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THE REACTION OF METHYL DIAZOACETATE WITH 4,4-DIMETHYL-2-PENTYN-1-AL AND RELATED COMPOUNDS

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Summary

Methyl 2-diazo-3-hydroxy-6,6-dimethyl-4-heptynoate was prepared by treating 4,4-dimethyl-2-pentyn-1-al with methyl diazoacetate at room temperature in the absence of catalysts. In the case of related aldehydes, $RC \equiv CCHO$ (R = n-Bu, Me_3Si , Et_3Si , Et_3Ge), this unusual reaction is partially or completely suppressed by a competing 1,3-cycloaddition process. The latter leads to a mixture of isomeric 3- and 4-formylpyrazole, one of which cyclodimerizes to form tricyclic hemiaminal.

Introduction

4,4-Dimethyl-2-pentyn-1-al (Ia) has broad synthetic applicability as a synthon since it posseses two reaction sites. For example, the carbonyl group and the carbon—carbon triple bond are reactive towards diazo compounds. The reaction of diazomethane and its derivatives with aldehydes has been shown to give corresponding epoxides and/or ketones [1]. Dipolar addition of diazo compounds to carbon—carbon multiple bonds is also well known [2]. Thus, when propynal was treated with ethyl diazoacetate in ether, only 3-formyl-4-(ethoxycarbonyl)pyrazole was isolated in 86% yield [3]. We describe below a study of the scope of the reaction between acetylenic aldehydes and methyl diazoacetate *.

Results and discussion

Methyl diazoacetate can react with aldehydes of general formula R—C= CCHO (Ia—Ie) in different ways: 1,3-cycloaddition and/or addition to the

^{*} Preliminary communication, see ref. 4.

carbonyl group (Scheme 1). Typical reaction conditions involve treatment of aldehyde with one equivalent of the diazo ester in ether at room temperature in the dark. Under these conditions 4,4-dimethyl-2-pentyn-1-al (Ia) was converted exclusively to an aldol-type adduct. The analytical data obtained for the adduct were compatible with the formulation of the product as methyl 2-diazo-3-hydroxy-6,6-dimethyl-4-heptynoate (Va). This reaction is particularly surprising in view of the stability of the triple bond towards the diazo ester. Moreover, this bond in Ia is apparently more stable to methyl diazoacetate than the carbon—oxygen multiple bond. For this compound the explanation may lie in shielding of the triple bond by the bulky tert-butyl group. In accord with this conclusion, addition of methyl diazoacetate to an ether solution of n-BuC=CCHO (Ib) under conditions similar to those used for Ia causes the formation of isomeric pyrazoles IIb and IIIb together with methyl 2-diazo-3-hydroxy-4-nonynoate (Vb) (see Table 1).

SCHEME 1

$$N_2$$
CHCOOMe + RCECCHO

(II)

(a) R = tert-Bu

(b) R = n-Bu

(c) R = Me₃Si

(d) R = Et₃Si

(e) R = Et₃Ge

(IV)

RCECCH(OH)C(N₂)COOMe

TABLE 1
REACTION OF 4,4-DIMETHYL-2-PENTYN-1-AL AND RELATED COMPOUNDS (Ia—Ie) WITH METHYL DIAZOACETATE

Starting reagent	Produ	cts (%) ^a			
	II	ш	IV	v	
t-BuC≡CCHO (Ia)		_		100	
n-BuC≡CCHO (Ib)	5	4	38	53	
Me ₃ SiC≡CCHO (Ic)	15	50	. 35	_	
Et3SiC≡CCHO (Id)		45	10	45	
EtaGeC=CCHO (Ie)	10	35	25	35	

^a Determined by ¹H NMR spectroscopy.

