[CONTRIBUTION FROM THE CHEMICAL AND BIOLOGICAL RESEARCH SECTION, LEDERLE LABORATORIES DIVISION, AMERICAN CYANAMID COMPANY]

Derivatives of 1-Piperazinecarboxylic Acid as Sedatives¹

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A series of esters of 4-acyl- and 4-aroyl-1-piperazinecarboxylic acid represented by formula II has been synthesized and characterized. Several have been found to possess sedative properties.

In studying the pharmacological properties of various derivatives of piperazine it has been noted² that some possess sedative properties when tested in animals; Nonas³ has reported that N,N-diethyl-1-piperazinecarboxamide⁴ (I) has possible clinical usefulness as a daytime sedative. Since various amides and urethans exhibit sedative and hypnotic properties,⁵ it was decided to synthesize derivatives of piperazine represented by formula II in which N⁴ is substituted with an acyl or aroyl group and N¹ is substituted with a carbalkoxy or carbaralkoxy group.



Among the 4-acyl and aroyl derivatives of ethyl 1-piperazinecarboxylate listed in Table I, a number were found to be active as sedatives when tested with the asymptomatic dose in rats in activity cages. In the normal acyl series there is a narrow range of activity with the peak (4+) reached at C_4-C_5 in ethyl 4-n-butyryl-1-piperazinecarboxylate (III) and in ethyl 4-n-valeryl-1-piperazinecarboxylate (XIII). In the branched-chain acyl series there is a wider range of activity with the peak



(4+) at isovaleryl⁶ (XIV), and 3+ activity for the 2-methylbutyryl (XV), 3-methylvaleryl (XVIII) and 2-ethylcaproyl (XXII) derivatives. The isocaproyl derivative (XVII) is much more toxic than the previously mentioned compounds and at the asymptomatic dose level (30 mg./kg. orally) the compound is inactive. The aroyl derivat (XXV, XXVI, XXVII, XXVII) are inactive. The aroyl derivatives

A comparison of the methyl, ethyl, n-butyl and benzyl esters (IX, III, X, XI) of 4-n-butyryl-1-

(1) Presented before the Division of Medicinal Chemistry, 126th National Meeting of the American Chemical Society, New York, N. Y., September 12-17, 1954.

- (2) R. W. Cunningham, et al., private communication.
- (3) G. Nonas, N. Y. State J. Med., 50, 1257 (1950).

(4) S. Kushner, L. M. Brancone, R. I. Hewitt, W. L. McEwen and Y. SubbaRow; H. W. Stewart, R. J. Turner and J. J. Denton, J. Org. Chem., 13, 144 (1948).

(5) S. Fränkel, "Die Arzneimittel-Synthese," Julius Springer, Berlin, 1927, p. 522.

(6) During the course of this investigation R. B. Keller and R. A. LaForge, J. Am. Pharm. Assoc., Sci. Ed., 41, 301 (1952), reported the synthesis of XIV and found that it does not show promise as a sedative.

piperazinecarboxylic acid reveals that the peak sedative activity is reached in the ethyl ester III.

The compounds listed in Table I were synthesized by reaction of an alkyl or aralkyl ester of 1piperazinecarboxylic acid with an anhydride (procedures A, B and C) or with an acyl or aroyl halide under Schotten-Baumann conditions in the presence of sodium hydroxide (procedure D) or sodium bicarbonate (procedure E); or with a molar excess of the ester of 1-piperazinecarboxylic acid in an anhydrous solvent (procedures F, G, H and I).

Experimental⁸

The intermediate methyl, ethyl, n-butyl and benzyl The intermediate methyl, ethyl, *n*-butyl and benzyl esters of 1-piperazinecarboxylic acid were prepared by the methods of Moore, et al.,⁷ Stewart, et al.,⁹ and Goldman, et al.¹⁰ Ethyl trans-2,5-dimethyl-1-piperazinecarboxylate was prepared as described previously.¹¹ 2-Methylbutyryl chloride was obtained in 92% yield as previously described.¹² 2-Ethyl-*n*-caproyl Chloride.—*n*-Butylethylmalonic acid

2-Ethyl-n-caproyl Chloride.—n-Butylethylmatonic acid was decarboxylated to 2-ethyl-n-caproic acid in 96% yield according to the method of Raper.¹³ Reaction with thionyl chloride according to Levene and Kuna¹⁴ for the (-)-isomer gave 2-ethyl-n-caproyl chloride, b.p. 73° (17 mm.), in 87% yield. Tiffeneau¹⁵ reports b.p. 85–90° (20 mm.); Levene and Kuna¹⁴ report b.p. 62–64° (10 mm.) for the -)-isomer.

