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SCHIFF BASES DERIVED FROM AMINOMETHYL-DIMETHYL-PHOSPHINE OXIDE

S. Varbanov^a; A. Georgieva^b; G. Hägele^{cd}; H. Keck^c; V. Lachkov^b

^a Institute of Polymers, Bulgarian Academy of Sciences, Sofia, Bulgaria ^b Department of Ecology, Forestry University, Sofia, Bulgaria ^c Department of Inorganic and Structural, Chemistry Heinrich-Heine-University-Düsseldorf;, Düsseldorf, Germany ^d Heinrich Heine Universität Diisseldorf. Inst. Für Anorg. Chemie Universitätsstr, Düsseldorf, Germany

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SCHIFF BASES DERIVED FROM AMINOMETHYL-DIMETHYL-PHOSPHINE OXIDE

S. VARBANOV^a, A. GEORGIEVA^b, G. HÄGELE^{c*}, H. KECK^c and V. LACHKOVA^b

^aInstitute of Polymers, Bulgarian Academy of Sciences, 1113 Sofia, Bulgaria, ^bDepartment of Ecology, Forestry University, 1756 Sofia, Bulgaria and ^cDepartment of Inorganic and Structural Chemistry, Heinrich-Heine-University-Düsseldorf, D-40225-Düsseldorf, Germany

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A group of Schiff bases 1 – 18 derived from aminomethyl-dimethyl-phosphine oxide with aromatic aldehydes have been synthesized. Structure and purity of the new compounds were determined by IR, ¹H NMR and ³¹P{¹H} NMR spectroscopy, mass spectrometry and elemental analysis.

Keywords: Tertiary phosphine oxides; aminomethyl-dimethyl-phosphine oxide; Schiff bases; synthesis

Dedicated to Professor Reinhard Schmutzler on occasion of his 65th birthday

INTRODUCTION

Aminomethyl-dimethyl-phosphine oxide (AMPO) is a very interesting functionalized tertiary phosphine oxide. It has been synthesized independently by L. Maier^[1] and the Sofia team^[2,3] from chloromethyl-dimethyl-phosphine oxide using different methods. It was shown that this phosphorus-containing primary monoamine has a pK_avalue of $6.23^{[3]}$ and possesses high reactivity and nucleophilicity to be used as an organic intermediate reagent, flame retardant for polymers and organic ligand for the synthesis of complexes with metal salts^[2,4–10].

^{*} Author to whom correspondence should be addressed Prof. Dr. G. Hägele – Heinrich Heine Universität Düsseldorf. Inst. für Anorg. Chemie Universitätsstr. 10 D-40225 Düsseldorf, Germany

TABLE I Molecular structures of Shiff bases derived from aminomethyl-dimethyl-phosphine oxide

No.

OH CH=NCH ₂ P(O)(CH ₃) ₂	10 CH=NCH ₂ P(O)(CH ₀) ₂ H	
roxy-1-naphthylmethylene-imino-methyl)-dimethyl-phosphine oxide OH CH=NCH ₂ P(O)(CH ₃) ₂	(2-Indolylmethylene-iminomethyl)-dimethyl-phosphine oxi	
oxyphenylmethylene-iminomethyl)-dimethyl-phosphine oxide	(4-Methyl-phenylmethylene-iminomethyl)-dimethyl-phospi	hine oxid

13

14

Compound

-CH=NCH2P(O)(CH3)2 xy-2-hydroxyphenylmethylene-iminomethyl)-dimethyl-phosphine oxide

rophenylmethylene-iminomethyl)-dimethyl-phosphine oxide

, -CH=NCH₂P(O)(CH₃)₂

rophenylmethylene-iminomethyl)-dimethyl-phosphine oxide

-CH=NCH₂P(O)(CH₃)₂ (4-Methoxy-phenylmethylene-iminomethyl)-dimethyl-phosphine ox

Compound

(3-Methoxy-4-hydroxy-ohenylmethylene-iminomethyl)-dimethyl-ph

(4-Nitro-phenylmethylene-iminomethyl)-dimethyl-phosphine oxide

olidinyl)phenylmethylene-imino-methyl]-dimethyl-phosphine oxide odidinyl)-phenylmethylene-imino-methyl]dimethyl-phosphine oxide		CH=NCH ₂ P(O)(CH ₃) ₂
ary		$(9-An thracenyl methyleneiminomethyl)-dimethyl-phosphine\ oxide$
CH=NCH ₂ P(O)(CH ₃) ₂	16	CH=NCH ₂ P(O)(CH ₃) ₂
⊕ H √Emethylene-iminomethyl)-dimethyl-phosphine oxide		(Phenylmethylene-iminomethyl)-dimethyl-phosphine oxide
p p (CH₃)₂N CH=NCH₂P(O)(CH₃)₂	17	CH=NCH ₂ P(O)(CH ₀) ₂
tgylamino-phenylmethylene-imino-methyl)-dimethyl-phosphine oxide		[3,4-(Methylenedioxy)phenylmethylene-iminomethyl)-dimethyl-pho oxide
	18	

No.