ANALYTICAL AND OTHER DATA FOR THE COMPOUNDS PREPARED IN THIS INVESTIGATION

TABLE 2

Compound	œ	Yield	m.p. (°C)	n20 nD	Analysis, for	Analysis, found (calcd.) (%)			
					N.	ວ	H	Si/Ge	
IIIc	Me3	43	152-153		12.47	47.62	6.32	12,53	
,	i	į			(12,37)	(47.74)	(6.24)	(12.41)	
IIId	Et3Si	.25	66-86		10,68	53.94	7.50	10.57	
111.0	0 1	· C	900		(10.44)	(53.70)	(7.51)	(10.46)	
1116	E13Ge	99	108—108		8.62	45.84	6.35	23.54	
IVb	n-Bu	19	89—90		13.21	57.08	6.78	(43.44)	
					(13.33)	(57.14)	(6.67)		
IVc	Me ₃ Si	23	52-53		12,18	47.86	6.44	12.24	
					(12.37)	(47.74)	(6.24)	(12.41)	
IVe	Et ₃ Ge	35	138 - 139		8.99	46.04	6.24	23.16	
					(8.95)	(46.07)	(6.40)	(23.22)	
Va	t-Bu	88	oil	1,4866	13.20	57.21	6.59		
					(13.32)	(57.13)	(6.71)		
Vb	n-Bu	33	Πο		13,01	57.48	6.82		
					(13.32)	(57.13)	(6.71)		
ρΛ	Et3Si	54	lio	1,5030	10,34	53,52	7.50	10.21	
					(10.44)	(53.70)	(7.51)	(10.46)	
Ve	Et ₃ Ge	28	tjo	1.5040	7.17	46.09	6.43	23.08	
					(6.95)	(46.07)	(6.40)	(23.22)	
VII	Et3Ge	100	8990		7.43	46.97	7.43	ø	
					(7.27)	(46.79)	(7.33)		
VIII	Et3Ge	26	180-182		14.01	53.31	6.31	18.12	
					(13.94)	(53.65)	(6.56)	(18.01)	• • • • • • • • • • • • • • • • • • • •
×	Et3Ge	92	93-94.5		7.93	46.73	7.08	19,99	
		- :			(4.80)	(46.85)	(7.30)	(20.22)	

a Si 7.28 (7.29) %; Ge 18.70 (18.85) %.

It is also well known that propynal reacts with ethyl diazoacetate [3], diazoethane [5] and phenyldiazomethane [6] to give mainly the products of dipolar addition of diazo compounds to the triple bond. These results indicate that the steric effect of the R group attached to the sp carbon of the acetylenic aldehyde plays an important role in the regioselectivity. Similarly, α-diazo-β-hydroxy esters (Vd, e) were formed along with mixtures of the corresponding pyrazoles in the reactions of triethylsilyl- and triethylgermylpropynal with methyl diazoacetate. The esters (Va—Ve) are new compounds and appear to be the first compounds isolated containing both a diazo group and a carbon—carbon triple bond. This type of reaction seems to be a convenient method for the preparation of diazo compounds containing four different functional groups. Compounds of type V probably result from initial attack of methyl diazoacetate at the carbonyl group of I to give the dipolar intermediate VI [cf. 1,7], which is followed by fast proton migration (Scheme 2)

SCHEME 2

The structure of the product Va could be assigned unambiguously from its IR and ¹H NMR spectra (Table 3). For example, the IR spectrum shows a strong band at 1700 cm⁻¹ attributable to the ester carbonyl group, a diazo group band at 2110 cm⁻¹, a strong band at 2250 cm⁻¹ [ν (C=C)], and a hydroxy group band at 3450 cm⁻¹.

Similar carbonyl addition reactions of diazo compounds with aldehydes and ketones have appeared in a number of reports [8]. For example, 2,2,2-trifluoro-diazoethane has been recently shown to react with F_3 CCHO and Cl_3 CCHO with an intermediate formation of relatively stable α -diazo- β -hydroxy alkanes [9]. Similarly, ethyl diazoacetate was shown to add to the carbonyl group of alloxane and related compounds [10].

