measurement was determined on 15 and was found to be consistent with the formula $C_{19}H_{16}^{16}O_2^{18}O$ within 4 ppm. The location of the ¹⁸O in warfarin was checked by the ratio of m/e 267/m/e 265 fragment ions (Table II).^{2a}

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Naphthylalkyl Lactamimides as Inhibitors of Blood Platelet Aggregation

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Hexahydro-2-[1-(1-naphthyl)ethylimino]azepine \cdot HCl (2) (RMI 7822) was found to inhibit human blood platelet aggregation induced by ADP and other agents with minimal release of procoagulant platelet factor 3. The compound was selected after careful modification of structural parameters, such as α -substitution, lactam ring size, aromatic substitution, as well as evaluation of its optical isomers.

Arterial thrombosis, especially in arteries supplying heart muscle and brain, is a leading cause of death and disability.¹ The blood platelets play a dominant role in such thromboses, both in the initial event and at the occlusive stage.² The so-called white thrombus that develops before fibrin clot formation is composed primarily of aggregated platelets. A number of tissue constituents can initiate platelet aggregation.³ Among these substances adenosine diphosphate (ADP) seems to be especially important, since it not only can initiate aggregation, but appears to mediate aggregation due to other agents as well.⁴ Physiologic functions of platelets include hemostasis and possible repair of vascular endothelium.⁵ In performing these functions platelets must undergo controlled adhesion and aggregation. Agents that would normalize abnormal platelet functions in the thrombosis-prone individual would have great therapeutic value. For these reasons we have adopted and developed techniques to evaluate large numbers of compounds available to us from earlier synthetic programs in our laboratories. Using human platelet-rich plasma (PRP) we measured in vitro inhibition of platelet aggregation induced by ADP, collagen, and several other agents.⁶ Compounds that showed activity were then checked for release of platelet factor 3 (PF3) or PF3like activity by measuring Stypven time. PF3 is a phospholipoprotein that acts as a cofactor in the coagulation process. Since we had established earlier that a normal breakfast causes PF3-like activity of from 0.1 to 0.3%, we adopted this as our limit of acceptability.7 Selected compounds were then evaluated further. We reported earlier our findings on certain piperidineethanols of benzyl- and benzylidenefluorene,^{8,9} and on a member of a series of anilines,¹⁰ obtained from Zellner.¹¹ We now wish to report our findings on certain lactamimides.

An exploratory series of lactamimides was prepared by one of us (E.M.R.) as potential antihypertensive agents. Later one of these, compound 2^{\dagger} in Table I, was found to inhibit ADP-induced platelet aggregation. We then explored systematically the effect of structural modifications on this activity by preparing and evaluating the compounds listed in Table I. This study revealed that activity is distributed broadly throughout the series. The structural parameter that most affects inhibition of platelet aggregation in this series was found to be the substituent R on the carbon atom adjacent to the lactamimide function (compounds) 1-6). The unsubstituted congener 1 was found to be less active than several of the alkyl-substituted congeners, and of these the methyl-substituted compound 2 showed least effect on PF3 release. The sterically hindered tert-butyl congener 5 and the phenyl-substituted congeners 16-19 were less active. Exploration of the influence of the lactam ring size (compounds 2, 7-10) showed that the 5- and 6membered lactam ring congeners are less active (cf. also 14 vs. 13 and 16 vs. 17), while the larger 8- and 13-membered ring congeners showed enhanced PF3 release. Several examples of aromatic substitution (compounds 11-14, 16, 17) revealed no trends, nor did the structural changes represented by compounds 15 and 20. No significant differences were found between the d and l isomers of 2.

Compound 2[†] also inhibited aggregation induced by thrombin, epinephrine, and serotonin. It did not inhibit clot retraction and this property may be advantageous. More detail on these and additional evaluations will be reported elsewhere.

