PREPARATION OF [1,2,4]TRIAZOLO[5,1-c][1,2,4]TRIAZINE DERIVATIVES FROM 3,4-DIAMINO[1,2,4]TRIAZINE

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<u>Abstract</u> - 3,4-Diamino[1,2,4]triazine <u>1</u> reacts with diarylcarbodiimides to give the 7-arylamino-3-methyl-8 \underline{H} -4-oxo[1,2,4]triazolo[5,1- \underline{c}][1,2,4]triazines $\underline{2}$ in good yields. The reaction of the iminophosphorane $\underline{4}$ with thioureas leads directly to 2 in moderate yields.

As a part of an investigation on fused heterocycles, we have been engaged in the preparation of bridgehead nitrogen heterocycles which contain the [1,2,4]triazolo moiety. In this context we have previously reported that the reaction of functionalized N-aminoheterocycles with carbodismides leads to fused [1,2,4]triazoles 1.

We now describe two general methods for the preparation of some derivatives of the $\{1,2,4\}$ triazolo $[5,1-\underline{c}][1,2,4]$ triazine ring system. The methods described so far for the preparation of [1,2,4]triazolo $[5,1-\underline{c}][1,2,4]$ triazines can be classified in three groups: a) from 5-hydrazino[1,2,4]triazines and carbon-inserting reagents 2,3,4; b) from 3-hydrazino[1,2,4]triazoles and a-dicarbonyl compounds 5,6,7; and c) from [1,2,4]triazole-5-diazonium salts and active-methylene compounds 8,9,10. Our approach is based on the reaction of 3,4-diamino[1,2,4]triazine with diarylcarbodiimides to give N-heteroaromatic quanidines as highly reactive intermediates which undergo cyclization under the reaction conditions to give fused [1,2,4]triazoles.

Thus, the N-aminoheterocycle, 3,4-diamino-6-methyl-5-oxo-4,5-dihydro[1,2,4]triazine 1, readily available from diaminoguanidine hydrobromide and pyruvic acid 1, reacts with diarylcarbodiimides in dry toluene at reflux temperature for 24 h to give the corresponding 7-arylamino-3-methyl-8 $\underline{\text{H}}$ -4-oxo[1,2,4]triazolo[5,1- $\underline{\text{c}}$][1,2,4]-triazines $\underline{\text{2a-c}}$ as crystalline solids in good yields (70-97%) (Method A).

TABLE 1. Preparation of 7-Arylamino-3-methyl-8 \underline{H} -4-oxo[1,2,4]triazolo[5,1- \underline{c}][1,2,4]-triazines $\underline{2}$.

Entry	Ar	Mp(°C)	Yield(%)		Found / Required			Molecular
			Α	В	С	н	N	Formula
2a	^С 6 ^Н 5	298-300	70	30	54.57 54.54		34.51 34.69	C ₁₁ H ₁₀ N ₆ O
2b	p-H ₃ C-C ₆ H ₄	337-339	75	36			32.68 32.79	C ₁₂ H ₁₂ N ₆ O
2c	р-H ₃ CO-С ₆ H ₄	334-335	97	41	53.12 52.94		30.68 30.87	^C 12 ^H 12 ^N 6 ^O 2
2 đ	p-Cl-C ₆ H ₄	335-337	55	33	47.61 47.75		30.42 30.37	C ₁₁ H ₉ C1N ₆ O
2e	p-Br-C ₆ H ₄	328-330	57	45	41.22		26.09 26.17	C ₁₁ H ₉ BrN ₆ O

The ir spectra of compounds 2 show a strong absorption band in the region 3336-3296 cm⁻¹ due to the NH group, and other one at 1699-1687 cm⁻¹ attributable to the C=O bond. In the 1 H-nmr spectra the chemical shift of the 2 C₃-methyl group is characteristic at 6 2.00 ppm. The mass spectra show the expected molecular ion peaks, and other significative is due to the fragment $[Ar-NH-CN]^{\frac{1}{2}}$.

