II}	18	Column chromatography (petrol. ether: $Me2 CO = 10:4$), $B =$ isomer, (petrol. ether:Me ₂ CO = 2:1), A:B (94:6)	61	30
14 _c		170-195	Column chromatography $(C_7H_{16}$: Me ₂ CO = 10:4), B -isomer, $(C_7H_{16}$:Me ₂ CO = 2:1), A:B (75:15) crystallization of A and B isomer mixture ($Et2O$), A-isomer	95	54
14 _c	\mathbf{II}	\overline{a}	Column chromatography (petrol. ether: $Me2CO = 10:4$), B - isomer, (petrol. ether: $Me2CO = 2:1$), A:B (75:15), precipitation $Et2O - A-isomer$	Quantitative	57
15a		140-200	Precipitation Et ₂ O-A:B (43:57), crystallization from C_6H_6 -B- isomer; mother liquor crystallization ($Et2O$), A and B isomer mixture (1:1)	75	51
15 _b		190-210	Column chromatography (petrol. ether: $Me2CO = 10:4$), crystallization from C_6H_6 , B-isomer)	55	21 ^b
15 _b	\mathbf{II}	32	Column chromatography $(C_6H_{14}$: Me ₂ CO = 10:4), A:B (63:37), crystallization from C_6H_{14} , A:B (47:53)	32	17
15c		210-220	Crystallization from C_6H_6 A-isomer, mother liquor crystallization (C_7H_{16}) , B -isomer	30	18
15c	\mathbf{H}	15	Reaction mixture crystallization from C_7H_{16} - A:B (74:26)	53	25
15d		202-210	Column chromatography $(C_6H_{14}$: Me ₂ CO = 10:4) - A -isomer	44	21
15d	Ш	22	Column chromatography $(C_6H_{14}$: Me ₂ CO = 10:4) - A-isomer	79	34
15e	\mathbf{H}	40	Column chromatography (petrol. ether: $Me2CO = 10:1$) - A:B (40:60), crystallization (C_6H_6 : Et ₂)- B -isomer	29	10
15f		193-210	Column chromatography $(C_6H_{14}:Me_2CO = 10:4) - A:B$ (36:54); crystallization (C_6H_6 : C_2H_{14}) - A:B (84:16)	19	6
15f	\mathbf{H}	16	Column chromatography (petrol. ether: $Me2CO = 10:4$) - A-isomer	Quantitative	24
15g		180-190	Column chromatography $(C_6H_{14}:Me_2CO = 10:4) - A:B$ (36:54); crystallization (C_6H_6 : Et ₂ O) - A:B (84:16)	15	6 ^h
15h	\mathbf{H}	25	Column chromatography (petrol. ether: $Me2CO = 10:1$) - A:B (45:55), repeated column chromatography (petrol. ether: $Me2CO = 10:1$) - A:B (73:27), 2:1 - A-isomer	64	22
15i	\mathbf{II}	100	Precipitation by $Et_2O - A:B$ (32:68), crystallization from $Et2O - B-isomer$	75	45

TABLE 9 Methods of purification and isolation of cis- and trans-diastereomers **14** and **15**

^aBased on both diastereomers.

^bThe compound decomposes during chromatographical purification yielding monobasic phosphonic acid EtO(HO)P(O)C(Me)(CN)CH2CH2SH (yield 5% after purification). 31P NMR (*^δ* ppm, DMSO-d6) 17.2. 1H NMR (*δ* ppm, **J** (Hz), DMSO-d6): 1.23 **^t** (3H, CH3CH2, ³**J**HH 7.2), 1.43 d (3H, CH3C(CN), ³**J**PH 14.4), 1.86–1.95, 2.06–2.15 two ^m (1H+1H, CH2CH2SH), 2.59–2.69 ^m (2H, C<u>H</u>₂CH₂SH), 4.02–4.11 m (2H, OCH₂).

