Synthesis and Hypotensive Properties of New 4-Aminoquinolines

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Received March 10, 1971

A series of 6,7-dimethoxy-4-(substituted amino)quinolines, several 6,7-dimethoxy-4-aminoquinolinium iodides, and some miscellaneous 4-substituted quinolines were synthesized and evaluated for hypotensive activity in dogs. Several of the simple 4-(alkylamino)-6,7-dimethoxyquinolines exhibited good hypotensive activity, equal to that of the parent 4-amino-6,7-dimethoxyquinoline (1).

Several 4-aminoquinolines (1, 54–57, Table I), a variety of 6,7-dimethoxy-4-(substituted amino)quinolines¹ (2–26, 27–29, 30–32, 33–35), four 6,7-dimethoxy-4-aminoquinolinium iodides (50–53), and a few 4-phenoxy-, 4-thio-, 4-hydrazino-, and 4-chloroquinolines (36–41, 42, 43, 46) were synthesized and screened for hypotensive activity in anesthetized dogs in the present work. Previously the hypotensive activity and the mechanism of action of the parent 6,7-dimethoxy-4-aminoquinoline HCl (1) and of 6,7-dimethoxy-4-veratrylideneaminoquinoline (33) were reported by Buckley, *et al.*²

Chemistry.—The 4-aminoquinolines described in this paper were prepared primarily by halogen displacement of the corresponding 4-chloroquinolines with amines in phenol (methods A1, A3, A4). Compd 1 was obtained through the reaction of 6,7-dimethoxy-4-chloroquinoline (47)³ with phenolic NH_3 .⁴ The 6,7-dimethoxy-4phenoxyquinoline (36, free base) was obtained by refluxing 47 in phenol (method B); 36 (free base) was easily converted to 1 (free base) by heating in excess NH_4OAc (method A2). This latter reaction indicates that 36 is a possible intermediate in the "phenolic reaction" of 47 and NH_3 to give 1.

The product of the reaction between 47 and *p*-aminophenol was 4-(*p*-hydroxyanilino)quinoline (26). The structure of 26 was verified through independent synthesis of the other possible isomer, the *p*-aminophenoxy ether (38), through catalytic reduction of 37.

The 4-mercapto- and 4-hydrazino-6,7-dimethoxyquinolines (**39**, **41**) were prepared through displacement reactions of **47**; methylation of **39** with Me_2SO_4 gave 6,7-dimethoxy-4-methylthioquinoline (**40**).

Various derivatives (22, 30–32, 33–35) of 1 (free base) were prepared. Examples are acetylation with Ac_2O to give 30; addition of ethyl isocyanatoacetate to give 31; condensation with veratrylaldehyde to give 33, followed by catalytic reduction to give 22.

Three 1-alkyl-6,7-dimethoxy-4-aminoquinolinium iodides (50-52), Table I) were synthesized through (1) alkylation of the 4-chloroquinoline with alkyl iodides

(3) B. Riegel, G. R. Lappin, B. H. Adelson, R. I. Jackson, C. J. Albisetti, Jr., R. M. Dodson, and R. H. Baker, J. Amer. Chem. Soc., **68**, 1264 (1946).



followed by displacement with amines, or (2) amine displacement of the 4-chloroquinoline followed by alkylation (Scheme I). While the ethylation of 4-chloro-6,-7-dimethoxyquinoline (47) with EtI was easily achieved, the ethylation of 42 for preparation of the projected intermediate 49 failed to occur. Although the synthesis of 3-carboxyl-1-ethyl-6,7-dimethoxy-4methylaminoquinolinium iodide (52) was carried out through alkylation of the 4-methylamino-3-quinolinecarboxylate (44) with EtI and base (NaOH), the nmr spectrum indicates that 52 may be contaminated with the product of ethylation at the 4-CH₃NH grouping.

4-Amino-6,7-dimethoxy-1-methylquinolinium iodide (50) was prepared by the reaction of 1 (free base) with MeI. Passage of 50 over a basic ion-exchange resin afforded the corresponding chloride 53. The physical properties of 53 were compared with those of the isomeric 6,7-dimethoxy-4-methylaminoquinoline hydrochloride (2) of known structure; the 2 compounds were different. The nmr spectrum of 2 shows that the (NCH₃) protons are split by the (NH) proton of the (4-CH₃NH) substituent; in the case of 53 the (NCH₃) protons shown only a singlet.

The ring closure of **19** (Scheme II) in PPA gave 2,3dehydro-8,9-dimethoxybenzo[*h*]-1,6-naphthyridin-4-

 ^{(1) (}a) F. F. Ebetino and G. C. Wright, U. S. Patent 3,272,824 (1966);
 Chem. Abstr., 63, P589b (1965). (b) A. Winterstein, U. S. Patent 3,272,806 (1966);
 Chem. Abstr., 65, P18567a (1966).

⁽²⁾ B. S. Jandhyala, G. J. Grega, and J. P. Buckley, Arch. Int. Pharmacodyn., 167, 217 (1967).

⁽⁴⁾ A modification of the procedure for the synthesis of 4-aminoquinaldines, by O. G. Backeberg and J. L. C. Marais, J. Chem. Soc., 381 (1942).



(1H)-one (58). This is similar to the reaction of 4aminoquinaldine and ethyl trifluoroacetoacetate in PPA to give the completely aromatic 5-methyl-2-trifluoromethylbenzo [h]- 1,6- naphthyridin- 4-ol.⁵

That ring closure of 19 did not occur on the benzo ring was established by the nmr spectrum, which contained 3 singlets for aromatic protons. The spectrum of a benzo ring closure product would exhibit a pair of aromatic ortho proton doublets, not observed in the spectrum of 58.