Lactamimides, also named cyclic or semicyclic amidines,¹⁹ exist in two tautomeric forms A and B. This tautomerism



has been studied by Kwok and Pranc.²⁰ It is not known, however, which tautomer prevails in the crystalline monohydrochloride salts, not to mention solutions under physio-

Table I. Naphthylalkyl Lactamimides. Effects on Human Blood Platelets



							[Effects on human blood platelets ^d			
								\sim			% inł	nibition	
No.	R	ł	۲ ^۱	n	х	Mp, ^a °C	Recrystn solvent ^b	% yield	Formula ^c	Concn, µg/ml	of aggre	gation vs. Collagen	Release of PF3
1	Н	н		5	Н	214-215	A	51e	C ₁₇ H ₂₀ N ₂ ·HCl	300			0.004 (2)
										100	46 (2)	89 (2)	0.002 (2)
										30	15 (2)	33(2)	
dl- 2	CH,	н		5	н	224-232	Α	56 <i>e</i>	C ₁₈ H ₂₂ N ₂ ·HCl	300	0	15(2)	0.037
	•								10 22 2	100	70 (2)	100 (2)	0.003
										30J	46 (2)	54 (2)	
d-2						252-253	В	38 <i>e,g</i>		300	0(2)	30 (2)	0.063
							-	•••		100	83 (4)	91 (3)	0.022
										30	20 (5)	72 (3)	
<i>l</i> -2						252-253	R	36 <i>e</i> ,h		300	15 (2)	42 (3)	0.150
• =							2	50 /		100	56 (8)	87	0.013
										30	14 (9)		
2	сн сн	ч		5	U	282-284 dea	٨	nci		10	11 (4)		
5	CI12CI13			5	14	205-204 400	A	70-	C19H24N2 HCI	30	18		
4	$CH(CH_3)_2$	н		5	н	290-291 dec	Α	62 <i>i</i>	$C_{20}H_{26}N_2 \cdot HCl$	300			0.46
										100	91	100	0.014
										30 10	14	94 63	
5	C(CH ₃) ₃	н		5	н	>300	Α	56 <i>k</i>	$C_{21}H_{28}N_2 \cdot HCl$	100	71	05	
										30	10		
6	СНл	ч		5	ц	200-201	C	711		10	0		2.0
v	C411971			5		200-201	C	/4/	C ₂₁ H ₂₈ N ₂ ·HCI	100	79 (3)	100	0.12
										30	31 (3)	97	
7	CU	т		2		200 201 4		07 <i>h</i>		10	18	22	
'	CH ₃	п		3	н	300-301 dec	A	3/"	$C_{16}H_{18}N_2 \cdot HCI$	300	55 (2)	88	<0.001
										30	29 (2)	81	20.001
8	CH,	Н		4	н	258-259	Α	28 <i>h</i>	$C_{17}H_{20}N_2 \cdot HCl$	300			0.002
										100	55 (2)	70	<0.001
9	CH.	н		6	н	234-236	Α	59h	C.H.N. HCl	300	9	21	0.30
	3			•					019-24-12 1101	100	87	100	0.037
10	CU	TT		1 1		217 210		3 0 h		30	16	86	• • •
10	Cn ₃	п		11	н	217-219	A	28"	$C_{24}H_{34}N_2$ ·HCI	300	100	100	2.60
										30	50	100	0.50
	CII			~	4 (7)					10	15	0	
11	CH ₃	н		3	4-C1	263-265	D	50 <i>m</i>	$C_{18}H_{21}CIN_2 \cdot HCI$	300	01 (2)	08 (2)	0.75
										30	22(3)	80 (2)	0.10
				_	_					10	10	43 (2)	
12	Н	н		5	2-CH ₃	223-228	Α	82 <i>e</i>	$C_{18}H_{22}N_2 \cdot HCl$	300	50 (0)		0.021 (2)
										30	32 (2) 29 (2)		0.004 (2)
13	CH3	н		5	5,8-Me ₂	265-267 dec	Α	5 n	C ₂₀ H ₂₆ N ₂ ·HCl	300	27 (2)		2.30
										100	96 (4)	100 (2)	0.088
										30	37 (4)	89 (2) 21 (2)	
14	CH3	н		4	5,8-Me ₂	205-207 dec	Е	66 <i>n</i>	C ₁ ,H ₂₄ N ₂ ·HCl	300	0(2)	21 (2)	0.3
					-					100	96	100	0.032
										30	16	89	
15	CH3	5-tert-	Bu	5	н	>300	Α	19 <i>e</i>	C ₂₂ H ₂₀ N ₂ ·HCl	100	93	22	
										30	0		
16	C ₆ H₅	н		4	4-F	197-199	Α	690	$C_{22}H_{21}FN_2 \cdot HC1$	100	87 11		
17	C₅H₅	н		5	4-F	278-279	Α	730	C₂₃H₂₃FN₂·HCl	100	90		
18	C,H.CH0	н		5	н	290-292	А	64 <i>P</i>	C.H.N.HCI	30 100	20 85		
10	снсп∼	н		5	цо	208 200 4		A 6 4	C U NI UCI	30	36		
17	CI14CH3-0	п 		5	П Ч	308-309 dec	A	40'	C24H26N2·HCI	30	66 18		
20	CH3	н		3	HY	251-252	D	23 <i>e</i>	$C_{18}H_{22}N_2$ ·HCl	300 100	80 (2)		0.15
										30	32 (2)		0.007
										10	5 (2)		

