578. Organic Complex-forming Agents for Metals. Part II.* Preparation of 5-Hydroxy- and 5:8-Dihydroxy-quinoxalines and Related Compounds.

By E. S. LANE and C. WILLIAMS.

5-Hydroxy- and 5:8-dihydroxy-quinoxalines containing a variety of 2- and 3-substituents have been prepared by dealkylation of the corresponding 5-methoxy- and 5: 8-diethoxy-quinoxalines, and characterised as the acetyl derivatives.

This paper reports the preparation of further new complex-forming agents containing the substituents recorded in Part I 1 but based on the less basic quinoxaline ring system. Some compounds of this series have been previously described (for references see Albert and Hampton,² and Adachi³). King, Clark, and Davis⁴ noted that 5:8-dihydroxyquinoxalines possessed indefinite melting points and gave inconsistent analyses. We confirmed this observation and ascribe it partly to the pronounced tendency of quinoxaline derivatives to form molecular complexes, to which we have drawn attention previously, and to the ease with which they are oxidised to p-quinonoid derivatives. To obtain the analyses reported below, repeated purification of the compounds was necessary by protracted recrystallisation from selected solvents or, preferably, by vacuum-sublimation. King 4 et al. used an isomeric mixture of diamines as starting material, but like Adachi we preferred to use pure o-diamines as the primary intermediates and this undoubtedly facilitated the purification of these compounds.

Well-established methods were used for the preparation of the 2:3-disubstituted 5-methoxy- and 5:8-diethoxy-quinoxalines from the appropriate o-diamines and α -diketones (an improved preparation of di-α-pyridyl diketone is described). Similar compounds singly substituted in the 2-position were prepared from the corresponding glyoxal derivative. 2: 3-Dihydroxyquinoxalines were prepared from the parent diamine and oxalic acid, and were smoothly converted into the 2:3-dichloro-compounds by phosphorus oxychloride or phosphorus oxychloride-dimethylaniline.⁵ For dealkylation either aluminium chloride in benzene (method A) or boiling aqueous hydrobromic acid (method B) was used, the latter being usually reserved for aryl-substituted quinoxalines.

The hydroxyquinoxalines were characterised as their O-acetyl derivatives prepared by use of acetic anhydride, optionally in the presence of pyridine. The base was omitted when acetylating 2:3-dichloro-hydroxyquinoxalines owing to side reactions with the reactive chlorine atoms. Similarly, it was not possible to use hydrobromic acid for the dealkylation of 2:3-dichloroquinoxalines because of replacement of the chlorine atoms by hydroxyl groups.

The 5-hydroxy- and 5:8-dihydroxy-quinoxalines exhibit chelating properties to a varied extent. Preliminary accounts have been given by Irving and Rossotti 6 and by Kawa, Kimura, and Furahata.7 Their main disadvantages are their low solubility in water and alcohol and their ready oxidisability.

Since hemipyocyanine (1-hydroxyphenazine) is a fungicide a representative selection of our analogous quinoxalines was kindly tested by Mr. W. H. Read of the Glasshouse Crops Research Institute, who reported, however, that they displayed only low fungicidal activity.

EXPERIMENTAL

Determination of the Equivalent Weight of Quinoxalines.—The methods used in Part I were applicable to the compounds described herein.

- * Part I, J., 1956, 569.
- ¹ Lane and Williams, J., 1956, 569.
- Albert and Hampton, J., 1952, 4985.
 Adachi, J. Chem. Soc. Japan, 1955, 76, 311.
 King, Clark, and Davis, J., 1949, 3012.
 Landquist, J., 1953, 2816.
 Irving and Rossotti, Analyst, 1955, 80, 245.

- ⁷ Kawa, Kimura, and Furuhata, Proc. Japan Acad., 1953, 29, 344.