The reaction of LiC(N₂)COR and LiC(N₂)COOR with aldehydes or ketones is the most fundamental one for the preparation of α -diazo- β -hydroxy ketones and β -hydroxy- α -diazo esters, respectively [11—16]. The best yields of α -diazo- β -hydroxy esters or α -diazo- β -hydroxy ketones are obtained when ethyl lithiodiazo-acetate (or lithiodiazoacetone) is generated in situ by adding to the mixture of aldehyde and ethyl diazoacetate (or diazoacetone) a solution of lithium diiso-propylamide in THF at low temperature [12,14,16—21]. The base-catalysed reaction between aldehydes and ethyl diazoacetate (or α -diazoacetone) has been reported to yield aldol-type products [18,22—26].

The reaction between trimethylsilylpropynal (Ic) and methyldiazoacetate does not give the corresponding β -hydroxy- α -diazo ester (Vc) but instead gives 3(5)-formyl-4-trimethylsilyl-5(3)-methoxycarbonylpyrazole (IIc) together with

SPECTROSCOPIC PROPERTIES OF THE COMPOUNDS PREPARED IN THIS INVESTIGATION TABLE 3

		он сно ин	10,48s 10,48s	10,48s	$4.90d^d$ $6.82d^d$		3.25s 5.48s		3.87s 5.54s		3.87s 5.54s	6.92s	7,98s	
12	1)	MeOCO	3.98s		3.92s	3,75s	3.80s		3.74s		3.77s	4.02s	3,86s	
TIGATION	NMR, chemical shifts (ppm)	MR3	0,96t 0,56m 0,95m	0,96s	1.05t	1,19s	0.88m	1.48m	2,20m 0.56m	0.95s	1.00m	1.15m	1.05m	
POUNDS PREPARED IN THIS INVESTIGATION	NMR, chemi	OH(NH)	3130 3150	3380	3370	3450	3380		3300		3360		3040	3230
PARED IN		Ω≡Ω				2250	2240		2190		2180			
UNDSPRE		NIIN				2110	2125		2120		2115			
IE COMPO		0=20	1730 1730	1720 1725	1725	1700	1695		1700		1710	1720	1735	
IES OF TH	n-1)	CH≃0	1685 1640											
o propert	IR, ν (cm ⁻¹)	Z II O	1525 1530	1540 1530	1530							1530	1625	
TABLE 3 SPECTROSCOPIC PROPERTIES OF THE COM	Compound		IIId	IIIe IVc	IVe	Va	Λρ		۸d		Ve	VII	VIII a	

 $a \ \delta$ (CII=) 7.61s, ν (Ph) 1690, b OSiMe $_{\rm s}$, $^c \delta$ (OCH $_{\rm 3}$) 3,26s, d $^3J=5.0$ Hz.

its isomer (IIIc) and a tricyclic hemiaminal (IVc). This reaction has been found to proceed in two stages: the addition of N₂CHCOOMe to the triple bond to give IIc and IIIc and subsequent cyclodimerization of pyrazole IIc to give hemiaminal (Scheme 1). Dimerization of this type is not unexpected. It is known [6] that 5-phenyl-3-formylpyrazole prepared from propynal and phenyl-diazomethane is converted to the corresponding hemiaminal. Moreover, it has been established that such cyclic hemiaminal formation might also occur with aldehydes of other N-substituted azoles such as 2-formylbenzimidazole [27—29], 3-formyl-1,2,4-triazoles [6,30,31]. With 5-formyl-5-methyl-1-pyrazolines the cyclodimerization is a highly stereospecific process [32—35].