3-Methylvaleryl Chloride.—sec-Butylmalonic acid,16 obtained in 76% yield by saponification of the ethyl ester,16 was decarboxylated¹⁷ to give a 93% yield of 3-methylvaleric acid,¹⁸ which reacted with thionyl chloride according to Colonge¹⁹ to give a 65% yield of 3-methylvaleryl chloride, b.p. 73-75° (80 mm.); Hommelen¹⁸ gives b.p. 142.5-143.0° (749 mm.).

n-Caprylyl Chloride.—Obtained in 96% yield by reaction of *n*-caprylic acid with phosphorus pentachloride. Averill, *et al.*,²⁰ had synthesized this compound by the reaction of n-caprylic acid with oxalyl chloride (method of Adams and Ulich²¹).

Procedure A. Ethyl 4-Acetyl-1-piperazinecarboxylate (IV).—A solution of 31.6 g. (0.2 mole) of ethyl 1-piperazine-carboxylate' in 100 ml. of glacial acetic acid was treated, with cooling, with 20.4 g. (0.2 mole) of 97% acetic an-hydride. The resulting solution was heated for 1 hour on a steam-bath and then evaporated in vacuo to remove the acetic acid. Distillation in vacuo of the residual sirup gave 33.7 g. (84%) of colorless liquid, b.p. 143-145° (0.3 mm.), n²⁵D 1.4865.

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⁽⁸⁾ All boiling points and melting points are uncorrected.

NCO2R'