Required (%)	z		1	14.8		12.2	17.8	•	14.0	13.1	0.71		I	16.1	13.0	18.7	9. 4.	1.01	14·0 13·1	·		I	13.0	11.0	16.4	8.2	12.3	11.6	10.9	
	Н]	I	I	2.6	4.5		0.9	6.5	1		I	5.7	ნ- <u>I</u>	4·0 '	4·7	4.0	6.55			1	5.55	2.3	4 ∙I	4.7	5.3		6.25	
	ပ		1	I]	47.2	72.6		72.0	72.9	2		I	0.69	44.65	75.0	9 9 1 1	0.17	0.25 23.0 23.0 23.0) }]	66.7	46.7	I	9.44	68.3	69.4	70.3	
Found (%)	Z]	14.8	J	12.0	17.5		14.2	13.0	9		I	15.75	12.9	18.7	9.7	1.01	13.2))		1	12.6	11.0	16.3	8.1	12.1	11.7	11.1	
	H		j]	ĵ	5 .8	4.4	1	0.9	6.9	к -		I		œ i	ဗ ဗ	6	0.0	6.1 6.4	1		1	5.65	2.25	4 ∙I	4.7	5.6	2.1	တ်	
ŭ	ပ		I]	J	47.6	72.3	1	71.9	72.5	2		I	8.89	45.2	71.1	90.0	0.01	73.0))]	9.99	47.1	1	77.8	68.2	0.69	8.69	
	Formula	noxalines.	I	C,1H,0N,	See Text	C,HON,CI,	C',H,'ON,	: :	C, H, ON,	C ₁₃ H ₁ ON ₂	~141116 ~1.2	oxalines.	j	Clot Hook	C.H.ON.C.	C18H13ON		C111110012	Cirtion,	8 87 07 -	oxalines.	J	$C_{18}H_{12}O_{2}N_{2}$	C10HO2N2C1,	C20 H 14 O2 N4	C23H16O2N2	$\mathrm{C_{13}H_{12}O_2N_2}$	C14H14O2N2	C15H16O2N3	
	Appearance *	TABLE 1. 5-Methoxyquinoxalines.	I	Plates	Powder	Yellow needles	Powder	Plates	ı	Pink prisms Vellow needles		. 5-Hydroxyquinoxalines.	J	Yellow cubes	Yellow needles	Fawn needles	Needles	rawn powder	Yellow powder Fawn powder	4	3. 5-Acetoxyquinoxalines.	Solid	Cubes	Yellow needles	Cream powder	Powder	Cream powder	Yellow powder	Needles	
Equiv. wt.	Reqd.		I	188	96	1	I	I	l	1 1		TABLE 2.	J	l	1 ;	300	867	190	214		TABLE 3	J	ļ	1	I	J	I	j	1	
Equi	Found		I	192	86	1	1	I	1	П			l	I	1	300	302	19/	212	!		J]	1	1	ļ	I	ļ	Ī	
	M. p.		100	121	272 - 274	144.5—145	175.5	1907	99.5 - 102	$95 \\ 110.5 - 114$	****		101	146	144	176—177	133—134	139—140	$118.5 \\ 139 - 140$			101	103	90—94	158 - 159	142	87—83	94—95	83	* Colourless unless otherwise stated.
	3-Subst.		, H	Me b	НО	ប៉	2-Py d, c	Ph	ethylene	: 3-Tetramethylene : 3-Pentamethylene			н	Me	: :	2-Py 4	Fn	ernylene	: 3-1 etrametnylene : 3-Pentamethylene			, H	Me	ວ ^ເ	2-Py a	Ph	ethylene	2:3-Tetramethylene	amethylene	urless unless o
	2-Subst.		Н	Me	НО	ರ	2-Py	Ph	2:3-Trim	2:3-Tetra			Н	We G	J :	z-Py	Fn 9 : 9 Tuim	11111-6:2	2:3-1 etr. 2:3-Pent			Н	Me	ວ¦	2-Py	r.	2:3-Trim	2:3-Tetr	2:3-Pent	* Colc

* Colourless unless otherwise stated.

* King, Clark, and Davis 4 reported m. p. 100—101. * Landquist and Stacey (J., 1953, 2822) obtained m. p. 118. * Found: Cl, 30.9. Read., 31.0%. * Fy = pyridyl. * Perchlorale, yellow, m. p. 254* (decomp.) (Found: N, 9.8; Cl, 12.4. Cl, 14.10, N, 31.04.) (Found: N, 9.8; Cl, 12.4. Cl, 14.10, N, 31.04.) (Found: N, 9.8; Cl, 12.4. Cl, 14.10, N, 9.8; Cl, 12.3%). * Medola and Eyre (J., 1902, 281, 992) give m. p. 191. * King et al.* reported m. p. 100—101°; Sorkin and Roth (He.c. Chin., Acta, 1951, 34, 27) reported m. p. 102—103°. * Did not titrate quantitatively in ethylenediamine owing to interaction of Cl with solvent. * King et al.* reported m. p. 106.5—108°. * Found: Cl, 27.9. Reqd., 27.6%. * Nietzki and Rechberg give m. p. 127° and 163° respectively. * King et al.* obtained m. p. 230°; Adachi * obtained m. p. 230°; Adachi * obtained m. p. 225° (decomp.) (Found: N, 7.1; Cl. 8-8. Cl, H1, Ob.N2Cl requires N, 7.2; Cl, 9-1%).