Table I (Continued)

	Effects on human blood platelets ^{d}				
Compound	Concn, µg/ml		% inhibition of aggregation vs. ADP Collagen		
α-[p-(Fluoren-9-ylidenemethyl)phenyl]-2-piperidineethanol glycolate (RMI 10,393) ^s	300			0.747 (6)	
	100	99 (4)	100(2)	0.162 (6)	
	30	28 (6)	39 (2)	0.003 (7)	
	10	1(6)	0 (2)		
$N-(2-(Diethylaminoethyl)-N-(2-hydroxy-2-phenylethyl)-2,5-dichloroaniline (AN162)^t$	300			0.52 (42)	
	100	62 (118)	75 (35)	0.10 (47)	
	30	10 (104)	11 (35)		
2,4,6-Trimorpholinopyrimido[5,4-d]pyrimidine (RA 433 BS] ^u	300				
	100	47 (2)		< 0.001	
	30	25		< 0.001	
	10	15			
2-(5,10-Dihydrothiazolo[3,2-b][2,4]benzodiazepin-3-yl)phenol·HBr ^v	300			< 0.001	
	100	85		< 0.001	
	30	45			

^aMelting points were detd on a Hoover capillary melting point apparatus and are corrected. ^bA = MeOH-Me₂CO, B = anhyd EtOH, C = MeCN, D = MeOH-Me₂CO, E = MeOH-Me₂CO-Et₂O. ^cAll compds were analyzed for C, H and one other element. Analytical results obtained for these elements were within ±0.4% of calcd values. ^dSee Experimental Section for method of biological evaluation. Values in parentheses refer to number of determinations. ^eFrom commercially available starting material as HCl salt. ^fThis corresponds to $1 \times 10^{-4} M$. ^gFrom dextrorotatory amine; for d-2, $[\alpha]^{25}D + 14.4^{\circ}$. ^hFrom levorotatory amine; for l-2, $[\alpha]^{25}D - 9.6^{\circ}$. ⁱFrom α -ethyl-1-naphthalenemethylamine·HCl, mp 269-270°, lit.¹² mp 281-282°. ^jDescribed in the Experimental Section. ^kFrom α -tert-butyl-1-naphthalenemethylamine·HCl, mp 276-278° dec, obtained in 9% yield from *tert*-butyl cyanide; the presence of the *tert*-butyl group was confirmed by nmr; the main product (41%) was *tert*-butyl α -naphthyl ketone, mp 78-81, lit.¹³ mp 73-74°. ^lFrom α -teutyl-1-naphthalenemethylamine·HCl, mp 246-247° dec; for free base cf. Schultz, et al.¹⁴ m See Experimental Section. ⁿFrom α -(4-fluoro-1-naphthalenemethylamine·HCl, mp 290-292° dec, obtained in 63% from 4,8-dimethyl-1-acetonaphthone, mp 49-51°. ^oFrom α -(4-fluoro-1-naphthyl)benzylamine·HCl, mp 290-292° dec, obtained in 52% yield from benzonitrile. ^pFrom α -o-tolyl-1-naphthalenemethylamine·HCl, mp 302-306°; for free base see Cervinka, et al.¹⁵ 9β-Naphthyl derivative. ^rFrom α -o-tolyl-2-naphthalenemethylamine·HCl, mp 306-309° dec, obtained in 49% yield from o-tolunitrile. ^sRef 8 and 9. ^tRef 10. ^wRef 16 and 17. ^vRef 18.

logic conditions. For the sake of convenience we have represented and named all compounds in the tautomeric form A.