TABLE I. RCON

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								1									Sec	lative
Cmpd.	R	R'	R.	Pro- cedure	د در B.	р. Мш.	M.p., °C.	1, ²⁵ D	Yield, %	Empirical formula	Calcd.	oon Found	Ana Hyd Calcd.	lyses, %~ rogen Found	Nitro Caled.	Found	E.E.B	ting ^a g./kg. rat)b
Λ	CH,	C_2H_b	Η	Y	143-145	0.3		1.4865	84	C9H16N2O3	54.0	53.7	8.1	8.4	14.0	14.1	+	500
Δ	CH ₃	C ₃ H ₆	CH3	щ	163	5			22	C ₁₁ H20N2O3	57.9	57.5	8.8	8.9	12.3	12.1	0	125 ip ^e
VI	CH,	CH ₃ C ₆ H ₅	Н	٩	198 - 203	0.3	42.5-43		87	C14H18N2O3	64.1	64.1	6.9	6.9	10.7	10.5	0	500
ΛII	C ₂ H ₆	C ₃ H ₆	Η	A'	145-149	¢.		1.4867	61	$C_{10}H_{18}N_2O_3$	56.0	56.0	8.5	8.5	13.1	12.7	0	500
VIII	C ₃ H,	CH ₂ C ₆ H ₆	Н	ව	184-188	с р .	50-50.5°		65	C16H20N2O2	65.2	65.4	7.3	7.1	10.1	10.1	0	85 ip
IX	$n-C_{3}H_{7}$	CH,	Н	υ	154 - 156	.7	35-37		77	C ₁₀ H ₁₈ N ₂ O ₃	56.0	56.1	8.5	8.6	13.1	12.9	+	200 ip
III	n-C ₃ H ₇	C ₃ H ₆	Η	'n	150 - 153	1.2		1.4849	84	C ₁₁ H ₂₀ N ₂ O ₂	57.9	57.5	8.8	8.8	12.3	12.3	$^{4+}$	400
				5	136-138.5	0.6			76			57.6		0.0		12.1		
x	n-C _a H ₇	$n-C_4H_9$	Η	υ	155 - 156	.4		1.4820	8	C ₁₃ H ₂₄ N ₂ O ₃	60.9	60.9	9.4	9.7	10.9	10.6	0	50 ip
XI	n-C ₃ H ₇	CH ₂ C ₆ H ₆	Н	٩	197 - 203	.3		1.5353	75	C16H22N2O3	66.2	66.0	7.6	7.8	9.6	9.5	0	200
IIX	iso-C ₃ H ₇	$C_{3}H_{6}$	Η	Έų	130 - 131	.5		1.4840	67	$C_{11}H_{20}N_2O_3$	57.9	57.7	8.8	8.8	12.3	12.0	$^{2+}$	500
XIII	n-C,H,	C ₃ H,	Η	J	157-158	9.		1.4840	87	C ₁₂ H ₂₂ N ₂ O ₃	59.5	59.2	9.2	9.2	11.6	11.7	4+	400
XIV	iso-C ₄ H ^s	C_2H_5	Η	Ċ	$121 - 122^{k}$.15		1.4839	71	C12H22N2O3	59.5	59.2	9.2	9.0	11.6	11.2	4+	500
XV	C ₂ H ₅ CH(CH ₁)	C ₃ H ₆	Н	Ċ	128-132	.08-0.1		1.4835	77	C12H22N2O3	59.5	59.3	9.2	8.8	11.6	11.4	3^{+}	500
IVX	n-C ₆ H ₁₁	C ₃ H,	Н	I	130-136	.05-0.08		1.4835^{l}	06	C ₁₃ H ₂₄ N ₂ O ₃	60.9	61.0	9.4	9.7	10.9	10.6	+	125
IIVX	iso-C ₆ H ₁₁	$C_{2}H_{5}$	Н	ы Ц	163 - 165	1.1		1.4823	92	C ₁₃ H ₂₄ N ₂ O ₃	60.9	60.3	9.4	9.8	10.9	10.9	0	30
IIIVX	C ₂ H ₅ CH(CH ₁)CH ₂	C ₃ H,	Н	Ċ	137 - 140	0.04		1.4840	87	C13H24N203	60.9	60.7	9.4	9.3	10.9	10.9	3+	500
XIX	(C ₃ H ₆) ₂ CH	$C_{2}H_{6}$	Η	г. Н	149 - 151	8.		1.4811	87	C ₁₃ H ₂₄ N ₂ O ₃	60.9	60.5	9.4	9.7	10.9	10.7	+	50
XX	n-CeH18	$C_{2}H_{6}$	Η	I	136 - 141	.05		1.4828	8 6	C14H26N2O3	62.2	61.8	9.7	9.8	10.4	10.3	0	125
IXX	n-C,H ₁₆	C ₃ H	Η	შ	168 - 172	.3-0.4	23 - 24.5	1.4810	82	C15H28N20	63.4	63.4	9.9	9.6	9.8	9.8	0	500
IIXX	C,H,CH(C,H,)	$C_{3}H_{6}$	H	ථ	138-143	.03-0.04		1.4791	68	C ₁₅ H ₂₈ N ₂ O ₂	63.4	62.9	9.9	9.5	9.8	9.6	33+	75
IIIXX	n-CuHm	C_2H_6	Η	μ			27 - 29		97	C19H36N2O3	67.0	66.8	10.7	10.8	8.2	8.0	0	1000
XXIV	n-CuHn	C ₃ H	Η	H			36.5-38"		96	$C_{21}H_{40}N_2O_3$	68.4	68.3	10.9	10.8	7.6	7.5	0	500
XXV	C,H,	$C_{3}H_{6}$	H	" Ц	186-187	0.9	80.5-82		67	C ₁₄ H ₁₈ N ₂ O ₂	64.1	64.0	6.9	6.8	10.7	10.4	0	500
				Ш			82.5-83.5		94			64.4		7.1		10.4		
XXVI	p-C ₆ H ₁ NO ₂	C ₃ H ₆	H	Э			$91.5 - 92.5^{\circ}$		45 (214Hr/N3O5	54.7	54.9	5.6	5.9	13.7	14.0	0	500
IIVXX	o-C,H,CI	C ₂ H	H	Ш	181-184	0.2	71-73, 77-78"		62	C ₁₄ H ₁₇ ClN ₂ O ₃	56.7	56.3	5.8	5.8	9.4	9.2		
														U U	1, 11.9	11.5	0	500
XXVIII	2,6-C,H,Cl	C ₂ H ₆	Η	т Ц			89.5-91		98	C14H16Cl2N2O2	50.8	51.0	4.9	5.0	8.5	8.2	0	500
														ΰ	1, 21.4	21.4		
Test for 2 hou 35%), pr	cd in rats in activity c rrs. • From ether. oduct extracted into	ages. ^b As: ^f Reaction benzene.	ympto run in React	matic benze ion n	dose admini ene on a ste nixture reflux	stered orally am-bath for ked 3 hours.	unless otherwis 3.5 hours. ^a R ⁱ Reaction tin	e indicated eaction tir ne 2 hours	ae 3 h K	is intraperito ours. ^h Run eller and LaF	neally. at -5 i orge ⁶ gi	^d Rea to -10 ve b.p.	ction r (whe 168-1	un in be in run at 70° (4 n	t +2 to $\frac{1}{100}$ mm.).	Tempe	tempe he yie	erature eld was re 24°.
Arom b From b	on mixture allowed to temperature for 2 da enzene-hexane. t Fr	ys. ^p Reac	tion m tion m	ixture lixture le con	n temperatu e allowed to npound melt	re. " Keacti stand at roo s at 71–73°, é	on mixture near m temperature and when the te	for 3 days	ng, coo	rom hexane. sed it solidifie:	r Moor s and re	LS IOF A Te, <i>et a</i> . melts a	11. 1.7 giv 1t 77–7.	keacuo e m.p. 8 8°. "	n mixtur 2° (from Obtained	e aulow I light I by trii	petro turat	o stand deum). ing the
undistille	d crude product with	1 5% sodiun	ı bicar	bonat	e and recrys	tallizing from	heptane and fi	rom aqueo	us etha	unol.						•		l