(%	z		12.8	11:4	0 0 0 0	15.1	9.5	10.8	12.1	10.85	9.65	8.6 8.6			14.7	19.1	1.7.1	11.0	2 i	10.7	10.6	13.0	10.01	8.9		;	11.4 10.0	200	14.0	8.7	6.4	8. 6.	တ တ		2
Required (%)	Н		1	7.3	4.2	5.4	6.1	7.7	6.9	7.0	7.65	7.7			5.5) F	-] (4. r		0 0	# 10 20 30 30 30 30 30 30 30 30 30 30 30 30 30	- -	4.5		•	4·1		4. 0.	4.3	j	4·9		7.0	÷
	ပ]	68·3	50.5	71.0	73.5	69.5	67.5	8.69	66.2	70.0	٠		63.9	2 5	0.14	6	9.00	7.70	4.10	66.7	67.0	76.4		ì	0.8.0 61.3	45.7	0.99	67.1	1	65.9	04.0	79.4	, 1
(%)	z		12.6	11:3 10:8	101	15.0	9.25	10.65	9.11	10.9	9.55	9.7			14.9	11.7	17.7	- 1	11.7	0.01	14.9	13.5	13.6	8.85			11.2	20.0	14.0	8.75	6.7	7.6	9.45	7 6.7	-
	н		1	7.3 5.0	. 4	5.4	6.5	7.7	7.5	6.75	7.65	7.95			6.5	9 -	0.1	9	4. 2				6.45	4.4		l.	4 r.	9 6	4.4	4.5	1	4·9	1.c	1.5	>
	ပ		1	68.2 58.1	49.9	8.02	73.4	69.1	8.99	69.7	1.99	69.5			63.35	41.0	0.14	9	0.07	0. * .0	64.0	6.4.0	68.4	9.92		6	29.5	45.7	65.5	67.1	1	62.6	64.3	71.7	:
	Formula	quinoxalines.	C12H14O2N2		C.H.O.N.CI.	C22H202N	CIRHISON,	C15H2002N2	C13H 16O2N	C15H18O2N3	C16H20O2N2,H2O	$\mathrm{C_{17}H_{22}O_2N_2}$		inoxalines.	C. H. O.N.			01811120214				C1111100217	O.H.O.N.	$C_{20}H_{14}O_{2}N_{2}$	noxalines.	¥ 0 # 0		CIATION OF THE COLUMN OF THE C	C23H1604N4	C18H1404N2	C15H16O4N2	C15H1404N2		C171180418	~24**1~4**3
Equiv. wt.	Appearance *	4. 5:8-Diethoxyquinoxalines	Yellow plates	Needles	Orange plates	Yellow needles		Cream powder		Yellow needles	Fawn powder	Yellow needles	Yellow prisms	5:8-Dihydroxyquinoxalines.	Vellow plates	Orange plates	Orange plates	Organigo powder	Orange powder	Orange needles	Organies needles	Orange necures	Vellow needles	Orange plates	5:8-Diacetoxyquinoxalines.	C	Cream plates	:	Needles "	Needles	Orange prisms	Yellow needles	r ellow needles	Needles	
	Reqd.	TABLE	J	11		1		j]	I]	1	l	TABLE 5.	I			ן נ	119	707	0	108	}	П	TABLE 6.				1	1	I	I	l]]	
nba	Found		j	1 1		1	1	1	1	I]		1		J			011	8118	104	00	107.5	;					!]	I]	1	I	l]]	
	M. p.		101.5-101°	127·5 971	199—199-5	175	114 - 116	47—49	70—73	$122 - 122 \cdot 5$	117	157	165		930—933	151 159	040 050	007-047	9/1	169	2016	198—199	177—178			000	200-202 $163.5-164$		285—287.5	157 - 158	40—45	190—192	277-278	130—138 228	
	3-Subst.		н	Me K		2-Py 4	, Н	Ħ	H	: 3-Trimethylene	amethylene	tamethylene	Ph '		Me m	*	, i.d.	7-1 y	##	4	TI	ietiiyieiie amethviene	amethylene	: o-rentamethylene Ph		;	r o) 	2-Py d	, Н	μ	nethylene	amethylene	: 3-Pentametnylene	
	2-Subst.		Н	Me	35	2 -Pv	Ph	돲	Me	2:3-Trin	2:3-Tetr	2:3-Pent	Ph		Me	ž į	ا 5 د	7-1-7 7-1-7	r.	£;	Me	2:3-1rm 9:3-Tetr	9 · 3 · Den	z: 3-ren Ph		;	ΗŞ	ž C	2-Pv	Ph	Pr	2:3-Trin	2:3-Tetr	2:3-Pen	rn L

Di-2-pyridyl Diketone (α-Pyridil).—The recorded method of oxidising α-pyridoin with fuming nitric acid, followed by heating on a water-bath, gave only a small yield of the required product. By maintaining the reaction mixture at -30° during the oxidation and eliminating the heating, the yield was increased to over 90%.