Compound $\underline{1}$ reacts with bis(p-chlorophenyl)- and bis(p-bromophenyl)carbodiimide to give guanidines $\underline{3d}$, as crystalline solids in good yields (78-80%). The structure of compounds $\underline{3}$ has been tentatively assigned on the basis of the chemical shift of the amino group at position 4 by comparison with the chemical shift of the same group in the ${}^1\text{H-nmr}$ spectra of $\underline{1}$ and $4\text{-amino-6-methyl-3-methylamino-5-oxo-4,5-dihydro[1,2,4]triazine, recorded in DMSO-d solutions.$

By heating in ethanolic solution, compounds $\underline{3}$ are converted into $\underline{2d,e}$ in high yields.

On the other hand the N-aminoheterocycle 1 reacts with triphenylphosphine dibromide in the presence of triethylamine to give the iminophosphorane 4 in excellent yield (94%). The ir spectrum of compound 4 shows two bands at 3318 and 3239 cm⁻¹ due to the amino group, and a strong band at 1659 cm⁻¹ attributable to the carbonyl group. In the 1 H-nmr spectrum recorded in CDCl $_3$ the methyl and the amino groups appear as singlets at 6 2.40 and 6 6.10 ppm respectively. The mass spectrum shows the expected molecular ion peak, and other significative appears at m/z 318 due to the fragment $[Ph_3P=N-NCO]^{\frac{1}{2}}$. The ^{31}P -nmr study confirms that the triphenylphosphoranylideneamino group is attached at position 4 of the [1,2,4]triazine ring: the value found for the compound 4 is similar to the iminophosphorane derived from 4-amino-6-methyl-3-methylthio-5-oxo-4,5-dihydro-[1,2,4]triazine and higher than iminophosphoranes derived from anilines 12 and C-aminoheterocycles.

Iminophosphorane $\underline{4}$ reacts with arylthioureas in dry toluene at reflux temperature for 24 h to afford the corresponding triazolotriazines $\underline{2}$ in moderate yields (30-45%) (Method B). We believe that the conversion $\underline{4} - \underline{2}$ involves initial aza-Wittig reaction between the iminophosphorane $\underline{4}$ and the thiourea to give a non-isolated intermediate guanidine $\underline{13}$ which undergoes cyclization and elimination of amine to give 2.

EXPERIMENTAL

Melting points were obtained on a Kofler hot-stage apparatus and are uncorrected. Ir spectra were run using NaCl plates on a Nicolet FT-5DX spectrophotometer in Nujol emulsions. $^1{\rm H}$ and $^{31}{\rm P-mmr}$ spectra were obtained on a Varian FT-80 spectrometer using tetramethylsilane as external reference for $^1{\rm H}$ and 85% ${\rm H_3PO_4}$ as external reference for $^{31}{\rm P-mmr}$. The EI-mass spectra were obtained with a Hewlett-Packard 5993 C spectrometer. Elemental analyses were performed with a Perkin-Elmer 240 C instrument.

3-Amino-6-methyl-5-oxo-4-triphenylphosphoranylideneamino-4,5-dihydro[1,2,4]triazine $\underline{4}$. Bromine (1.44 g, 9 mmol) in dry benzene (15 ml) was added dropwise to a stirred solution of triphenylphosphine (2.36 g, 9 mmol) in dry benzene (20 ml) at 0-5°C under nitrogen. The mixture was stirred for 1 h and then allowed to warm to room temperature. A solution of triethylamine (1.82 g, 18 mmol) in dry benzene (20 ml) and 3,4-diamino-6-methyl-5-oxo-4,5-dihydro[1,2,4]triazine 1 (1.27 g, 9 mmol) were added to the above mixture; after heating under reflux for 24 h, the deposited triethylammonium bromide was separated by filtration and the filtrate was concentrated to dryness to afford a crude product which was recrystallized from chloroform/cyclohexane (1:1, v/v) to give the iminophosphorane 4 (3.39 g, 94%) as colourless prisms, mp 214-216°C (Found: C, 65.71; H, 4.96; N, 17.67. $C_{22}H_{20}N_5OP$ requires C, 65.83; H, 5.02; N, 17.45); ir v max (Nujol) 3318, 3239, 1659, 1545, 1500, 1404, 1206, 1109, 1087, 1008, 959, 923, 753, 725 cm⁻¹; ¹_{H-nmr δ} $(CDCl_3)$ 8.3-7.4 (15H,m), 6.12 (2H,s), 2.35 (3H,s); m/z(%) 401(M⁺, 35), 400(67), 318(15), 304(18), 303(27), 301(38), 288(26), 276(12), 262(23), 261(16), 240(10), 200(15), 185(21), 183(100), 108(10).