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Procedure B. Ethyl trans-4-Acetyl-2,5-dimethyl-1-piperazinecarboxylate (V).—A mixture of 18.6 g. (0.1 mole) of ethyl trans-2,5-dimethyl-1-piperazinecarboxylate¹¹ and 25 ml. of acetic anhydride was refluxed for 8 hours. The reml. of acetic anhydride was refluxed for 8 hours. The re-sulting solution was distilled *in vacuo*. After removal of acetic acid and acetic anhydride two fractions were ob-tained: (a) 7.2 g. of colorless liquid, b.p. 128–143° (15 mm.) (mainly ethyl *trans-2*,5-dimethyl-1-piperazinecar-boxylate); (b) 10.2 g. of nearly colorless viscous liquid, b.p. 120-163° (5 mm.) (mainly b.p. 163°). Fraction b was re-distilled and after removal of 3.0 g. of forerun, b.p. 113-163°(5 mm.) 5.1 g. (2292) of V was obtained as a viscous vellow (5 mm), 5.1 g. (22%) of V was obtained as a viscous yellow liquid, b.p. 163° (5 mm.). After standing for a long time the material solidified.

Procedure C. n-Butyl 4-n-Butyryl-1-piperazinecarboxylate (X).—To 37.2 g. (0.2 mole) of *n*-butyl 1-piperazinecarboxyl-ate,⁹ 31.6 g. (0.2 mole) of *n*-butyl 1-piperazinecarboxyl-with mixing and cooling. The resulting solution was heated on a steam-bath for 1 hour and then distilled through a Vigreux column. After removal of the *n*-butyric acid, 40.8 g. (80%) of X was obtained as a colorless liquid, b.p. 155-156° (0.4 nnm.), n^{25} D 1.4820.

Procedure D. Benzyl 4-n-Butyryl-1-piperazinecarboxyl-ate (XI).—To a stirred mixture of 44 g. (0.2 mole) of benzyl 1-piperazinecarboxylate¹⁰ and ice were added simultaneously and dropwise 32 g. (0.3 mole) of *n*-butyryl chloride and 100 ml. of 4 N sodium hydroxide; ice was added as required to maintain an excess. A colorless oil separated. The mix-ture was kept cold by means of an ice-bath and stirring was continued for 3 hours. The oil was extracted into chloroform and the chloroform extract was washed with 0.5 N hydrochloric acid and then with water and dried over magnesium sulfate. The dried extract was evaporated in vacuo to remove the chloroform and the residual liquid was distilled *in vacuo* to yield 43.2 g. (75%) of nearly color-less liquid, b.p. 197-203° (0.3 mm.), n²⁵D 1.5353.

Procedure E. Ethyl 4-p-Nitrobenozyl-1-piperazinecarboxylate (XXVI) .-- A mixture of 31.6 g. (0.2 mole) of ethyl 1-piperazinecarboxylate, 37.1 g. (0.2 mole) of p-nitrobenzoyl chloride and 33.6 g. (0.4 mole) of sodium bicarbonate in 300 ml. of water was stirred at room temperature for 7.5 hours and then heated on a steam-bath for 30 minutes. The resulting precipitate was removed by filtration and crystallized twice from absolute ethanol (using Norit), yielding 27.5 g. (45%) of very pale yellow crystals, m.p. $91-92^{\circ}$. When recrystallized from absolute ethanol the