A second tricyclic compound, 7,8-diethoxy-2,3-dihydropyrrolo-1*H*-[3,2-g]quinoline HCl (**59**), was obtained through ring closure of 4-chloro-3-(2-chloroethyl)-2-methyl-6,7-diethoxyquinaldine (**43**) with NH₃ in phenol (Scheme II). Compd **43** was synthesized through the reaction of 3,4-diethoxyaniline and 2-acetylbutyrolactone⁶ to give the intermediate 1-(tetrahydro-2-oxo-3-furyl)ethylidene-2,4-diethoxyaniline (**60**), followed by chlorination of **60** with POCl₃ to give **43**. The 3,4-diethoxyaniline was obtained by catalytic reduction of 3,4-diethoxynitrobenzene.⁷

Pharmacology.—All compounds were evaluated for hypotensive activity in barbiturate-anesthetized mongrel dogs. Blood pressure was recorded from a cannulated femoral artery. Experimental materials were administered iv in H_2O when the solubility permitted or ip when the solubility was such that iv administration was not feasible. The hypotensive activity was evaluated on the basis of the maximum decrease in blood pressure and the duration of action of the respective

(6) Purchased from Columbia Organic Chemical Co.



effective doses. Generally the lowest dose is cited which caused the highest rating using the following classifications: minimal activity, + (<40% decrease

⁽⁵⁾ A. S. Dey and M. M. Joullie, J. Heterocycl. Chem., 2, 120 (1965).

⁽⁷⁾ D. F. Page and R. O. Clinton, J. Org. Chem., 27, 224 (1962).