The compounds IVc—IVe prepared by us represent the first organometallic hemiaminals. Their structure has been determined by elemental analysis and IR and NMR spectroscopy. Thus, the IR spectrum of 4H,9H-2,7-dimethoxy-carbonyl-3,8-bis(triethylgermyl)dipyrazolo[1,5-a:1',5'-d]pyrazine-4,9-diol (IVe) shows no formyl group, i.e. (C=O), absorption. There are, however, characteristic frequencies at 1530 cm⁻¹ (C=N), 1725 cm⁻¹ (the ester carbonyl group), and a broad band with a maximum at 3370 cm⁻¹ attributed to OH stretching vibrations [cf. 36]. The ¹³C NMR spectrum of the above compound containing a natural abundance of ¹³C shows signals at δ 166.1 (CO), 151.2, 145.6, 117.5 (C of pyrazole cycles), 79.0 (CHOH), 54.0 (OCH₃), 11.4 (CH₃ attached to C) and 7.4 ppm (CH₂). The mass spectrum of IVe shows the fragment ions at m/e 757 (M⁺ – CH₃) and 743 (M⁺ – C₂H₅).

On the other hand, the ¹H NMR and IR spectroscopic data indicate that a reversible conversion of the dimeric hemiaminal to a monomeric pyrazole IVe = IIe occurs in solution. We have found that the IVe = IIe reversible conversion occurs even at room temperature under appropriate polar conditions, as observed previously for 3-formyl-5-phenylpyrazole [6]. The data on the IVe: IIe ratio in organic solvents are summarized in Table 4. These data indicate that the pyrazole IIe content of solvents such as tetrahydrofuran, chloroform, dioxan does not exceed 28%. More polar solvents (methanol, hexamethylphosphoramide, dimethylsulfoxide) contain predominantly pyrazole IIe. Heating of IVe in dimethylsulfoxide to 60°C is accompanied by disappearance of the signals

TABLE 4
INFLUENCE OF THE REACTION CONDITIONS ON THE IVe : IIe RATIO

Solvent	IVe : IIe ratio (:	10%)	
	Determined by ¹ H NMR	Determined by IR	
tetrahydrofuran	90:10		
chloroform	80 : 20	87:13	
dioxan	80 : 20	72:28	
pyridine		74:26	
dimethylsulfoxide	25:75		
methanol	25 : 75		
hexamethylphosphoramide	10:90		

associated with the CH—OH group and an increase in the aldehyde proton signal intensity. Subsequent cooling of the solution to room temperature reduces the IVe: He ratio observed before heating. This demonstrates reversibility of the IVe \rightleftharpoons He conversion. In agreement with the above conclusion, hemiaminal IVe exhibits a dual reactivity. Thus, its reaction with excess hexamethyldisilazane in THF produces bistrimethylsilyl ether (VII) in almost quantitative yield. In contrast, the reaction of IVe with phenylhydrazine in diethyl ether provides a phenylhydrazone (VIII). Treatment of IVe with methanol in the presence of a catalytic amount of hydrochloric acid gave dimethylacetal (IX), which was isolated and characterized.

$$II = IV = PhNHNH2 Et_3Ge CH(OMe)_2$$

$$MeOOC N H (VIII)$$

$$II = IV = PhNHNH2 Et_3Ge CH(OMe)_2$$

$$MeOOC N H (VIII)$$

The hydrolysis of acetal IX by 5% HCl in THF leads to the dimer IVe.

Experimental

Triethylgermyl-, trimethyl- and triethylsilyl-propynals were prepared by oxidation of the corresponding acetylene carbinols with pyridinium chlorochromate [37].

The IR spectra were recorded on a UR-20 spectrometer. NMR spectra were

run on Tesla BS 487C (1 H, 80 MHz) or Varian XL 100/12 (13 C, 25.2 MHz) spectrometers. All chemical shifts are measured in ppm (δ scale) relative to hexamethyldisiloxane as internal standard. The 13 C NMR spectra are proton noise decoupled. Mass spectra were obtained on an MX-1303 spectrometer.

All starting materials and solvents were distilled before the experiments. All reactions were performed under an atmosphere of dry argon. Typical experiments are given below. Analytical data are collected in Table 2 and infrared and ¹H NMR spectroscopic data in Table 3.