15

Compound

______CH=NCH₂P(O)(CH₃)₂

(3-Pyridinyl-methylene-iminomethyl)-dimethyl-phosphine oxide

Compound

CH=NCH2P(O)(CH3)2

CH=NCH2P(O)(CH3)2

peridinyl)-phenylmethylene-imino-methyl]-dimethyl-phosphine oxide

A part of the compounds reported previously exhibits biological activity, e.g. the nitrosourea derivative^[5] and the platinum complexes^[9] exert antitumor activity being of low toxicity, while the phenoxyphenylalkylphosphine oxides, prepared by L. Maier are proved to be active herbicides^[4].

The present work is a continuation of our investigations on synthesis and characterization of functionalized tertiary phosphine oxides, based on aminomethyl-dimethyl-phosphine oxide^[2,3,5-10] and reports the preparation and characterization of its Schiff bases with aromatic aldehydes **1-18** (Table I). They are expected to show biological activity and complex-forming properties with metal ions as well. They could be used as organic intermediates for preparation of new organophosphorus-compounds similarly to known azomethyne derivatives^[1,11-14].

RESULTS AND DISCUSSIONS

The Schiff bases 1–18 were prepared by condensation of AMPO with corresponding aldehydes in benzene solution at room temperature according to the reaction:

(1)
$$R \cdot CHO + H_2N \cdot CH_2P(O)(CH_3)_2 \longrightarrow R \cdot CH=N \cdot CH_2P(O)(CH_3)_2 + H_2O$$

The yields of the majority of the reaction products proved to be very high, in general about or more than 90%. No catalyst is required. Most of the reactions observed were exothermic. The azomethyne derivatives crystallized from the reaction mixture after mixing the reagents and stirring for 3–4 hrs, usually without azeotropic distillation of the water.

Some preparative and analytical data on Schiff bases 1–18 are given in Table 2. The compounds are colourless or coloured crystalls with characteristic melting temperatures. Most of the compounds 1–18 are hygroscopic, easily dissolved in DMFA, methanol, ethanol, dichloromethane, chloroform, but sparingly soluble in diethyl ether, dioxane, tetrahydrofurane, aliphatic and aromatic hydrocarbons.

Molecular structure and purity for 1–18 were confirmed by IR, ¹H NMR and ³¹P{¹H} NMR spectroscopy and mass spectrometry. In addition results from the elemental analysis (phosphorus) are given in Table II.

TABLE II Preparative and analytical data on Schiff bases of aminomethyl-dimethyl-phosphine oxide

No Yield		M.p., °C / (Recr. solv.) / Colour	General formula Mol. mass	Phosphorus Content %	
	70		moi. mass	Found	Calcd.
1	86	153-155 / (Ethyl acetate) / White	C ₁₄ H ₁₆ NO ₂ P	11.72	11.86
			261.26		
2	86	102-105 / (Ethyl acetate) / Light yellow	$C_{10}H_{14}NO_2P$	14.47	14.67
			211.20		
3	87	120-122 / (Ethyl acetate) / Yellow	$C_{12}H_{19}NO_3P$	11.86	12.09
			256.26		
4	69	45-48 / (Ethyl acetate) / Cream	C ₁₀ H ₁₃ FNOP	14.45	14.53
			213.19		
5	82	124-126 / (Ethyl acetate) / White	C ₁₀ H ₁₃ CINOP	13.36	13.49
			229.05		
6	94	111 146 / (Ethyl acetate) / Yellow orange	$C_{14}H_{21}N_2OP$	11.55	11.72
			264.30		
7	85	142-144 / (Ethyl acetate) / Beige	$C_8H_{13}N_2OP$	16.73	16.82
			184.18		
8	97	135–137 / (Ethyl acetate) / White	$C_{12}H_{19}N_2OP$	12.82	13.00
			238.27		
9	50	134–137 / (Ethyl acetate) / White	$C_{15}H_{23}N_2OP$	11.05	11.13
			278.33		
10	98	205–207 / (Dichloromethane) / White	$C_{12}H_{15}N_2OP$	13.08	13.22
			234.24		
11	96	43–45 / (Benzene) / White	C ₁₁ H ₁₆ NOP	14.67	14.80
	00	046 046 (CD) 1 \ 1370 '.	209.23		10.55
12	90	245-246 / (Ethyl acetate) / White	C ₁₁ H ₁₆ NO ₂ P	13.62	13.75
12	05	190 101 / (Damana) / 377 /	225.23	10.72	12.04
13	95	189–191 / (Benzene) / White	C ₁₁ H ₁₆ NO ₃ P	12.73	12.84
			241.23		