$ \begin{array}{cccccccccccccccccccccccccccccccccccc$		2	Method	Yield,	Recrystn ^a solvent	qi.Jo u.M	L'Annual a		Hypote Dose, 	ansive activ	ty in dogs
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	II N		politatat	0/	solvent	Mp, C	Formula	Analyses	mg/kg	Route	$\operatorname{Rating}^{c}$
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	1121		A1 A2	97 97	ŗ	273-276 $273-276$	$C_{II}H_{I2}N_2O_2 \cdot HCI \cdot H_2O$	C, H, N, CI	10	Iv	+++
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	CH ₃ NH		$\mathbf{A1}$	31	К	265-267	$C_{12}H_{14}N_{3}O_{3}\cdot HCI$	C. H. N	06	L.v.	-
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	CH ₃ CH ₂ NH		$\mathbf{A3}$	60	Μ	236 - 237	ClaH ₁₆ N ₂ O ₂ ·HCl	C H N C	0 i 1	, ÷	} }
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	(CH3);CHNH		$A3^d$	16	ზ	240-242	$C_{14}H_{18}N_2O_2 \cdot HCl$	C, H, N	51	Ια	}- }∔ ∳∔
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	CH ₃ CH ₂ CH ₂ U ₃ N	Н	A3	62	Z	244 - 245	C ₁₄ H ₁₈ N ₂ O ₂ ·HCl	C. H. N. CI	10	Iv.	
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	CH ₃ (CH ₂) ₃ NH		$\mathbf{A3}$	34	М	206 - 208	$C_{15}H_{20}N_2O_2\cdot HCl$	C, H, N, CI	10	Ŋ	⊢ - 4 ├ -†
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	CH ₃ (CH ₂),NH		$\Lambda 3$	40	К	198 - 200	$C_{16}H_{22}N_2O_2 \cdot HCl \cdot 0.5H_2O$	C, H	10	Ιv	- + - +
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	NHCH2CH2NI	F	$A3^{e}$	47	Z	315-319 dec	C24H26N4O4 2HCl	C, H. N	20	In I	Presor
$ \begin{array}{lcccccccccccccccccccccccccccccccccccc$	HONH		$\mathbf{A4}$	35	S	270 - 274	$C_{11}H_{12}N_2O_3 \cdot HCl$	C,r H, N, Cl	10	Iv	+
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	HOCH2CH2NF	ł	A3	51	ſ	238 - 239	$C_{13}H_{16}N_2O_3 \cdot HCI \cdot H_2O$	C, H, N, CI	10	Iv	++
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	HO(CH ₂) ₃ NH		A3	43	L	235 - 236	C ₁₄ H ₁₈ N ₂ O ₃ ·HCl	C, H, N, CI	10	Iv	- - +
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	CH 3CHOHCH	2CH2NH	$\mathbf{A3}$	52	ඊ	200-201	$\mathrm{C}_{15}\mathrm{H}_{20}\mathrm{N}_{2}\mathrm{O}_{3}\cdot\mathrm{HCl}$	C. H. N. CI	10	Iv	+ + + -
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	HOCH2CHOF	ICH2NH	$\mathbf{A3}$	50	L	232 - 233	C ₁₄ H ₁₈ N ₂ O ₄ ·HCl·H ₂ O	C, H, N	10	Iv	- - - -
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	CH ₃ O(CH ₂) ₃ N	Η	$\mathbf{A3}$	99	റ	228 - 229	$C_{1_3}H_{20}N_2O_3 \cdot HCl$	C. H. N. CI	01	Iv	+ +
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	CH3CH2OCH	2CH2NH	$A3^{e}$	44	Μ	191 - 193	$\mathrm{C_{15}H_{20}N_2O_3}$	C, H, N	20	L u	- †
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	H2NCH3CH2N	H	A3	56	L	245-246	$C_{13}H_{17}N_{3}O_{2}\cdot 2HCI\cdot 2H_{2}O$	C, H, N, Cl	10	Iv	+ + -
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	(CH ₃) ₂ NCH ₂ C	HIN211	$A3^{e}$	21	М	262-264	$C_{15}H_{21}N_2O_9\cdot 2HCI\cdot H_2O$	C, H, N, Cl	10	Ĭv	- }
	C2H500CCH	^{2}NH	A3	27	Μ	228 - 230	C ₁₅ H ₁₈ N ₂ O ₄ ·HCl·0.5H ₂ O	C. II. N. CI	10	Iv	- +
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	HOOCCH2C	H_2NH	$A3^{e}$	57	Т	272 - 274	$C_{14}H_{16}N_2O_4 \cdot 1.5H_2O$	C. H. N	100	L L	- +
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	C ₂ H ₅ OOCH ₂	CH2NH	_ م	52	8	237 - 239	C ₁₆ H ₂₀ N ₂ O ₄ · HCl	C, H, N, CI	ю	Ιv	+ + + -
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	C6H5CH2NH		A3	69	L	250 - 251	C ₁₈ H ₁₈ N ₂ O ₂ ·HCl	C, H, N	10	Iv	. +
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	3,4-(CH ₃ U) ₂ C	36H3CH2NH	g	75	K	199 - 201	$\mathrm{C}_{20}\mathrm{H}_{22}\mathrm{N}_{2}\mathrm{O}_{4}\cdot\mathrm{HCl}$	C, H, N	20	Iv	+++++++++++++++++++++++++++++++++++++++
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	CHINH CHINH		$\Lambda 3^h$	17	I	252-254	$\mathrm{C}_{16}\mathrm{H}_{16}\mathrm{N}_{2}\mathrm{O}_{3}\cdot\mathrm{HCl}$	C, H, N, CI	10	Iv	+++++++++++++++++++++++++++++++++++++++
H $i = i = 41$ $Q = 248-551$ $C_{17}H_{ab}N_{2}O_{2} + HC1 - 0.5H_{2}O = C_{1}H_{3}N_{1} + 1$ $Q = 10$ $V_{1} + 1$ $A_{3}^{i} = 47$ $X = 255-256$ dec $C_{17}H_{ab}N_{2}O_{2} + HC1 - 0.5H_{2}O = C_{1}H_{3}N = 10$ $V_{1} + 1$ $A_{3} = 54$ $K = 243-245$ $C_{18}H_{ab}N_{2}O_{2} + HC1 - 0.5H_{2}O = C_{1}H_{3}N = 10$ $V_{1} + 1$ $A_{3} = 74$ $Q = 243-245$ $C_{18}H_{3}N_{2}O_{3} + HC1 + H_{2}O = C_{1}H_{3}N = 0$ $V_{1} + N, CI = 10$ $V_{1} + 1$ $A_{3} = 74$ $Q = 212-216$ $C_{18}H_{3}N_{2}O_{3} + HC1 + H_{2}O = C_{1}H_{3}N = 0$ $V_{1} + N, CI = 10$ $V_{1} + 1$ $A_{3} = 74$ $Q = 212-216$ $C_{18}H_{3}N_{2}O_{3} + HC1 + H_{2}O = C_{1}H_{3}N = 0$ $V_{1} + N, CI = 10$ $V_{1} + 1$ $C_{1} = 47$ $U = 251-257$ $C_{18}H_{3}N_{2}O_{3} + HC1 + H_{2}O = C_{1}H_{3}N = 0$ $V_{1} + 0$ $V_{1} = 10$ $V_{1} + 1$ $C_{2} = 45$ $R = 223-235$ dec $C_{16}H_{4}N_{2}O_{3} + HC1 + H_{2}O = C_{1}H_{3}N = 0$ $C_{1}H_{3}N_{3}O_{3} = 0$ $V_{1}H_{3}N_{3}O_{3} = 0$ $V_{1}H_{3}N_{3}O_{$	C6H1NH		A3	26	ßr	941949	$C_{n}H_{n}N_{n}O_{n}HO_{n}O_{n}$ and O	M II U	01		
I $A3^{j}$ 47 X $255-256 dec$ $C_{irH_{0}N_{2}O_{1}} + ICI \cdot H_{2}O$ $C_{j}H_{j}N_{j}$ $10^{-1}V_{j}$ $+$ $A5$ 54 K $243-245$ $C_{ia}H_{ia}N_{2}O_{2} + ICI \cdot 0.5H_{2}O$ $C_{j}H_{j}N_{j}$ $10^{-1}V_{j}$ $+$ $A3$ 28 L $251-257$ $C_{ia}H_{ia}N_{2}O_{2} + ICI \cdot H_{2}O$ $C_{j}H_{j}N_{j}CI$ $10^{-1}V_{j}$ $+$ $A3$ 74 Q $212-216$ $C_{ia}H_{ia}N_{2}O_{3} + ICI \cdot H_{2}O$ $C_{j}H_{j}N_{j}CI$ $10^{-1}V_{j}$ $+$ $A3$ 74 Q $212-216$ $C_{ia}H_{ia}N_{2}O_{3} + ICI \cdot H_{2}O$ $C_{j}H_{j}N_{j}CI$ $10^{-1}V_{j}$ $+$ $C1$ 47 U $254-259$ $C_{ia}H_{ia}N_{2}O_{3} + ICI \cdot H_{2}O$ $C_{j}H_{j}N_{j}CI$ $10^{-1}V_{j}$ $+$ $C2$ 45 R $223-235$ dec $C_{ia}H_{ia}N_{2}O_{3} + ICI \cdot H_{2}O$ $C_{j}H_{j}N_{j}CI$ $10^{-1}V_{j}$ $+$ $C2$ 45 R $223-235$ dec $C_{ia}H_{ia}N_{2}O_{3} - ICI \cdot H_{2}O$ $C_{j}H_{j}N_{j}CI$ $10^{-1}V_{j}$ $+$ $H_{c}CH=N$ $C3$ 59 Q $I_{5}-167$ $C_{ia}H_{ia}N_{2}O_{3} - C_{j}H_{j}N_{j}CI$ $10^{-1}V_{j}$ $10^{-1}V_{j}$ $+$ $H_{c}CH=N$ $C3$ 54 V $218-224$ $C_{ia}H_{ia}N_{3}O_{3} - C_{j}H_{j}N_{j}O_{3} - C_{j}H_{j}N_{j}O_{j} - C_{j}H_{j}N_{j}$	C ₆ H ₅ NH		į	41	ð	248-251	CLAHEN, O. HCI	C, H, N	10	1v 1.	+
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	p-HOC ₆ H,NI	I	$A3^{j}$	47	X	$255-256 \deg$	CirHisNaO2 HCI H.O	C. H. N	10	IV I	⊦ -
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	$(CH_3)_2N$		$\Lambda 5$	54	К	243 - 245	$C_{13}H_{16}N_2O_2 \cdot HCl \cdot 0.5H_2O$	C, H, N	5 5	Iv	- +-
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	CH _a N		$\Lambda 3$	28	L	251 - 257	$\mathrm{C}_{16}\mathrm{H}_{21}\mathrm{N}_{3}\mathrm{O}_{2}\cdot 2\mathrm{HCl}\cdot\mathrm{H}_{2}\mathrm{O}$	C, H, N, Cl	10	Iv	+
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	Morpholinyl		$\mathbf{A3}$	74	ð	212 - 216	$C_{15}H_{18}N_2O_3\cdot HCI\cdot H_2O$	C, H, N	10	Ĭv	· +
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	CH3CONH		CI	47	U	254 - 259	C ₁₃ H ₁₄ N ₂ O ₃ ·HCl·H ₂ O	C. H. N. CI	10	L,	i
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	C ₂ H ₅ 00CCH	2NHCONH	C2	45	ы	223–235 dec	$C_{16}H_{19}N_{3}O_{5}$	C, H, N	100	, u	Pressor
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	C2H5NHCON	ICONHC2H5	C2	37	2	185-187	$C_{17}H_{22}N_4O_4$	С, Н, N	100	Ip	+
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	3,4-(CH ₃ O) ₂ C	6H3CH=N	C3	59	ზ	165-167	$\mathrm{C}_{20}\mathrm{H}_{\mathrm{E0}}\mathrm{N}_{2}\mathrm{O}_{4}$	C, II, N	10	Iv	+ + +
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	$p-(CH_3)_2NC_6$	H,CH=N	3 3 3	49	M	199-202	C18H21N3O2	C, H, N	15	Ĭv	- - - +
15 40 M 201-204 $C_{17}H_{15}NO_{3} \cdot HCI \cdot 1.5H_{2}O$ $C_{1}H, N, CI$ 20 I_{V} +	p-U2NC6H4C	H=N	3 a	54	> ;	218 - 224	$C_{18}H_{15}N_3O_4$	C, H, N	25	$_{\rm Ip}$	+-
	U ₆ H ₅ U		я	40	M	201 - 204	$C_{17}H_{15}NO_8 \cdot HCl \cdot 1.5H_2O$	C, H, N, Cl	20	Iv	· +