Preparation of 2:3-Disubstituted 5-Methoxy- and 5:8-Diethoxy-quinoxalines.—Equimolecular quantities of the appropriate diketones and o-diamines (see Part I) were refluxed together in ethanol for 1 hr. The mixture was poured into water, and the insoluble quinoxaline filtered off, distilled *in vacuo* when possible, and recrystallised.

The compounds are listed in Tables 1 and 4.

- 2-Substituted 5: 8-Diethoxyquinoxalines.—These were prepared as above, from the appropriate glyoxal in place of the α -diketone (see Table 4).
- 2: 3-Dihydroxy-5-methoxyquinoxaline.—Equimolecular quantities of 2: 3-diaminoanisole and oxalic acid were refluxed together for 12 hr. in 4N-hydrochloric acid. The mixture was neutralised with ammonia, and the precipitated base filtered off. It formed colourless crystals (from ethylene glycol), m. p. 272-274°. The equivalent weight of this compound was determined in ethylenediamine and gave a satisfactory value but microanalyses were unsatisfactory (Found: C, 50·35; H, 5·6; N, 12·8%; equiv., 98. C₉H₈O₃N₂ requires C, 56·25; H, 4·2; N, 14.6%; equiv, 96). Satisfactory microanalyses were obtained from its derivatives (see below).
- 5: 8-Diethoxy-2: 3-dihydroxyquinoxaline was similarly prepared from oxalic acid and 2: 3-diaminoquinol diethyl ether (see Table 4).
- 2: 3-Dichloro-5-methoxyquinoxaline.—2: 3-Dihydroxy-5-methoxyquinoxaline (4 g.) was refluxed for 1 hr. with dimethylaniline (5 ml.) and phosphorus oxychloride (20 ml.). Excess of phosphorus oxychloride was removed in a vacuum and the residue poured on crushed ice (200 g.). The base was filtered off and dried. It formed pale yellow needles, m. p. 144.5— 145° (from benzene) (see Table 1).
- 2: 3-Dichloro-5: 8-diethoxyquinoxaline.—This was similarly prepared from 5: 8-diethoxy-2: 3-dihydroxyquinoxaline (16 g.) and phosphorus oxychloride (400 ml.) (see Table 4).
- 2:3-Disubstituted 5:8-Dihydroxyquinoxalines.—(A) The 5:8-diethoxy-compound was refluxed for 3 hr. with anhydrous aluminium chloride (2.4 equivs.), in benzene. Water was then added and the mixture extracted with hot benzene. The benzene extract was evaporated to small bulk and the material which separated was recrystallised.
- (B) The 5: 8-diethoxy-compound was refluxed for 6 hr. in hydrobromic acid (48%). After cooling, the mixture was filtered and the solid material hydrolysed with warm water, washed with sodium hydrogen carbonate solution and with water, and recrystallised.
- 2: 3-Disubstituted 5-Hydroxyquinoxalines.—Analogous procedures to the above, but using 1.2 equivs. of anhydrous aluminium chloride, were used.
- 2: 3-Disubstituted 5-Acetoxy- and 5: 8-Diacetoxy-quinoxalines.—The hydroxy-compounds were refluxed in acetic anhydride, excess of reagent was removed under reduced pressure, and the residue recrystallised from ethanol (see Tables 3 and 6).
- 10: 13-Dihydroxydibenzo[a,c]phenazine.—10: 13-Diethoxydibenzo[a,c]phenazine 8 (2 g.) was refluxed with 48% hydrobromic acid (60 ml.) for 6 hr. On cooling, the deposited solid was hydrolysed by boiling water to an orange powder (1.6 g.). Recrystallisation from benzyl alcohol gave the phenolic base as an orange-red powder, m. p. 280—283° (Found: C, 76.6; H, 3.9; N, 8.8. $C_{20}H_{12}O_2N_2$ requires C, 76.9; H, 3.8; N, 9.0%). A prior preparation attributed to Kawai and Kosaka 9 is mentioned by Kawal, Kimura, and Furahata 7 but without details. The diacetate was obtained as yellow-green needles, m. p. 273—273-5° (Found: C, 72.5; H, 4.2; N, 6.85. $C_{24}H_{16}O_4N_2$ requires C, 72.7; H, 4.0; N, 7.1%).

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⁸ Nietzki and Rechberg, Ber., 1890, 23, 1212.

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