No Yiel	Yield	M.p., °C/(Recr. solv.)/Colour	General formula Mol. mass	Phosphorus Content %	
	70		MOL Mass	Found	Calcd.
14	90	159-160 / (Ethylacetate) / White	C ₁₀ H ₁₃ N ₂ O ₃ P 240.20	12.80	12.90
15	98	130-134 / (Ethyl acetate) / Yellow	C ₁₈ H ₁₈ NOP 295.31	10.38	10.49
16 ^a	92	30–32 / (Benzene) / White	C ₁₀ H ₁₄ NOP 195.20	15.72	15.87
17	74	36-37 / (Ethyl acetate) / Yellow	C ₁₁ H ₁₄ NO ₃ P 239.21	12.81	12.95
18	82	119-120 / (Ethanol) / White	C ₉ H ₁₃ N ₂ OP 196.19	13.36	13.49

a. The compound was reported before us by L. Maier^[1]: b.p. 133–138°C/0.06 torr.

The infrared spectra (Table III) showed characteristic bands assigned to the phosphoryl group (P=O) at 1138–1199 cm⁻¹, methyl (CH₃-P) and methylene (CH₂P) groups bonded to a phosphorus atom at 1288–1305 cm⁻¹ and 741–769 cm⁻¹, respectively. Bands at 1459–1521 cm⁻¹ and 1550–1615 cm⁻¹ are typical for aromatic rings. The Ar-O-R bonds of 3, 12, 13 and 17 were confirmed by bands at 1024–1047 cm⁻¹ and 1252–1269 cm⁻¹ while the Ar-OH group in 1, 2, 3 and 13 was supported by bands at 1204–1213 cm⁻¹ Very strong, sharp bands at 1608–1644 cm⁻¹ are characteristic for the monoconjugated CH=N groups^[15,16]. Additional bands of the phosphoryl group (P=O) have been observed in the spectra of most of the compounds. This phenomenon was reported by many authors and could be ascribed to different spatial isomers^[17,18].

¹H NMR spectra of **1–18** (Table IV) showed resonance signals for all protons in the compounds: doublets for a) methyl protons $C\mathbf{H}_3P=O$ at $\delta=1.29$ to 1.63 ppm and $^2J_{PH}=-12.6$ to -13.2 Hz, b) methylene protons $C\mathbf{H}_2P=O$ at $\delta=3.80$ to 4.42 ppm and $^2J_{PH}=-11.9$ to -13.8 Hz, c) $C\mathbf{H}=N$ protons at $\delta=8.08$ to 9.55 Hz and $^4J_{PH}=1.9$ to 4.7 Hz. The complex resonance signals for the aromatic ring protons were found in the region of

 δ = 6.00 to 8.70 ppm, while the Ar-O**H** protons in **1–3** occurred as singlets at δ = 9.82 to 14.48 ppm. The relatively high δ -values of Ar-O**H** in 1–3 is indicative for intramolecular hydrogen bonds formed with the nitroger atom of -CH=N-groups, an observation which is typical for o-hydroxysubstituted Schiff bases^[16,19–21]. The resonance signal of the Ar-O**H** proton in **13** was found at 1.78(s) ppm. No intramolecular hydrogen bond is formed, since the OH group is in p-position to the CH=N group^[21]. It is worth mentioning that ²J_{PH} of C**H**₂P=O for all **1–18** were found to higher absolute values than reported previously for similar units having ²J_{PH} in a range –3 to –8 Hz^[2.6,22–25].

 $^{31}P\{^{1}H\}$ NMR spectra of **1–18** were singlet resonance signals at $\delta = +41.01$ to +45.03 ppm typical for tertiary phosphine oxides binding two methyl and one methylene groups to the phosphorus atom^[25,26].