+ + + + +			Rating ^e +	$\begin{array}{c} + \\ + \\ + \\ + \\ Pressor \end{array}$		Rating ^c	+++			Rating ^e	+ + + + +	+ + +	+++++++++++++++++++++++++++++++++++++++
I v Iv V I v I			Route Ip	d d d ∧		Route	Iv Ip Iv			Route	Iv Iv	Iv Iv	Iv
100 50 30 30			Dose, mg/kg 20	2 10 5 20 2 10 2 7		Dose, mg/kg	10 100 10			Dose, mg/kg	25 30	10 10	10
L N S N C L N S N L L L N S S N S S S S S S S S S S S S S S			Analyses C, H, Cl	C, H, N, Cl C, H, N C, H, N, Cl C, H, Cl		Analyses	С, Н, И С, Н, N С, Н, С	5		Analyses	C, H," N, Cl	C, H, N, Cl C, H, N, Cl	C, H, N, Cl
4 ₂ 0 ₅ HCl C, H 4 ₀ 0 ₃ 2HCl 2H ₂ O C, H 40 ₅ S HCl 2H ₂ O C, H 40 ₅ S HCl H ₈ O C, H 40 ₅ S HCl H ₈ O C, H			Formula C14H14CINO4	$C_{16}H_{19}Cl_2NO_2$ $C_{13}H_{14}N_2O_4$ $C_{15}H_{18}N_2O_4 \cdot HCl \cdot 0 \cdot 5H_2O$ $C_{13}H_{14}ClNO_2 \cdot HCl \cdot H_2O$		Formula	C ₁₂ H ₁₅ IN ₂ O ² C ₁₄ H ₁₅ IN ₂ O ² C ₁₅ H ₁₅ IN ₂ O ₄ ·H ₂ O C ₁₂ H ₁₅ CIN ₂ O ₂			Formula	$C_9H_8N_2 \cdot HCI$ $C_{10}H_{10}N_2O \cdot HCI$	С ₁₃ H ₁₆ N ₂ O ₂ HCl С ₁₃ H ₁₄ N ₂ O ₂ HCl	C ₁ ,H ₁₄ N*O ₃ ·HCl
C ₁₇ H ₁₆ N C ₁₇ H ₁₆ N C ₁₂ H ₁₁ N C ₁₂ H ₁₃ N C ₁₁ H ₁₃ N	\mathbb{R}_4	$\mathbf{N} - \mathbf{R}_{2}$	M _P , °C ^b 159–160	$\begin{array}{c} 158{-}159\\ 248{-}250\\ 215{-}216\\ 180{-}200\end{array}$	$+ R_2 I^-$	$M_{\rm p}, {}^{\circ}{\rm C}^{b}$	272–276 dec 240–244 171–174 280–283	\mathbf{H}_{2}	R	$M_{\mathbf{p}}, \ ^{\circ}\mathrm{C}^{b}$	303–311 dec 250	273-275 292-293	266-270
215-220 215-218 232-235 227-229 283-288		CH ₃ O-H3	Recrystn ^a solvent M	он я	H ^O H ^O	Recrystn ^a Solvent	L W 66% EtOH 75% MeOH	Z –		A8 Recrystn ^a solvent	сIJ	Γ	S
${\rm P}_{\rm P}$			Yield, $\%$ 46	48 54 15		Yield, %	51 42 43			Yield, %	29 58	54 63	41
46 43 68 21 21			Method l	l m Ale,n l		Method	~~~~			Method	0 A1 ^p	A1 A3 ^d	1
ч ч К			$\mathbf{G}^{\mathbf{R}}$	CI CH _s NH CH _s NH CI		ž	H2N CH3NH CH3NH H2N			${ m R_8}$	нн	н	
			R3 COOC ₂ H5	СІСН ₂ СН ₂ СООН СООС ₂ Н5 Н		R3	Н Н СООН Н			\mathbf{R}_{7}	H $CH_{3}O$	C ₃ H ₅ O CH ₃ O	
$p-0_2\mathrm{NC_6H_4O}$ $p-\mathrm{H_2NC_6H_4O}$ HS CH_3S NH_2NH			R _i R ₂ H	(OC ₂ H ₅) ₂ CH ₃ H (OC ₂ H ₅) ₂ H		\mathbf{R}_{2}	нннн	ion is Cl-)		R	Н	C ₂ H ₅ O , CH ₃ O	
37 33 40 41			t c	6,7- 6,7-(R	C C C H	(Ani		R.	нн	н СЩ	CH ³ C
			Compd-) s No. 42	44 44 45 46		Compd ss No.	I 50 52 51 53			Compd ss No.	11 54 55	56 57	58
			VI VI			Cla	ΙΛ			Clar	ΙΛ		IX