General Procedure for the Formation of 7-Arylamino-3-methyl-8H-4-oxo[1,2,4]-triazolo[5,1-c][1,2,4]triazines 2. Method A. To a solution of 3,4-diamino-6-methyl-5-oxo-4,5-dihydro[1,2,4]triazine $\frac{1}{2}$ (1 g, 7 mmol) in dry toluene (25 ml) the appropriate diarylcarbodiimide (7 mmol) was added. The reaction mixture was stirred at reflux temperature for 24 h; after cooling the precipitated solid was collected by filtration, dried and recrystallized from ethanol to give $\frac{2}{2}$ as colourless prisms.

In the cases of bis(p-chlorophenyl)- and bis(p-bromophenyl)carbodiimide the reaction products were found to be the corresponding heteroaryl guanidines 3.

TABLE 2. Spectral Data of Compounds 2.

Compound	Ir v (cm ⁻¹)	1 _{H-nmr} a 6 (ppm)	Ms ^b <u>m/2</u> (%)
	3336,1699,1619,	10.00(1H,s);	242(M ⁺ ,96),144(18),119
	1580,1359,742,	8.0-7.0(6H,m);	(18),118(100),104(12),91
	691.	2.00(3H,s).	(19),77(40),68(11),65
			(16),51(29).
2b	3330,1687,1648,	10.05(lH,s);	256(M ⁺ ,66),158(24),133
	1614,1574,1557,	8.1-7.0(5H,m);	(21),132(100),131(34),118
	1518,826,747,	2.00(6H,s).	(24),117(15),106(14),92
	719.		(21),91(89),77(29),65(35).
2c	3296,3183,1687,	10.00(1H,s);	272(M ⁺ ,100),257(43),159
	1631,1585,1512,	8.0-7.0(5H,m);	(20),148(28),133(89),122
	1246,1223,1178,	4.13(3H,s);	(26),105(27),92(21),77
	1030,798,742.	2.10(3H,s).	(22),65(21).
2d	3336,1687,1625,	10.05(1H,s);	278(M ⁺ +2,31),276(M ⁺ ,100),
	1609,1563,1042,	8.1-7.1(5H,m);	154(31),152(99),140(6),
	1008,826,804.	2.05(3H,s).	138(16),128(4),126(12),
			113(11),111(36),83(21).
2e	3336,1687,1636,	10.10(1H,s);	323(M ⁺ +2,14),321(M ⁺ ,19),
	1614,1585,1552,	8.1-7.5(5H,m);	198(77),196(87),184(15),
	1087,1008,838,	2.05(3H,s).	182(15),172(11),170(10),
	799,744.		157(37),155(35),143(30),
			117(51),91(67),90(99),83
			(37),76(48),75(57).

 $^{^{\}rm a}$ Obtained as solutions in DMSO-d $^{\rm 6}$. $^{\rm b}$ Recorded at 70 eV.

When an ethanolic solution of $\underline{3a}$ or $\underline{3b}$ was heated at reflux temperature for 5 h, the corresponding triazolotriazine $\underline{2d}$ or $\underline{2e}$ were obtained in 70% or 72% yield respectively.

Method B. To a solution of 3-amino-6-methyl-5-oxo-4-triphenylphosphoranylidene-amino-4,5-dihydro[1,2,4]triazine $\underline{4}$ (2 g, 5 mmol) in dry toluene (50 ml) the appropriate diarylthiourea (5 mmol) was added. The reaction mixture was stirred at reflux temperature for 24 h. Similar work-up to the above method led to pure compounds $\underline{2}$.

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