All compounds were characterized by mass spectrometry and significant mass spectrometric data are presented in Table V. EI mass spectra that show signals for [M]⁺were obtained for compounds 1–3, 6–10, 13 and 15 only. Two fragmentations are typical for the molecular ions. Loss of (CH₃)₂PO radicals (-77u) takes place that leads *via* α-cleavage to even-electron ions [RCHNCH₂]⁺. Base peaks in all the EI mass spectra are signals for [(CH₃)₂P⁺(OH)CH₂⁻] (m/z 92). This ylidion with positive charge located on the phosphorus and the unpaired electron on the adjacent carbon atom is formed from [M]⁺. by elimination of RCN *via* McLafferty rearrangement. It should be noted that this ion is an isomeric form of the molecular ion of trimethylphosphine oxide [(CH₃)₃PO]^{+*}. It has been shown by *ab initio* MO calculations that the unconventional ylidion lies 86 kJ/mol lower in energy than the ion with conventional structure and is separated from the latter by a substantial isomerization barrier of about 195 kJ/mol^[28].

Not all the Schiff bases could be measured by EI mass spectrometry possibly due to thermal decompositions of the compounds prior to ionization. However, in case of the remaining compounds **4**, **5**, **11**, **12**, **14**, **16–18** FAB mass spectrometry proved to be a powerful method. The FAB mass spectra that have been measured with 3-nitrobenzyl alcohol as matrix are characterized by intense signals for [M+H]⁺ (base peaks). A typical fragmentation under FAB conditions is the elimination of (CH₃)₂POH (-78 u) leading to cations [RCHNCH₂]⁺. The formation of the latter ions has been observed in the EI mass spectra as well.

TABLE III Characteristic infrared frequences (ν cm $^{-1}$) of Schiff bases derived from aminomethyl-dimethyl-phosphine oxide

No	P=O	CH ₃ P	CH ₂ P	C=N	Ar	Ar-O-R	Ar-OH
1	1145(w)	1298(m)	763(m)	1624(s)	1497(w)		1211(m)
	1169(vs)		(,	(-)	1595(w)		3350(w)
					` '		3500(w)
2	1157(s)	1292(w)	750(s)	1629(s)	1489(m)	_	1213(w)
	1164(vs)	. ,		, ,	1574(m)		• •
3	1173(vs)	1297(m)	750(m)	1629(s)	1495(w)	1047(w)	1205(w)
					1582(w)	1252(vs)	
4	1138(vs)	1295(m)	756(s)	1633(s)	1484(s)	_	_
	1147(vs)				1557(w)		
5	1144(m)	1288(w)	769(vs)	1630(s)	1467(m)	_	_
	1167(vs)				1591(w)		
6	1178(vs)	1299(m)	768(w)	1610(vs)	1491(w)	_	_
					1550(m)		
7	1156(vs)	1295(s)	759(w)	1635(vs)	_	-	-
	1193(s)						
8	1175(vs)	1300(w)	743(w)	1612(vs)	1449(w)	-	-
					1551(w)		
9	1162(s)	1302(m)	750(w)	1608(vs)	1452(w)	-	-
	1179(vs)				1560(w)		
10	1156(vs)	1301(m)	750(s)	1627(vs)	1459(m)	-	
	1169(s)				1579(w)		
11	1161(vs)	1304(m)	750(w)	1647(m)	1461(w)	-	-
	1189(s)				1573(w)		
	1199(s)						
12	1166(vs)	1303(m)	759(w)	1642(m)	1511(w)	1027(m)	_
	1179(vs)				1577(w)	1253(vs)	
13	1154(vs)	1288(vs)	759(w)	1635(m)	1521(s)	1039(m)	1204(w)
	*****	1000()	= • • • •		1591(m)	1256(s)	
14	1166(vs)	1298(w)	746(s)	1634(m)	1511(s)	_	-
15	1171(vs)	1205()	741(-)	1620()	1602(m)		
15	1139(s)	1295(m)	741(s)	1629(m)	1520(w)	_	-
	1158(m) 1174(vs)				1615(m)		
16		1200(m)	754(m)	1638(s)	1/01(w)		
10	1163(vs)	1290(m)	754(m)	1030(8)	1491(w) 1579(w)	-	-
17	1139(s)	1291(m)	750(m)	1636(s)	1579(w) 1513(vs)	1024(s)	_
1,	1161(vs)	1271(111)	/30(III)	1050(8)	1575(vs) 1599(m)	1024(s) 1269(vs)	_
18	1161(vs)	1305(s)	749(s)	1611(s)	1339(m) 1478(m)	-	_
10	1101(vs) 1194(s)	1202(3)	177(3)	1011(8)	1586(m)	_	_
	1174(2)				1200(111)		