in blood pressure or a duration of action of $\langle 45 \text{ min} \rangle$; moderate activity, ++ (>40% decrease in blood pressure with a duration of action >45 min but $\langle 270 \text{ min} \rangle$; maximum activity, +++ (>40% decrease in plood pressure with a duration of action >270 min).

Results

This paper has dealt primarily with 6,7-dialkoxy-4-(substituted amino)quinolines (I, II, VIII). The hypotensive activity of these compounds as determined in anesthetized dogs and evaluated by the previously described rating system is delineated in Table I. Although a wide variation of these 4-(substituted amino)quinolines exhibited various degrees of activity, the essential fact is that the basic 4-(substituted amino)quinoline molecule generally elicits hypotensive activity in anesthetized dogs. Also lesser degrees of hypotensive activity were observed (a) where the 4 position of the quinoline was substituted by amido (III), methyleneamino (IV), phenoxy, thio, and hydrazino (V), and (b) where the 1 position of the 4-(substituted amino)quinoline was quaternized (VII).

Experimental Section

4-Amino-6,7-dimethoxyquinoline \cdot HCl \cdot H₂O (1) (Method A1³).—A 5-l. flask fitted with stirrer, thermometer, condenser, and gas-inlet tube was charged with molten PhOH (2000 ml) and 47³ (600 g, 2.68 moles). The mixt was heated to 100° with a Glas-Col mantle, then satd with anhyd NH₃ in 10 min. The NH₃ addn was interrupted, while the mixt was heated to reflux (166–176°), and then resumed at a moderate rate at refux for 3.5 hr. The reaction soln was air-cooled to 50° and poured rapidly into stirred Et₂O (12 l). The product was collected (on paper over cloth) by filtration and rinsed with *i*-PrOH and Et₂O: mp 261°; yield, 613 g (88%). Recrystn of 306 g from boiling 85% aq MeOH (3000 ml) with slow addn of H₂O (200 ml) with charcoal gave white cryst 1.

Method A2.—NH₄OAc (12 g, 0.16 mole) was heated in an open flask at 110–173° in 18 min and cooled in air for 7 min. To the preheated NH₄OAc was added the free base of 36 (2.0 g, 0.0071 mole), and the mixt was heated at 178–180° for 1.1 hr. The cooled reaction mixt was dissolved in *i*-PrOH, then treated with 10% HCl (5 ml) in the cold. The cryst product was collected by filtration and washed with *i*-PrOH and ether.

6,7-Dimethoxy-4-propylaminoquinoline \cdot HCl (5) (Method A3).—To a warm soln of 47 (34 g, 0.15 mole) in PhOH (110 ml) was added PrNH₂ (10 g, 0.17 mole) with mechanical stirring. The reaction soln was refluxed for 2 hr, and then added to anhyd Et₂O (1250 ml); the resultant sticky material gradually solidified upon trituration. The solid was collected by filtration and immediately recrystd from a mixt of 95% EtOH (250 ml) and MeOH (140 ml) to give cryst 5.