Reaction of 1-(trimethylsilyl)prop-1-yn-3-al (Ic) with methyl diazoacetate A solution of 2.3 g (23 mmol) of methyl diazoacetate and 2.9 g (23 mmol) of propynal Ic in 6 ml of diethyl ether was kept in the dark at room temperature under argon for 45 days. The precipitate formed was filtered off and washed with ether to give 1.2 g (23.1%) of the product (IVc) in an analytically pure condition, m.p. 52–53°C. The filtrate was concentrated at reduced pressure to afford crude pyrazole (IIIc) as a thick yellow oil. The product was chromatographed on an alumina column with elution with CHCl₃ to yield 2.2 g (42.5%) of pyrazole IIIc as a white solid, m.p. 152–153°C.

Reaction of 1-(triethylgermyl)prop-1-yn-3-al (Ie) with methyl diazoacetate A solution of 3.11 g (14.6 mmol) of propynal Ie and 1.46 g (14.6 mmol) of methyl diazoacetate in anhydrous ether (10 ml) was kept under argon for 14 days at 20–25°C. The crystalline precipitate was filtered off, washed with ether and dried to give 1.6 g (35.0%) of the hemiaminal IVe. Workup as above and chromatography on an alumina column, eluting with chloroform, afforded diazo ester Ve as a yellow clear oil (1.3 g, 28.1%), n_D^{20} 1.5040. Continued elution gave the pyrazole IIIe as colorless crystals (1.5 g, 32.8%), m.p. 108–109°C.

Methyl 6,6-dimethyl-3-hydroxy-2-diazo-4-heptynoate (Va)

A solution of 3.1 g (31 mmol) of methyl diazoacetate and 3.4 g (31 mmol) of 4,4-dimethylpent-2-yn-1-al (Ia) in 15 ml of ether was kept in the dark at room temperature under argon for 60 days. The solution was filtered and concentrated to an oil. This oil was chromatographed on an alumina column with elution by chloroform to yield 5.7 g (87.8%) of diazo ester Va as a yellow viscous oil, n_D^{20} 1.4866.

4H,9H-2,7-Dimethoxycarbonyl-3,8-bis(triethylgermyl)dipyrazolo[1,5-a:1',5'-d]-4,9-bis(trimethylsiloxy)-pyrazine (VII)

A solution of 0.62 g (1.0 mmol) of cyclic hemiaminal IVe was treated with 2 ml of hexamethyldisilazane. The resulting mixture was kept under the same conditions for 1 day. The solvent and excess of (Me₃Si)₂NH were evaporated under reduced pressure, and the resulting solid was triturated with hexane to provide 0.74 g (ca. 100%) of disilylated hemiaminal VII as a solid, m.p. 89—90°C.

3(5)-Formyl-4-triethylgermyl-5(3)-methoxycarbonylpyrazolephenylhydrazone (VIII)

A mixture of 0.91 g (1.45 mmol) of hemiaminal IVe and 0.31 g (2.9 mmol)

of phenylhydrazine in anhydrous ether (10 ml) was stirred for 22 h at room temperature under argon. The solution was concentrated to give a solid. The solid was recrystallized from benzene/heptane to give phenylhydrazone VIII (0.31 g, 26.3%), m.p. 180—183°C.

3(5)-Dimethoxymethyl-4-triethylgermyl-5(3)-methoxycarbonylpyrazole (IX) A solution of 0.81 g (1.3 mmol) of hemiaminal IVe and a catalytic amount of concentrated hydrochloric acid in 10 ml of methanol was refluxed for 6 h and cooled. The resulting crystalline precipitate was collected by filtration and recrystallized from pentane to yield 0.71 g (75.5%) of dimethylacetal IX, m.p. 93–94.5°C.

Hydrolysis of dimethylacetal (IX)

A mixture of 0.41 g (1.1 mmol) acetal (IX) obtained as above and refluxed with 7 ml of 5% aqueous hydrochloric acid for 5 h. The resulting mixture was cooled and extracted with ether. The extract was washed with water, dried over magnesium sulfate and concentrated to a solid. Recrystallization from ether gave 0.14 g (39.4%) of hemiaminal IVe, m.p. 136—138°C.

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