H-NMR $C\underline{H}_2P=O$

 δ_H

4.17(d)

4.09(d)

3.80(d)

4.08(d)

4.06(d)

4.06(d)

4.16(d)

 $^{2}J_{PH}$

11.9

12.0

12.6

13.2

12.6

12.6

12.6

 $C\underline{H}_3P=O$

 δ_H

1.63(d)

1.29(d)

1.56(d)

1.56(d)

1.57(d)

1.61(d)

¹00 1.59(d)

 $\overline{^{2}J}_{PH}$

12.6

12.6

13.2

13.2

12.6

12.6

12.6

1.58(d)	13.2	4.09(d)	13.2	8.43(d)	3.8	6.00 - 7.00(m)	12.82(bs)	-
ទី 1.58(d)	13.2	4.14(d)	13.2	8.63(d)	4.4	7.00 - 8.00(m)	-	-
[∞] 1.58(d)	13.2	4.16(d)	13.2	8.77(d)	3.8	7.26 – 8.05(m)	_	-
∺ 1.54(d)	12.6	4.03(d)	13.8	8.15(d)	3.8	6.50(d), 7.60(d) ^c	-	
∷ d 1.53(d)	12.6	3.98(d)	12.6	8.08(d)	3.8	-		4
ଚ୍ଚୁ 1.54(d)	13.2	4.03(d)	13.2	8.16(d)	3.8	6.69(d), 7.59(d) ^c	-	
ਰੂ 8 1.54(d)	12.6	4.04(d)	13.2	8.17(d)	4.7	6.68(d), 7.59(d) ^c	_	-

TABLE IV ^{1}H and $^{31}P\{^{1}H\}$ NMR data of Schiff bases of aminomethyl-dimethyl-phosphine oxides

 $C\underline{H} = N$

 δ_H

9.27(d)

8.43(d)

8.31(d)

8.27(d)

8.25(d)

8.19(d)

8.43(d)

 $^{4}J_{PH}$

2.5

3.1

3.1

3.8

3.7

3.8

3.8

Ar-OH

 δ_H

14.47(bs)

12.47(s)

1.78(s)

Ar-<u>H</u>

 δ_H

7.10 - 8.10(m)

6.85 - 7.40(m)

6.90 - 8.10

7.23(d), 7.63(d)^c

 $6.93(d), 7.68(d)^{c}$

6.91 - 7.34(m)

7.91(d), 8 28(d)^c

					u	սս (ծ)չը չ pue	ասս (၁)ՀՑ չ	A te anotora l
							.mqq (s) č8	.otons at 8=2.
		.zH 2.2=HH	l ^{(c} ,mqq (1)9	protons at 3.2	$C\overline{\mathbf{H}}^{S}NC\overline{\mathbf{H}}^{S}$	pas mqq (m)27	7.1-03.1=8 1	3-C protons a
11			•		·wc	iq (2)50.5=6 is	protons was	al of C H 3-N
Swdd (sq)	oton at 6=9.28	74 lovyq H N 101 bas (s)26.9=8	;zH T.E=HH	le, ,mqq (b)42.	3-8 ;5H 1.E	(1) bbw [,] 21 ^{HH} =	92.9 = 6:26	ons from pyro
5								blets.
ua.		$sH \ \delta. \xi = HH \ U^{\epsilon} \ bas \ (wp) + \xi. \xi$	Hz and 5=3	1.£=HHIc ,(wp)	e at <i>8</i> =2.02(ne protons wen	ine methyle	als of pyrolid
Jan		;zH 6.3=HHl ^E ,(p)E1.4=	=8 bns sH e.	9=HH[c (1)wd	q 94. l=8 is	bnuoi sisw end	AO ₂ Toto	als of $\mathbf{CH}_3\mathbf{CI}$
80		·		·				
6								riplet, q -quar
ան-ա ,191ժ	singlet, d- dou	n as absolute values, bs-broad	J _{PH} are give	tive. Data for ² ,	legan zi _{Hq} l	² To ngis adT .[sH] t .[mqq] _q 8 bns _H 8 :s
7 4-1	_	(m)07.8 – 0£.7	6'1	(b)09.8	12.6	4.12(d)	12.6	(b)8č.1
¥ †		(m)0L 8 - 0E L	01	(P)00 8	9 61	(P)CI V	961	(1 58(4)
[6]+	_	(m)02.7 - 28.8	8.£	(b)£2.8	15.6	(b)70.4	13.2	(b)72.1
ਰ								