product had m.p. 91.5–92.5°. Procedure F. Ethyl 4-Isobutyryl-1-piperazinecarboxylate (XII).—To a cold stirred solution of 150 g. (0.95 mole) of ethyl 1-piperazinecarboxylate in 1 1. of ethyl acetate, 50 g. (0.47 mole) of isobutyryl chloride was added dropwise. After standing for 1.5 hours at room temperature the mixture was filtered to remove 91.9 g. (100%) of colorless

crystals of ethyl 1-piperazinecarboxylate hydrochloride.22 The filtrate was distilled to remove the ethyl acetate and the residual red-brown liquid was distilled in vacuo. The prod-

restuar red-brown inquid was distined in vacuo. The product, 89 g. (97%), was obtained as a colorless liquid, b.p. 130-131° (0.5 mm.), n²⁵D 1.4840.
Procedure G. Ethyl 4-(3-Methylvaleryl)-1-piperazine-carboxylate (XVIII).—3-Methylvaleryl chloride (13.5 g., 0.1 mole) was added dropwise to 34.8 g. (0.22 mole) of ethyl 1-piperazinecarboxylate in 250 ml. of ether with cooling. After stending coverside to a remember of the stendard ste cooling. After standing overnight at room temperature the reaction mixture was filtered to remove 19.6 g. (100%) of ethyl 1-piperazinecarboxylate hydrochloride. The filtrate was washed with 1 N hydrochloric acid, water and 5% sodium bicarbonate, and dried over magnesium sulfate. The ether was removed on a steam-bath and the residual liquid was distilled *in vacuo* to yield 22.3 g. (87%) of XVIII as a colorless liquid, b.p. 137-140° (0.04 mm.), n^{25} D 1.4840.

Procedure H. Ethyl 4-Myristoyl-1-piperazinecarboxylate (XXIV).—To a solution of 63.3 g. (0.4 mole) of ethyl 1-piperazinecarboxylate in 350 ml. of ether, 49.4 g. (0.2 mole) of myristoyl chloride was added in portions with shaking and cooling. After standing overnight at room temperature the mixture was filtered to remove 38.4 g. (99%)b) of ethyl 1-piperazinecarboxylate hydrochloride The filtrate was washed with 5% sodium bicarbonate and dried over Drierite. The dried solution was heated on a area over Drierite. The area solution was heated on a steam-bath to remove the ether, leaving 71.1 g. (96%) of XXIV, m.p. 35-36.5°. Recrystallization from hexane gave colorless crystals, m.p. 36.5-38°.
Procedure I. Ethyl 4-n-Caproyl-1-piperazinecarboxylate (XVI).—To a cold solution of 63.3 g. (0.4 mole) of ethyl 1-piperazinecarboxylate in 350 ml. of ether, 26.9 g. (0.2 mole) of the procedure of a correct of bloride vac added corrective immediately.

producing a precipitate of ethyl 1-piperazinecarboxylate hydrochloride. After standing at room temperature for 2 hours 100 ml. of water was added to dissolve the precipitate. The layers were separated and the ether layer was washed successively with 5% hydrochloric acid, water and 5%sodium bicarbonate, and dried over magnesium sulfate. The ether was removed on a steam-bath and the residual The product was obtained as a nearly colorless liquid, b.p. $130-136^{\circ} (0.05-0.08 \text{ mm.}), n^{23.8} \text{D} 1.4835.$

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[CONTRIBUTION FROM THE DEPARTMENT OF CHEMISTRY OF HARVARD UNIVERSITY AND THE DIVISION OF NUTRITION AND PHYSIOLOGY OF THE PUBLIC HEALTH RESEARCH INSTITUTE OF NEW YORK CITY]

The Synthesis of 4-Amino-2(3H)-oxo-5-imidazolecarboxamide

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4-Amino-2(3H)-oxo-5-imidazolecarboxamide has been synthesized by the action of base on carboxamidoaminocyanoacetamide. Its structure has been proved by hydrolytic degradation to 2,4-dioxo-5-imidazolecarboxamide and to hydantoin. Biological testing of C¹³-labeled material gave no evidence of its being a precursor of uric acid.

Introduction

In 1945, Stetten and Fox³ discovered a new diazotizable amine in cultures of E. coli whose growth was inhibited by sulfadiazine or sulfapyridine. The amine was subsequently identified by Shive, et al.,⁴

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as 4-amino-5-imidazolecarboxamide (I). Isotopic studies have demonstrated this compound to be a purine precursor in a number of biological systems including yeast,⁵ the pigeon,⁶ the rat⁷ and man.⁸ In man, with a ureotelic nitrogen metabolism, uric

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