4-Hydroxyamino-6,7-dimethoxyquinoline \cdot HCl (9) (Method A4).—To a soln of 47 (60 g, 0.27 mole) in PhOH (640 ml) was added 54% NaH (14.6 g, 0.33 mole) in mineral oil at 55-70°, with mechanical stirring; the reaction was exothermic. After 10 min H₂NOH \cdot HCl (15.0 g, 0.22 mole) was added to the mixt, which was heated on the steam bath for 0.5 hr. A second portion of H₂NOH \cdot HCl (15.0 g) was added, and the reaction mixt was heated at 94-103° for 2.5 hr. The cooled mixt was added to *i*-PrOH and cooled in an ice bath, and the resultant brown solid was collected by filtration and washed with *i*-PrOH (90 ml) and Et₂O; yield, 58 g.

Recrystn of the product (25 g) from a mixt of 25% MeOH (4 1.) and 10% HCl (80 ml) with charcoal gave **9**.

6,7-Dimethoxy-4-dimethylaminoquinoline \cdot HCl \cdot 0.5H₂O (27) (Method A5).—A soln of 47 (70 g, 0.31 mole) in DMF (450 ml) was satd with dry Me₂NH at 40–85° over 35 min with mechanical stirring. The satd soln was heated to 140° in 40 min, then the addn of Me₂NH was contd at 145–150° over 6.5 hr. The cooled soln was added to anhyd Et₂O (2300 ml) and filtered. The filtrate was evapd to dryness under reduced pressure. In order

to remove residual DMF the crude residue was treated with *i*-PrOH (150 ml) and evapd to dryness. Treatment of a soln of the residue in *i*-PrOH (500 ml) and 10% HCl (110 ml) gave the hydrochloride **27**.

6,7-Dimethoxy-4-phenoxyquinoline \cdot HCl \cdot 1.5H₂O (36) (Method B).—A soln of 47 (70 g, 0.31 mole) in PhOH (240 ml) was refluxed for 2.3 hr. The cooled soln was treated with H₂O (500 ml) and C₆H₆ (650 ml), and the mixt was neutralized with 10% NaOH to pH 8–9. The aq layer was further extd with C₆H₆ (800 ml). The combined exts were dried over a mixt of MgSO₄ and charcoal, filtered, concd to a vol of 115 ml, and cooled in the refrigerator. The resultant light yellow, cryst free base of 36 was collected by filtration and washed with cold C₆H₆ and Et₂O: mp 115–120°; yield, 35 g (40%).

A soln of the free base of **36** (46 g, 0.16 mole) in C_6H_6 (700 ml) was treated with dry HCl with cooling. The hydrochloride was collected by filtration and recrystn from *i*-PrOH (290 ml) to give **36**.

4-Mercapto-6,7-dimethoxyquinoline (39).—Dry H_2S was bubbled through a soln of Na₂S (54 g, 0.69 mole) in DMF (1300 ml) at 25–30° for 3.5 hr. To the soln was added the free base of 47 (136 g, 0.6 mole) with mechanical stirring. The reaction soln was refluxed at 132–139° for 1.2 hr. The cooled mixt was added the J2 (61.). The resultant yellow, cryst solid was collected by filtration, washed with H₂O (375 ml), and dried in air. Recrystn from 12% EtOH–DMF gave 39.

6,7-Dimethoxy-4-methylthioquinoline $HCl H_2O$ (40).—To a soln of 39 (60 g, 0.27 mole) in 5% NaOH (258 ml) was added Me₂SO₄ at 3-10° in 20 min with rapid stirring. The ice-cooled mixt was stirred for 1 hr, then warmed in the air for 2.5 hr. The resultant yellow, cryst solid was collected by filtration, washed with H₂O (200 ml), and dried in air at 65°. Recrystn from *i*-PrOH (700 ml) with charcoal gave the free base of 40: mp 174–175°; yield, 29.8 g. The free base in 95% EtOH (1500 ml) was treated with dry HCl in the cold. The hydrochloride 40 was collected by filtration and washed with EtOH and Et₂O.

 $\begin{array}{l} \label{eq:4-Hydrazino-6,7-dimethoxyquinoline \cdot 2HCl (41). A suspension of 47 (5.0 g, 0.022 mole) and N_2H_4 \cdot H_2O (20 ml) was refluxed for 75 min. The reaction soln was cooled, and the resultant solid was collected by filtration. Recrystn from 5% HCl gave 41. \end{array}$

4-Acetamido-6,7-dimethoxyquinoline \cdot HCl \cdot H₂O (30) (Method C1).—A soln of the free base of 1 (30 g, 0.15 mole) in (MeCO)₂O (450 ml) was refluxed for 6 hr. The reaction soln was concd under reduced pressure, the residue was heated on the steam with *i*-PrOH (60 ml) and cooled, and the resultant solid was collected by filtration and washed with *i*-PrOH (3 × 10 ml). A mixt of the solid product was treated with dry HCl in *i*-PrOH (200 ml). Recrystn of the resultant hydrochloride (26 g) from a mixture of 95% ethanol (1400 ml) and coned HCl with charcoal gave 30.