3-Cyano-2-oxo-1,2-thiaphosphacyclanes and 3-Alkyl-3-cyano-2-oxo-1,2-thiaphosphacyclanes 14, and 15.

Method I: The Intramolecular Cyclization of ω-Chloro-2-thiophosphorylalkanonitriles (General Procedure). ω-Chloro-2-thiophosphorylalkanonitriles **5–7** produced according to the aforementioned procedure as crude products without further purification [34] were distilled *in vacuo*. After low boiling alkyl halide (R^3Hlg) had been removed, the fraction was collected, boiling, in the ∼160–210◦ C (1 mm Hg) range. The mixture of 3-cyano-2-oxo-1,2-thiaphosphacyclanes **14** and **15** diastereomers obtained was further purified and resolved by column chromatography, fractional crystallization, or by the combination of the previously mentioned methods (Table 9).

Method II: The Intramolecular Cyclization of ω-Iodo-2-thiophosphorylalkanonitriles (General Procedure). The mixture of *ω*-chloro-2-thiophosphorylalkanonitrile **5–7** (3.0 mmol) and sodium iodide (6.0 mmol) in absolute acetonitrile (20 mL) was heated under reflux for 2–40 hours (the course of the cyclization being monitored by the 31P NMR spectroscopy). After cooling, the reaction mixture was filtered. The filtrate was concentrated *in vacuo*, and the residue was taken up in CH_2Cl_2 (15 mL) and filtered again. After concentration, the residue was purified as described in Table 9.

3-Cyano-2-ethoxy-3-methyl-2-oxo-1,2 thiaphosphinane 15i

The solution of 5-bromo-2-diethylthiophosphoryl-2 methylpentanonitrile **7a** (0.15 g, 0.46 mmol) in absolute acetonitrile (10 mL) was heated under reflux for 100 hours (by then the amount of the cyclic product was 75% according to ³¹P NMR data and was not increasing under further heating). After cooling, the reaction mixture was concentrated *in vacuo* and crystallized from ether to give diastereomer **B** (45 mg, 45%) of the compound **15i**. The **15i (B)** was filtered off and the filtrate was concentrated and dried *in vacuo* (1 mm Hg) to give a 2:1 mixture of **A** and **B** diastereomers (40 mg).

3-Cyano-3-methyl-1,2-thiaphospholane Acid Sodium Salt 16

The mixture of 4-chloro-2-diethylthiophosphoryl-2-methylbutanonitrile **6a** (2.5 g, 9.78 mmol) and sodium iodide (3.0 g, 20,0 mmol) in absolute acetonitrile (15 mL) was heated under reflux for 10

hours. The solid that precipitated by then hampered the ability of the reaction mixture to boil uniformly. Thus, it was filtered, and heating was continued for 10 hours. After the standard work-up (see method II) the organic fraction (0.3 g) contained no phosphorus compounds (according to $31P$ NMR). The solid obtained (2.5 g) representing a mixture of the sodium salt **16** and inorganic sodium salts was washed with CH_3CN (25 mL) and CH_2Cl_2 $(2\times25 \text{ mL})$, and the extract was dried *in vacuo*. ³¹P NMR *δ* 47.89 (DMSO-d₆), 51.36 (Py). IR and ¹³C and 1H NMR spectroscopy data of the compound **16,** confirming the heterocycle formation, are shown in Tables 5 and 6. All attempts to isolate the corresponding acid by acidification, followed by extraction of the organic phase, resulted in the destruction of the ring compound.

In the same manner, the sodium salt of 3 cyano-1,2-thiaphosphinanic acid **17** (2.2 g) was obtained from 5-chloro-2-diethylthiophosphorylpentanonitrile **5a** (2.0 g, 7.42 mmol) and sodium iodide (2.36 g, 15.7 mmol). ³¹P NMR *δ* 25.43 (CH₃CN), 25.75 $(CH_2Cl_2).$

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