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ADVANTAGES OF USING DI(p-METHYLBENZYL) HYDROGEN PHOSPHITE IN SYNTHESIS OF AMINOPHOSPHONATES FROM ALDIMINES

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Aminophosphonic acids containing groups sensitive to hydrolysis and hydrogenation are obtained easily from aldimines by addition of di(p-methylbenzyl) hydrogen phosphite followed by selective removal of p-methylbenzyl groups by solvolysis with formic acid. Properly chosen substituens at nitrogen are also removed under these conditions and the synthesis of aminophosphonic acids with a free NH₂ group was achieved by a two-step procedure.

The p-methylbenzyl group, first proposed by Miyano and Funahashi for protection of phosphate function, ¹⁾ appears not to be widely used in syntheses of phosphates or phosphonates, although it offers some advantages over the most commonly used benzyl group. Both these groups are removed by hydrogenolysis at comparable rates, but in reaction sequences starting with secondary phosphites it is advantageous to use crystalline di(p-methylbenzyl) hydrogen phosphite instead of dibenzyl phosphite which is difficult to purify. Moreover, we found that the p-methylbenzyl group is more readily removed by comparatively mild acidolysis with formic acid than the unsubstituted benzyl group.

In this communication we demonstrate the usefulness of di(p-methylbenzyl) hydrogen phosphite ($\underline{2}$) in synthesis of aminophosphonic acids containing functions sensitive to hydrogenation and hydrolysis. Like other secondary phosphites, $\underline{2}$ readily adds to the C=N bond of aldimines ($\underline{1}$), giving high yields of esters($\underline{3}$). Removal of p-methylbenzyl groups, achieved by brief formolysis, affords aminophosphonic acids ($\underline{4}$) and p-methylbenzyl formate ($\underline{5}$). Selective removal of a p-methylbenzyl group is demonstrated by successful prepa-

ration of products <u>4a</u> and <u>4b</u> (Table 1) with intact methoxycarbonyl or diethoxyphosphinyl group. The presence of nitro and ester functions in these products obviously prevents removal of protecting group by hydrogenolysis or silylation.

$$R-CH=N-R^{+} (R^{\circ})_{2}PHO \xrightarrow{110^{\circ}C}_{30 \text{ min.}} R-CH-PO(OR^{\circ})_{2} \xrightarrow{98\% \text{ HCOOH}}_{105^{\circ}C,1 \text{ min.}} R-CH-PO_{3}H_{2} + 2HCOOR^{\circ}$$

$$\frac{1}{NHR^{\circ}} \xrightarrow{2} \frac{4}{105^{\circ}C,1 \text{ min.}} H^{-}R^{\circ}$$

$$R = m-NO_{2}C_{6}H_{4} R^{\circ} = CH_{2}COOMe \ \underline{1a}, \underline{3a}, \underline{4a} R^{\circ} = CH_{2}CH_{2}PO(OEt)_{2} \ \underline{1b}, \underline{3b}, \underline{4b}$$

$$R^{\circ} = p-CH_{3}C_{6}H_{4}CH_{2}$$

Preparation of the biochemically more interesting aminophosphonic acids with free NH₂ group from aldimines requires removable substituents at nitrogen.^{2,3)} Particularly useful are tertiary benzyl groups, like 1-phenyl-1-cyclopentyl,³⁾ removable by acidolysis under conditions necessary for removal of p-methylbenzyl from the phosphonate function. Thus, using aldimines prepared from 1-phenyl-1cyclopentylamine and benzaldehyde, isobutyraldehyde, furfural or 2-thiophenecarbaldehyde, we were able to obtain high yields (see Table 1) of aminophosphonic acids <u>8a</u> - <u>8d</u> in two steps, involving addition of <u>2</u> to aldimines <u>6a</u> - <u>6d</u>, followed by simultaneous formolytic removal of substituents from amino and phosphonate functions. Preparation of <u>8a</u> by methods involving acid hydrolysis or hydrogenolysis would obviously be difficult because of the presence of acid-sensitive furan ring.⁴⁾ Similarly, the thiophene ring in <u>8b</u> would make hydrogenoly-



$$\xrightarrow{\text{R-CH-PO}_{3}\text{H}_{2}} + 5 + \text{C}_{6}\text{H}_{5} - \boxed{8}$$

