Antiradiation Agents. 3-[(Alkylthio)alkyl]thiazolidines and Substituted 2-{[(3-Thiazolidinyl)alkyl]thio}pyridines and -quinolines[†]

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5-Halo- and 3,5-dihalo-2-{[(3-thiazolidinyl)alkyl]thio} pyridine hydrochlorides [alkyl = $(CH_2)_{3-8,10}$] and substituted and unsubstituted 2-{[(3-thiazolidinyl)alkyl]thio}quinoline hydrochlorides [alkyl = $(CH_2)_{3-6,10}$] have been prepared by alkylation of 2(1H)-pyridinethiones and 2(1H)-quinolinethiones with 3-(chloroalkyl)thiazolidines. The series was extended to 3-[(alkylthio)alkyl]thiazolidine hydrochlorides [$CH_3(CH_2)_{n_1}$ -S-($CH_2)_{n_2}$, involving 27 combinations of $n_1 = 0-9$ and $n_2 = 2-7$] by alkylation of alkanethiols. Several compounds exhibited promising antiradiation activity, either by ip or po administration. Generally the compounds were more active in the po test. 5-Chloro-2-{[7-(3-thiazolidinyl)heptyl]thio}pyridine hydrochloride (19) given po at 75 mg/kg (ca. 0.13 LD₅₀) afforded 60% survival in the 30-day test. 3-[5-(Pentylthio)pentyl]thiazolidine hydrochloride (43) resulted in 92% survival (30-day) of the mice when given po at 125 mg/kg (0.2 LD₅₀).

A principal objective in the search for antiradiation agents has been to develop a drug which is effective on oral administration. In contrast with the many compounds effective by parenteral dosing, few agents have been reported active when given orally.¹⁻⁷ Selected pyridyloxyalkyl derivatives of thiazolidine were active in the po test⁴ and, in fact, 5halo-2-{[5-(3-thiazolidinyl)pentyl]oxy}pyridine and the corresponding hexyl derivative were more active po than ip. We now report a related series of thioethers, substituted 2-{[(3-thiazolidinyl)alkyl]thio}pyridines I (Table I) and -quinolines II (Table I), and a new group of simple alkyl thioethers, 3-[(alkylthio)alkyl]thiazolidines III (Table II). The same type of potentially useful antiradiation activity has been found.

3-(Chloroalkyl)thiazolidines were synthesized for alkylations of the 2(1H)-pyridinethiones, 2(1H)-quinolinethiones, and alkanethiols. 3-Thiazolidinealkanol hydrochlorides IV suspended in THF were readily converted (SOCl₂) to alkyl chlorides V. Other solvents or lack of solvent led to decomposition of the sensitive thiazolidine ring. Alkylation of thiols using these alkyl chlorides was accomplished in DMF (NaH).

Figure 1 shows correlations between the length of the thioether alkyl chain of the 5-halopyridine compounds and antiradiation activity in mice in the ip and po tests.⁸ The



activity is expressed as protective index values[‡] which incorporate both dose response and therapeutic index factors. In both test systems the 5-chloro and 5-bromo derivatives were more active than the 5-iodopyridines, with 5-chloro substitution being preferred for optimum activity by the oral route. With appropriate halogen substitution on the pyridine ring, good activity was obtained with pentyl, hexyl, or heptyl thioethers, regardless of route (ip or po) of administration. 5-Chloro-2-{[7-(3-thiazolidinyl)heptyl]thio}pyridine ·HCl (19) is the compound of choice in the



Figure 1. Effect of halogen substitution and chain length on antiradiation activity (activity expressed as protective index values[‡]) of 5-halo-2-{[(3-thiazolidinyl)alkyl]thio}pyridine hydrochlorides: 1, X = Cl; 2, X = Br; 3, X = I.

pyridine series because of its strikingly good peroral activity, but it is rated inactive when given ip. Perorally a dose of 75 mg/kg (ca. 0.13 LD₅₀) of **19** resulted in 60% survival in the 30-day test. The corresponding hexyl (15) and octyl (21) ethers also were highly active perorally. The 5-bromo derivative as a pentyl ether 10 is a good agent, in view of its effectiveness in both test systems. Fair activity was obtained with the 3,5-dichloropyridine derivatives (Table I), but corresponding dibromo compounds were only slightly active. Results using pyridine substituents other than halogen in the oxygen ethers discouraged us from employing those substituents in this thioether series. Of the quinoline thioether derivatives (Table I) 2-{[5-(3-thiazolidinyl)pentyl]thio]quinoline 2HCl (26) was the most active compound, although the hexyl thioether 29 compared favorably in the po test.