Ethyl 5-(6,7-Dimethoxy-4-quinolyl)hydantoate (31) (Method C2).—A mixt of hydrated free base of 1 (90 g, 0.41 mole on anhyd basis) and C₆H₆ (900 ml) was refluxed with Dean–Stark trap and mechanical stirring for 6 hr, until the H₂O (7.0 ml) was removed. To the mixt was added dropwise, a soln of ethoxycarbonylmethyl isocyanate (63 g, 0.49 mole) in dry C₆H₆ (225 ml) in 2 hr with heating on a steam bath. The reaction mixt was further heated for 1.5 hr, then cooled at 8–15° for 1 hr. The resultant brown solid was collected by filtration and washed with C₆H₆ (100 ml) and Et₂O: yield, 100 g. Recrystn of the product (67 g) from MeCN (3800 ml) with charcoal gave **31**.

Ethyl 4-Chloro-6,7-dimethoxy-3-quinolinecarboxylate (42).—A mixt of 48^3 (18 g, 0.065 mole) and POCl₃ (150 ml) was refluxed for 30 hr. The reaction mixt was worked up in the usual manner.

4-(3,4-Dimethoxybenzylideneamino)-6,7-dimethoxyquinoline (33) (Method C3).⁶—A mixt of hydrated free base of 1 (80 g, 0.34 mole on anhyd basis) and PhMe (1500 ml) was refluxed (a Dean–Stark trap) until the H₂O (10 ml) was removed. Then veratraldehyde (66 g, 0.40 mole) and piperidine (35 ml) were added to the mixt, which was refluxed for 18.5 hr. The hot reaction soln was decanted from an insoluble solid (4 g) and cooled in an ice bath, and the product was collected by filtration and washed with petroleum ether: mp 155–157°; yield, 95 g. Recrystn from EtOH (1700 ml) with charcoal gave 33.

4-Chloro-6,7-diethoxyquinoline \cdot HCl \cdot H₂O (46).—To Dowtherm A (780 ml) at 250° was added portionwise 6,7-diethoxy-4hydroxyquinoline-3-carboxylic acid⁹ (78 g, 0.25 mole), and the mixt was refluxed 1 hr. The cooled mixt was treated with hexane (1000 ml) then decanted from the amorphous residue, which was triturated with petr ether and C₆H₆, resp. Recrystn from H₂O (1200 ml) gave 6,7-diethoxy-4-hydroxyquinoline: mp 95°; yield, 39.5 g. The compd was chlorinated in the usual manner with POCl₃ to give 46.

4-Amino-6,7-dimethoxy-1-methylquinolinium Iodide (50).—To a soln of 1 (50 g, 0.25 mole) in EtOH (1300 ml) was added MeI (60 ml, 0.96 mole) at 21° over 4 min with mechanical stirring. The reaction mixt was refluxed for 1 hr and then cooled in an ice bath. The resultant white, cryst solid was collected by filtration and washed with i-PrOH-ether: yield, 78 g. Recrystn of the product (30 g) from MeOH (35 ml) with charcoal gave 50.

1-Ethyl-6,7-dimethoxy-4-methylaminoquinolinium Iodide (51). —A mixt of 47 (112 g, 0.50 mole), acetone (1500 ml), NaI (1.0 g, 0.0067 mole), and EtI (80 ml, 1.0 mole) was refluxed for 5 days. The resultant cryst iodide (101 g) was collected by filtration of the cooled reaction mixt. Dry MeNH₂ was passed through a mixt of the iodide (75, g 0.20 mole) and MeNO₂ (1350 ml) at 24–41° for 9 hr with mechanical stirring; the reaction was exothermic. The reaction mixt was stirred for 15 hr at 24–26°, and then cooled in an ice bath. The yellow cryst product was collected by filtration and recrystn from MeNO₂ (600 ml) with charcoal.

3-Carboxy-1-ethyl-6,7-dimethoxy-4-methylaminoquinolinium Iodide H_2O (52).—To a soln of NaOH (17 g, 0.42 mole) in 66%EtOH (900 ml) was added 44 (50 g, 0.19 mole) at 30° with mechanical stirring. Next, EtI (100 ml, 1.25 mole) was added at 25-27° in 2 min. The reaction mixt was refluxed 17 hr, then cooled in an ice bath. The resultant tan solid was collected by filtration and recrystd from 66% EtOH (600 ml) with charcoal.

4-Amino-6,7-dimethoxy-1-methylquinolinium Chloride (53).— A soln of 50 (3.0 g) in a mixt of MeOH (200 ml) and H₂O (150 ml) was passed slowly through a 44 \times 150 mm column, of Dowex 1-X8 (ionic Cl⁻ form) resin. A center cut of the effluent (150 ml) was evapd to dryness under reduced pressure; the resultant white crystals (mp 277-279° dec) were recrystd from 75% MeOH (H₂O), mp 280-283° dec, mmp 252-255° dec with 2 (mp 254-257° dec). The ir absorption of 53 and 2 differed greatly. Nmr spectrum (δ) showed: (DMSO) 3.98, singlet (CH₃); 4.04, 4.08, singlets (2CH₃O); 6.73, 6.85 and 8.30, 8.41, pair doublets (2 heterom); 7.31, 8.06, singlets (2 arom); 8.8, broad absorption (NH₂, exchanged with D₂O).

6,7-Dimethoxy-4-methylaminoquinoline \cdot HCl (2) was prepd by method A1. Nmr spectrum (δ) showed: (DMSO) 3.05, 3.13, doublet (CH₃, a singlet at 3.1 with D₂O exchange); 3.96, 3.98, singlets (2CH₂O); 6.55, 6.67, and 8.27, 8.38, pair doublets (2 heterom); 7.50, 8.05, singlets (2 arom); 9.4, broad absorption (NH, exchanged with D₂O).