<u>6a</u>, <u>7a</u>, <u>8a</u>: R = 2-furyl <u>6b</u>, <u>7b</u>, <u>8b</u>: R = 2-thienyl <u>6c</u>, <u>7c</u>, <u>8c</u>: R = isopropyl <u>6d</u>, <u>7d</u>, <u>8d</u>: R = phenyl <u>6e</u>, <u>7e</u>: R = H

Tabl	le l. Aminophosphonic aci	Lds obte)ib vd by di	p-methylbenzyl) hydrogen pho	sphite addition to aldimine and formolysis
No	Formula	Yield [#] (%)	(Do) dM	IR major bands (¹ H-NMR S,ppm
<u>4</u> 8	Q → CHP0 ₃ H ₂ 02 ^N NHCH ₂ COOCH ₃	67	241-3(đec)	1100,1190-1210,1355,1530, 1605,1745,2300-3000	$3.45(s, 2H, CH_2), 3.70(s, 3H, CH_3), 4.26(d, 1H, CH, ^2J_{PH}^{=17Hz}), 7.81-8.80(m, 4H_{ar});$ (in $D_2O + D_2SO_4$)
44	O2 ^N NHCH ₂ CHPO ₃ H ₂	79	200-202	1020,1085,1150,1220,1350, 1530,2400-3000	1.02-1.38(m,6H,0CH ₂ CH ₃),2.13-2.78(m,2H,CH ₂ P) 3.13-3.79(m,2H,CH ₂ N),3.86-4.47(m,4H,0CH ₂ CH ₃) 4.76-5.25(m,1H,CHP),7.45-8.63(m,4Har);
8 8	PO ₃ H2	82	225 -7 (đec)	920,1070,1225,1545, 2600-3000,3100	(in CF ₃ COOH) 4.90(d,1H,CHP, ² J _{PH} =16Hz),6.74-7.00(m, ^{2H} _{furyl}) 7.88(b.s, ^{1H} _{furyl}); (in D ₂ O)
8	CH PO _{3H2}	85	248 - 50(đec)	720,940,1075,1155,1190,	5.20(d,1H,CHP, ² J ⁼ 16Hz),7.37-7.96(m,3H _{thien}) (in D ₂ 0 + D ₂ S0 ₄)
80	CH ₃ CH ₃ PO _{3H2}	84	2713 ^{##}		
89	CHNH2 CHNH2	77	277–9 ***		
	NHCH ₂ PO ₃ H ₂	74	257 9	890,1085,1115,1180,1210, 1225,1290,1610,2100-2700, 2960,3100	1.87-2.50(m,8H,CH ₂ x4),2.65(d,2H,CH ₂ P, ² J _{PH} = 14Hz),7.82(b.s,5H _{ar}); (in D ₂ O + NaOD)
X ie Kre fai	ald based on aldimines. sutzkampf ⁶)(<u>8d</u>). ^{xxxx} Prepε led to remove substituer	MMKIden ared fro it from	utified by community of the second of the se	omparison with authentic sam r conditions described in ge ore drastic hydrolysis was r	ples prepared after Berlin ⁵)(<u>ac</u>) or meral procedure. In this case formolysis ecessary to prepare aminomethanephosphonic
aci	id from <u>7e</u> in one step."				

<u>General procedure:</u> A mixture of $\underline{2}$ (0.01 mole) and an aldimine (0.01 mole)³⁾ was heated at 110-120^o C for about 30 minutes. After cooling the crude reaction mixture was treated with 20 ml of formic acid (98% commercial grade, FLUKA) and was heated to boiling for 1 minute. Volatiles were then removed under reduced pressure and oily residues were extracted with anhydrous ether (3x25 ml). Resulting solid or semisolid aminophosphonic acids were recrystallized from ethanol to give pure specimens.

Unsubstituted benzyl group is not removed completely by formolysis under conditions specified above. Thus, formolysis of the dibenzyl ester <u>9</u> yielded the monobenzyl ester <u>10</u>.



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