Certain 3-[(alkylthio)alkyl]thiazolidines III (Table II) also

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 $[\]pm$ Protective index = (protection factor) × (LD₅₀/min effective dose), where doses are in mg/kg and the protection factor is 1.4 for 40% survival, 1.5 for 50% survival, etc. 30% survival is the smallest value used for the calculations. 2-Aminoethanethiol (MEA) is the standard for comparison. At 150 mg/kg ip of MEA, 87% survival of mice can be obtained in the 30-day test. Its ip LD₅₀ is *ca*. 250 mg/kg and it is rated ++. The po LD₅₀ is *ca*. 625 mg/kg. At 300 mg/kg, 73% survival can be obtained in the po test giving MEA a rating of ++.

Antiradiation activity									
Intraperitonea	l data		Peroral data						
Drug dose, mg/kg	Survival, %	Rating	LD 50, <i>ca.</i> mg/kg	Drug dose, mg/kg	Survival %				
100	7	0	950	500	7				
50	0	0	625	170	0				
150	13	0	>600	300	0				
150	7	0	>650	300	7				
100	7	0	650	300	26				
100	13	0	550	300	20				

Table I. S	Substituted 2	2-{[(3	-Thiazolidiny	vl)alk	vilthio	vridines	(I) ar	nd -a	uinolines ((II)	
			T TTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTTT	, . ,			~ / ~ ~			~~/	



^aAll compounds were analyzed for C, H, N, and S. ^bThe antiradiation data represent the lowest dose of drug for which a high rate of survival in the 30-day test was obtained. For each test (see ref 7), usually 15 mice were treated with drug and irradiated either 15 or 30 min later. The radiation dose was 950 rads (30-50 rads/min) of γ radiation from a Cobalt-60 source. Faster radiation doses are noted. Ratings are based on the following ranges of protective indices (footnote t in the text): 0. 0-1:+, 2-5:++, 6-10:+++, 11-14. cCa. 200 rads/min. 975 rads total. dS: calcd, 14.41; found, 15.18. A high S analysis is attributed to the interference of free I, in the sample preparation. Additional analysis for Cl. eS: calcd, 14.38; found, 14.84. Additional analysis for H₂O by nmr showed 1 molar equiv.

Table II. 3-[(Alkylthio)alkyl]thiazolidines (III)