2,3-Dihydro-8,9-dimethoxybenzo[h]-1,6-naphthyridin-4(1H)one ·HCl (58).—To warm PPA (800 g) was added 19 (80 g, 0.29 mole) with mechanical stirring; the resultant soln was heated at 85-90° for 2 hr. The cooled soln was added to H₂O (3500 ml) at 20-30° with stirring. The crude product was collected by filtration and washed with *i*-PrOH (125 ml) and Et₂O. Recrystn from 10% HCl (7 l.) with charcoal gave 58. Nmr spectrum (δ) showed: (DMSO) 3.70 singlet (2CH₃O); 6.96, singlet (1 heterom); 7.54, 8.30 singlets (2 arom); the CH₂CH₂ protons were unresolved.

4-Chloro-3-(2-chloroethyl)-6,7-diethoxyquinaldine (43).—A mixt of 3,4-diethoxynitrobenzene⁹ (30 g, 0.14 mole), 5% Pd/C (3 g), and EtOH (200 ml) was hydrogenated in a Parr apparatus. The catalyst was removed by filtration. The process was repeated, and the combined 3,4-diethoxyaniline soln were treated with 2-acetylbutyrolactone⁸ (36 g, 0.28 mole) and refluxed for 2 hr. The reaction soln was concd to 0.5 vol and cooled, and the resultant anil **60** was collected by filtration: mp 113–116°; yield, 60 g (74%).

To a soln of POCl₃ (112 ml) in PhMe (100 ml) at 40–50° was added portionwise **60** (94 g, 0.32 mole) in PhMe (700 ml). The reaction mixt was heated at 85–90° for 1 hr then refluxed for 3 hr. Excess solvents were removed under reduced pressure, addnl PhMe (250 ml) was added and again removed, and the residue was dissolved in PhMe and treated with ice H₂O (2000 ml). The mixt was made fully basic with NH₄OH and filtered. The solvent

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was removed from the org layer under reduced pressure, and the residue was recrystd from i-PrOH to give **43**.

Acknowledgments.—The authors are grateful to Dr. Harry Snyder for the preparation of ethyl 4-chloro-6,7-

dimethoxy-3-quinolinecarboxylate; to Mr. Grant Gustin and Mr. Marvin Tefft for the elemental analyses; to Mrs. Patricia Curtis for the nmr analyses; and to Mr. Frank Wessels and Mr. Paul Bowes for pharmacologic data.

Quaternary Pilocarpine Derivatives Acting as Acetylcholine Antagonists

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Received November 12, 1970

Several quaternary *d*-pilocarpine derivatives have been prepared in order to investigate the influence of structural changes on the biological activity of this alkaloid. The effect of the substituents in the reagent, as well as of the temp and the solvent (its dielectric constant), on the rate of the quaternization has been studied, and the products have been analyzed by various spectroscopic means. The anticholinergic activities of the compounds are reported, and a relation has been sought in connection with the structural changes.

Pilocarpine (I) is the main alkaloid obtained from the leaves of the South American shrubs *Pilocarpus jabor*andi and *Pilocarpus microphyllus* Stapf. The structures of pilocarpine and its isomer, isopilocarpine, were determined by Jowett¹ and both were synthesized by several routes.² The absolute configuration of pilocarpine has been established as being $7R,8S.^3$ *d*-Pilocarpine, one of the oldest parasympathomimetic drugs,⁴ may act as an anticholinergic in certain systems.⁵

The purpose of this study was: (a) to develop methods for the addition of various groups to the alkaloid by quaternization at N-3 and determine the various conditions influencing the reaction and the stability of the products; (b) study some aspects of the relative reactivity of the alkaloid with various halo organic reagents; (c) test the pharmacological activity of the new compds as a function of structural change. It has been reported that quaternization of atropine and scopolamine with different substituted phenacyl bromides induces changes in their pharmacological activities.⁶

Results and Discussion

The free base of d-pilocarpine (I) was treated with different halo organic compds producing a series of quaternary deriv with the general structure II (Table I).

The effect of the substituents in the halo organic reagents, the temp, and the solvent influence the optimal time of the reaction. The data collected in Table I show a marked decrease in the rate of quaternization in Me₂CO medium passing from Et to *n*-Pr (1-3), but in contrast to previous observations,^{7,8} with *n*-BuBr prac-

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tically no reaction took place. In a solvent with higher polarity (2-methoxyethanol) only 6 days were required for completion of the reaction. It was observed that n-PrI was about twice as reactive as the bromide, whereas with i-PrBr no quaternization would take place. It is therefore difficult to distinguish between electronic and steric effects in these reactions.

In the case of benzyl halides the reactivity is relatively greater, and is influenced by the character and the position of the substituent. Electron-releasing groups in the para position (7, 10, 13) enhance the displacement of the halogen, the reaction becoming more sluggish with a Me group. With ortho substituents of the same character (9, 11) steric hindrance makes the reaction slower by far. An electron-attracting group, such as NO_2 , at the para position induces a decrease of the rate of the reaction, bromide 15 being more reactive than chloride 16. In contrast, when NO₂ is at the meta position (14) the reaction is faster. When Ph is further away from the side-chain halogen atom (5), no conjugation between the ring and the side-chain halogen is possible, and the reaction becomes sluggish; it could be accelerated, however, by using a solvent with high dielectric constant (8) such as 2-methoxyethanol ($\varepsilon \sim 40$). The comparatively high reactivity with the phenacyl bromides (19-23) may be explained by the activating effect of the carbonyl group.⁹

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