The compounds listed in Table I were prepared, with one exception, from the appropriate primary amine hydrochloride C, and the appropriate *O*-methyl lactim D by the method



first described by Benson and Cairns.²¹ Yields were generally good (Table I).

Two examples are described in the Experimental Section. We preferred the method using no, or very little, solvent particularly for reactions with sterically hindered amines. Compound 10 was prepared by the procedure of Bredereck^{22,23} in which the complex formed from a lactam and POCl₃ was used with a primary amine as the free base or the hydrochloride salt. No effort was made to improve the low yield obtained with this procedure.

The primary amines were prepared either by Leuckart reaction from the corresponding ketones or by *in situ* LiAlH₄ reduction of the addition products of Grignard reagents to nitriles. An example of each of these reactions is given in the Experimental Section and the footnotes to Table I give appropriate detail and references.

The ir spectra of lactamimides show C=N stretching vibrations that vary with ring size: 7 (1675 cm⁻¹), 8 (1650), 2 (1640), 9 (1645), and 10 (1650), as expected.

Experimental Section[‡]

dl-Hexahydro-2-[1-(1-naphthyl)ethylimino] azepine · HCl (2). A mixt of 176.4 g (0.85 mole) of dl- α -(1-naphthyl)ethylamine · HCl and 118.5 g (0.932 mole) of O-methylcaprolactim in 950 ml of MeOH was refluxed for 3 hr. The solvent was evapd and the residue crystd and recrystd from MeOH-Me₂CO to give 143.1 g (56%) of dl-2 (Table I).

2-[1-(1-Naphthyl)ethylimino]azacyclotridecane • HCl (10). To 21.7 g (0.11 mole) of 2-azacyclotridecanone in 200 ml of C_6H_6 was added dropwise 15.3 g (0.10 mole) of POCl₃, and the mixt was stirred at room temp for 4 hr under exclusion of moisture. Then 17.1 g (0.10 mole) of $l-\alpha$ -(1-naphthyl)ethylamine was added and the mixt was stirred at room temp for 1 hr and at reflux temp for 4 hr. The mixt was allowed to stand overnight, 2 N HCl was added, and the organic phase, after addn of some CH₂Cl₂, was dried (Na₂SO₄) and evapd to dryness. The residue was crystd and recrystd from MeOH-Me₂CO to give 10.8 g (28%) of 10 (Table I).

2-{[2,2-Dimethyl-1-(1-naphthyl)propyl]imino} hexahydroazepine-HCl (5). A slurry of 7.5 g (0.03 mole) of powdered α -tert-butyl-1naphthalenemethylamine • HCl in 8 ml of O-methylcaprolactim was allowed to stand at room temp for 5 days with occasional stirring. The mixt became nearly homogeneous and then solidified, Small portions of anhyd EtOH were added to keep the mixt stirrable. When the mixt ceased to further solidify, it was cooled and the product was collected and recrystd (Table I). This method is particularly suited for reactions with sterically hindered amines.

2-Methyl-1-(1-naphthyl)propylamine • HCl. To Grignard reagent, prepd from 161.0 g (1.31 mole) of BrCHMe₂ and 31.8 g of Mg turnings in 300 ml of Et₂O, was added 800 ml of PhMe, Et₂O was distd off, 50.0 g (0.327 mole) of 1-cyanonaphthalene was added, and the mixt was refluxed overnight. (In some analogous prepns, this soln was added directly to LiAlH₄ in Et₂O to obtain the amine.) This mixt was decompd with hot 6 N HCl, sepd, washed (H₂O), and dried (Na₂SO₄), solvent was distd off from the PhMe layer, and the residue was distd to obtain 2-methyl-1-propionaphthone, 44.6 g (69%), bp 160–170° (0.6 mm), $n^{25}D$ 1.5925.