		·				CH	₃ (CH ₂) _{n1} -S-(CH	$I_2 n_2 - N $	CI					
										Antiradia	tion activity ^b			
								Intraperitor	neal data			Peror	al data	
No.	<i>n</i> ₁	n_2	Recrystn solvents	Yield, %	Mp,°C	Formula ^a	LD ₅₀ , <i>ca</i> . mg/kg	Drug dose, mg/kg	Survival, %	Rating	LD _{so} , <i>ca</i> . mg/kg	Drug dose, mg/kg	Survival, %	Rating
31	5	2	EtOAc	46	166-168	C11H22NS2.HCl	250	62	33	+	>600	300	67	+
32	6	2	Me,CO	34	167-170	C ₁₂ H ₂ , NS ₂ HCl	250	100	33	+	550	250	33	+
33	7	2	Me,CO	36	1 66- 170	C, H, NS, HCI	150	100	13	0	>600	400	53	+
34	8	2	Me	39	166-169	C ₁ , H ₂ , NS ₂ , HCl	250	100	7	0	>350	300	0	0
35	3	3	Me,CO	11	134-137	C, H, NS, HCi	300	80	0	0	>600	300	7	0
36	5	3	Me ₂ CO	41	147-154	C ₁ ,H ₂ ,NS ₂ · HCl	120	75 c	67	+	>300	200 ^c	60	+
37	6	3	Me ₂ CO, EtOAc		149-156	C, H, NS, HCl	1.75	50	13	- 0	500	300	60	+
38	7	3	Me ₂ CO, EtOAc	5	158-162	C ₁₄ H ₂₉ NS ₂ ·HCl	125	50	7	0	>600	4 0 0	80	+
39	8	3	Me,CO, EtOAc	10	159-165	C, H, NS, HCI	250	100	47	+	>600	60 0	73	+
40	6	4	Me ₂ CO, EtOAc	17	161-165	C, H, NS, HCI	175	80	67	+	600	300	40	+
41	2	5	EtOAc	16	145-147	C ₁₁ H ₂ NS ₂ HCl	300	75	0	.0	>600	300	7	0
42	3	5	EtOAc	43	151-155	C ₁₂ H ₂₅ NS ₂ ·HCl	160	60	73	+	900	500	40	+
43	4	5	EtOAc	24	160-163	C ₁₃ H ₂₇ NS ₂ ·HCl	125	50	67	+	650	125,63	92, 40	+++
44	5	5	Me,CO	38	162-166	C ₁ H ₂ NS ₂ HCi	75	30	73	+	500	125	87	++
45	6	5	EtŌAc	55	161-166	C ₁ ,H ₃ ,NS ₂ ,HCl	100	50	80	+	500	150	60	+
46	7	5	EtOAc	35	164-169	C ₁₅ H ₃₃ NS ₂ ·HCl	120	60	33	+	>525	400	80	+
4 7	9	5	EtOAc, MeCN	43	171-174	C ₁₈ H ₃₇ NS ₂ ·HCl	125	80	0	0	>900	300	0	0
48	1	6	Me ₂ CO	34	133-139	C ₁₁ H ₂₃ NS ₂ ·HCl	300	100	0	0	>600	300	0	0
49	2	6	EtŐAc	22	109-125	C12H25NS2·HCl	160	75	13	0	>500	200	0	0
50	3	6	EtOAc	7	139-143	C13H29NS2 HCl	>150	50	7	0	>300	300	33	0
51	4	6	EtOAc	39	159-161	C14H29NS2 HCl	150	80, 40	60, 38	+	500	200, 100	67, 47	++
52	5	6	Me ₂ CO	27	158-162	$C_{15}H_{31}NS_2 \cdot HCl^d$	125	60, 30	50, 47	++	175	90	53	+
53	6	6	Me ₂ CO, EtOAc	14	161-164	C16H33NS2 HCl	1 30	50	13	0	>1000	400	27	0
54	7	6	Me ₂ CO	33	159-164	C17H35NS2 HCl	>200	100	0	0	650	300	13	0
55	8	6	Me ₂ CO, MeCN	11	166-169	C18H37NS2 HCl	150	80	7	0	> 600	200	0	0
56	0	6	EtOAc-Me ₂ CO	12	116-121	C10H21NS2 HCl	100	60	13	0	500	300	0	0
5 7	3	7	EtOAc	54	150-153	C14H29NS2·HCl	175	80, 40	73, 46	++	650	300	67	+

^aAll compounds were analyzed for C, H, N, and S. ^bSee footnote b, Table I. ^cCa. 200 rads/min, 975 rads total. ^dS: calcd, 19.67; found, 19.12.



Figure 2. Effect of chain length on antiradiation activity (activity expressed as protective index⁺) of 3-[(alkylthio)alkyl] thiazolidine hydrochlorides given perorally.

Table III. 3-Thiazolidinealkanol Hydrochlorides, HO(CH₂) $_{n}$ -N $_{/}$ ·HCl (IV)

	3			
n	Yield, %	Mp, °C	Formula	Analyses
2	49	80-84	C.H. NOS · HCl	C, H, N, S
3	55	92-94	C.H. NOS HCI	C, H, N, S
4	13	93-99	C,H, NOS HCI	C, H, N, S
5	50	96-98	C.H. NOS HCI	C, H, N, S
6	71	112-116	C H, NOS HCI	C, H, N, S
7	50	101-105	C, H, NOS HCI	C, H, N, S
8	53	128-130	C, H, NOS HCI	C, H, N, S
10	32	140-143	C ₁₃ H ₂₇ NOS · HCl	C, H, N, S

Table IV. 3-(Chloroalkyl)thiazolidine Hydrochlorides, Cl(CH₂) $n \cdot N \longrightarrow HCl (V)$

	5				
n	Recrystn solvents	Yield, %	Mp, °C	Formula	Analyses
2	THF	94	129-131	C _s H ₁₀ CINS · HCl	$H, N, S; C^a$
3	n-BuOH	88	133-135	C ₆ H ₁₂ CINS HCi	C, H, N, S
4	THF,	27	120-124	C,H,CINS HCI	C, H, N, S
	Me ₂ CO				
5	THF	89	133-135	C ₈ H ₁₆ CINS HCi	C, H, N
6	THF	4 0	117-122	C ₀ H ₁₈ CINS HCI	C, H, N, S
7	MeCN	35	1 32-1 34	C ₁₀ H ₂₀ CINS · HCl	C, H, N, S
8	EtOAc	50	130-132	C ₁₁ H ₂₂ CINS HCi	C, H, N, S
10	Me ₂ CO	92	144-147	C13H26CINS HCi	C, H, N, S