(b)££.8

(b)&&.9

 $H_{\mathbf{Q}}$

C**H**=N

13.2

15.6

Hdfz

4.11(d)

(b)24.4

 H_{Q}

 $O=d^{\overline{L}}\overline{H}O$

8.€

4.4

 $^{Hd}I_{p}$

 $(m)08.\Gamma-04.\Gamma$

(m)08.8 - 04.7

 $H_{\mathbf{Q}}$

 \overline{H} -1V

 $H_{\mathbf{Q}}$

₩O-1¥

.mqq (2)EQ.£ bns mqq (2)Z8.£ δ is snotong .zH ϵ . ζ = H_H L bns mqq (b) ϵ 9. ϵ is snotong

13.2

13.2

Hd [Z

(b)72.1

(b)69.1

 H_Q

 $O=d^{\xi}\overline{H}\mathcal{I}$

TABLE V Significant mass spectrometric data (m/z / rel. Int. %) of Schiff bases 1-18 derived from aminomethyl-dimethyl-phosphine oxide (results of El-MS are indicated by $^{a)}$, those of FAB-MS by $^{b)}$)

compound	method	[M+H] ⁺	[M] ⁺	[RCHNCH ₂] ⁺	[(CH ₃) ₂ P ⁺ (OH)CH ₂ *] (m/z 92)
1	a)		261/10	184/13	/100
2	a)		211/9	134/13	/100
3	a)		255/23	178/12	/100
4	b)	214/100		136/41	
5	b)	230/100		152/39	
6	a)		264/20	187/92	/100
7	a)		184/12	107/47	/100
8	a)		238/14	161/58	/100
9	a)		278/19	201/76	/100
10	a)		234/7	157/20	/100
11	b)	210/100		132/44	
12	b)	226/100		148/39	
13	a)		241/6	164/6	/100
14	b)	241/100		163/19	
15	a)		295/8	218/11	/100
16	b)	196/100		118/46	
18	b)	197/100		119/35	

EXPERIMENTAL

Starting materials

Aminomethyl-dimethyl-phosphine oxide was prepared according to reference^[2]. Aromatic aldehydes were commercially available products. Liquid aldehydes were purified by distillation prior to use. Solvents were dried by standard procedures.

Characterization of the prepared compounds 1-18

The elemental analysis for phosphorus content was performed according to reference^[27]. Melting point temperatures, measured on a Boetius microheating plate PHMK 05 (Germany), were uncorrected. Infrared spectra

 $(400-4000~{\rm cm}^{-1})$ were recorded on a Bruker IFS-113V spectrometer using KBr pellets. The ^{1}H NMR spectra for solutions of 1-9 and 11-18 in CDCl₃ were obtained from a Bruker DRX 500 NMR spectrometer operating at 500.13 MHz. 10 was dissolved in DMSO-d₆. The chemical shifts are referenced vs. int. TMS. $^{31}P\{^{1}H\}$ NMR spectra were obtained at 202.45 MHz. Chemical shifts $\delta_{\rm P}$ are given vs. ext. 85% $H_{\rm 3}PO_{\rm 4}$. The electron impact (EI) mass spectra were measured on a Varian MAT 311A at 70 eV using the direct inlet system. Fast atom bombardment (FAB) mass spectra were obtained with a Finnigan MAT 8200 mass spectrometer using 3-nitrobenzyl alcohol as matrix. Argon was used as the particle source. The FAB gun was operated at 2 mA discharge current with an acceleration voltage of 8 kV.

General procedure for the preparation of 1-18

To a stirred solution of the aldehyde (2.0 mmol) in dry benzene (5 ml) at room temperature was added dropwise a solution of AMPO (2.0 mmol) in dry benzene (5 ml). The mixture was stirred at room temperature for 3-4 hrs. In some of the cases the reaction water was distilled azeotropically. The precipitate formed was isolated by filtration, washed with fresh solvent and dried. The prepared crude product was recrystallized from the corresponding solvent till a constant melting temperature. The preparative and analytical data of 1-18 are presented in Table II.

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