This material (30.3 g, 0.153 mole) was mixed with 30.7 g (0.486 mole) of HCOONH₄ and was slowly heated to 150° with stirring. After the initial foaming had subsided, the temp of the heating bath was raised to $185-190^{\circ}$ and stirring was continued for 3 hr. Upon cooling, the mixt was washed with H₂O, the washes were extd with a small amt of C₆H₆ and the ext was added to the residue along with 150 ml of concd HCl. This mixt was refluxed for 6 hr, and on cooling the product pptd and was collected, 34.6 g (85%), mp 287-288°. Recrystn from H₂O raised the mp by 1°. Anal. (C, H, N, HCl) C, H, N.

A nal. ($C_{14}H_{17}N \cdot HCI$) C, H, N. 4-Chloro- α -methyl-1-naphthalenemethylamine \cdot HCl. A mixt of 50.0 g (0.24 mole) of 4'-chloro-1'-actonaphthone²⁴ and 49.0 g (0.78 mole) of HCOONH₄ was treated as described in the preceding paragraph, 43.1 g (73%), recrystd from MeCN-MeOH, mp 293-294°. Anal. ($C_{12}H_{12}CIN \cdot HCI$) C, H; N: calcd, 5.79; found, 5.22.

Biological Methods. Blood Collection and Isolation of Plasma. Whole blood was obtained from voluntary, experienced donors before breakfast. Donors were instructed to take no drugs, specifically aspirin, for 5 days before giving blood. If the plasma was lipidemic or, in a preliminary aggregation experiment, showed no second phase aggregation (aspirin-like effect), this plasma was not used. Blood was collected by the 2-syringe technique. It was decalcified with 3.8% sodium citrate soln (1:9 with blood). The citrated blood was centrifuged at 100g for 10 min and citrated platelet-rich plasma (PRP) was isolated. Platelet-poor plasma (PPP) was isolated by recentrifuging the blood residue at 1500g for 15 min.

Inhibition of Platelet Aggregation. Compounds were tested for inhibition of ADP- and collagen-induced aggregation in a Bryston platelet aggregometer by the procedure of Mustard, *et al.*⁶ Human platelet-rich plasma (PRP) was diluted with autologous plateletpoor plasma to 400,000 platelets/mm³. Solns of test compd were prepd and added to obtain the indicated concs. Saline was added to another sample of the same plasma to serve as control. After incubation for 20 min at 37°, ADP (2 µg/ml of final concentration) was added to induce aggregation. Platelet aggregation produces an increase in light transmittance (ΔT) through the plasma sample in the aggregometer and this response was recorded on a Bausch and Lomb VOM-5 chart recorder. The maxima of the ΔT responses for control and test sample were then used to calculate per cent inhibition of platelet aggregation by the test compound.

Collagen was prepared by the method of Hovig²⁵ and was standardized.⁹ The values given in Table I refer to inhibition of the initial slope of the aggregation curve, as discussed elsewhere.¹⁰

Platelet Factor 3 Activation. A soln of the test compd was added to human citrated PRP and incubated at 37° for 20 min, and a modified Stypven test was performed. The plasma was diluted 1:10 for this modified test.⁷

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Treloxinate and Related Hypolipidemic 12*H*-Dibenzo[*d*,*g*][1,3]dioxocin-6-carboxylate Derivatives¹

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Synthetic studies on o-phenylenedioxyacetic acids led to preparation of 2,10-dichloro-12H-dibenzo[d,g]-[1,3] dioxocin-6-carboxylic acid, and its methyl ester, treloxinate, was found to be a potent hypolipidemic agent. Structural modifications of treloxinate and the effect of these on hypocholesterolemic and hypotriglyceridemic activity were explored in rats (Wistar strain). Variation of aromatic substitution patterns and the size of the central heterocyclic ring resulted in decrease or loss of activity. A general synthetic method was developed to prepare analogous tricyclic acids from bisphenols and excess potassium dichloroacetate in hydroxylic solvents. Yields were affected by the size of the central ring and by aromatic substitution but surprisingly little by steric hindrance.

A large number of alkylcarboxylic acids with aryl or aryloxy substituents have been reported to have hypolipidemic activity.² Clofibrate, ethyl 2-(*p*-chlorophenoxy)-2-methylpropionate, is one of the more effective of these agents, and is the most widely used for control of hyperlipidemias associated with atherosclerotic cardiovascular diseases. This