^aC: calcd, 31.92; found, 31.42.

were highly effective when given by mouth. Generally these are more effective perorally than parenterally. Figure 2 illustrates the effect of variations in chain length on po antiradiation activity. A total of 10-12 carbons in the side chain is the optimum chain length. As can be seen from Figure 2, however, even within this range large differences in activity are evident with various combinations of n_1 and n_2 . The best compound is 3-[5-(pentylthio)pentyl] thiazolidine \cdot HCl (43). Given orally 30-min preirradiation at 125 mg/kg (0.2 LD_{50}), 43 afforded 92% survival of the mice in the 30-day test. At 63 mg/kg (0.1 LD_{50}) 40% survival was obtained in the same test. 3-[5-(Hexylthio)pentyl]thiazolidine \cdot HCl (44) had similar activity.

Thiazolidine substituted in the 3 position with (5-halo-2pyridyl)thioalkyl, (substituted-2-quinolyl)thioalkyl, and alkylthioalkyl groups resulted in effective antiradiation agents in mice. A 5-chloropyridyl thioether 19 and a (pentylthio)pentyl derivative 43 can be considered candidates for further study in the search for an antiradiation agent which is effective by the oral route.

Experimental Section §

3-Thiazolidinealkanol hydrochlorides IV (Table III) were prepared from 1-aziridinealkanols using the general procedure described⁴ previously. Crude products were recrystallized from MeCN.

3-(5-Chloropentyl)thiazolidine Hydrochloride (V, n = 5) (General Procedure for V, Table IV). To a slurry of 30.5 g (0.14 mol) of IV (n = 5) in 500 ml of THF was added in one portion 20 g (0.17 mol) of SOCl₂. The stirred mixture was purged continuously with a rapid stream of N₂, gradually heated to 45°, and kept at that temperature for 2-4 hr. The mixture, with continued stirring, was chilled and filtered. The solid was washed (as a slurry) in cold THF, dried, and recrystallized to give V (n = 5) (Table IV). In some cases Et₂O was added to the cold mixture to effect precipitation.

Substituted 2(1*H*)-pyridinethiones and 2(1*H*)-quinolinethiones (general procedure⁹): 5-chloro-,⁹ 5-bromo-,⁹ and 5-iodo-2(1*H*)pyridinethione;⁸ 3,5-dibromo-2(1*H*)-pyridinethione, mp 155–159° (methyl cellosolve at 120° was substituted for propylene glycol used in the general procedure,⁹ this solvent was preferable to propylene glycol in most cases); 3,5-dichloro-2(1*H*)-pyridinethione, mp 153– 156° [*Anal.* ($C_5H_3Cl_2NS$) C, H, N, S]; 6-chloro-4-methyl-2(1*H*)quinolinethione, mp 296–311° [*Anal.* ($C_{10}H_6CINS$) C, H, N]; 4methyl-2(1*H*)-quinolinethione, mp 258–262° [*Anal.* ($C_{10}H_9NS$) C, H, N]; 4,6-dimethyl-2(1*H*)-quinolinethione, mp 298–302°.

Alkylation of Thiols (Preparation of I-III, Tables I and II). A mixture of the substituted 2(1H)-pyridinethione, substituted 2(1H)-quinolinethione, or alkanethiol and 2 molar equiv of NaH in DMF was treated portionwise in the cold with 1 molar equiv of solid V. Mixtures containing the heterocycles were kept at $80-90^{\circ}$ for 4-6 hr, whereas those involving alkanethiols were kept on a steam bath for 4-5 days. The diluted (Et₂O) mixtures were washed with H₂O, dried (MgSO₄), and treated with 1 equiv of dry HCl giving I and II after recrystallization. The free bases of the aliphatic analogs were distilled under high vacuum before conversion to HCl salts (III).

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Melting points (uncorrected) were determined using a Thomas-Hoover melting point apparatus. Where analyses are indicated only by symbols of the elements, analytical results obtained for these elements are within $\pm 0.4\%$ of the theoretical values.