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3,801,609

N-SUBSTITUTED FATTY ACID AMIDE LUBRICANTS

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1 Claim

ABSTRACT OF THE DISCLOSURE

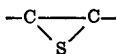
This invention relates to N-acylmorpholines and N-mono and N,N-disubstituted fatty acid amides and to similar derivatives of epithioamides which are useful as base and extreme pressure lubricants and additives.

This is a division, of application Ser. No. 176,734, filed Aug. 31, 1971, now Pat. No. 3,746,644.

A non-exclusive, irrevocable, royalty-free license in the invention herein described, throughout the world for all purposes of the United States Government, with the power to grant sublicenses for such purposes, is hereby granted to the Government of the United States of America.

This invention relates to certain compounds which are N-fatty acyl derivatives of primary and secondary amine and N-substituted acyl derivatives of primary and secondary amines which have exhibited utility as base lubricants, extreme pressure lubricants or lube additives. More particularly, this invention relates to N- and N,N-disubstituted long chain aliphatic amides the acyl component of which is a normal, branched, or substituted alkenoic or alkanolic acyl group containing from 16 to 22 carbon atoms the amide nitrogen of which may be derived from an alkyl amine, dialkyl amine, alkylalkoxyalkylamine, dialkoxyalkylamine or nitrogen heteroalicyclic. Typical amines are butylamines, dibutylamines, N-methyl-alkoxyethyl amine, di-ethoxyethyl amine, and morpholine. The acyl substituent referred to may be divalent sulfur or pentavalent phosphorous.

We have discovered that many of the simple N-alkyl and N,N-dialkyl or N-alkyl-N-alkoxyalkyl amides of the C₁₆ to C₂₂ alkenoic or alkanolic fatty acids are good base lubricants. We have further discovered that the introduction of divalent sulfur into the fatty acid moiety imparts excellent extreme pressure lubricant characteristics to such compounds. In particular we have found the epithio group



to be most effective. We have also observed that the introduction of both hydroxy and alkyl phosphato groupings alpha to each other also imparts extreme pressure lubricant characteristics to the compound.

In addition it was noted that the above hydroxy-phosphato compound and its mercapto analog N-[9(10)-mercapto-(9)10-dibutylphosphato]stearamide were effective antiwear compounds and additives.

Included among the specific compounds operable as base

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lubricants are N-methyl-N-butyleamide, N-ethoxyethoxy propyleamide, N-ethoxyethoxy propyleamide, N-methoxyisopropyleamide, N-methoxyethylamide. Compounds showing particular promise as extreme pressure lubricants or additives are N,N-dibutyl-9,10-epithiostearamide, N,N-dibutyl-9,10-12,13-diepithiostearamide, 9,10-epithiostearoyl morpholine, N-ethoxyethoxyethoxypropyl-9,10-epithiostearamide, N-9,10-12,13-diepithiostearoylmorpholine and N-[9(10)-hydroxy-9(10)-dibutylphosphato]stearamide.

Compounds showing particular promise as antiwear additives are N,N-dibutyl-[9(10)-hydroxy-(9)10-dibutylphosphato]stearamide and N,N-[9(10)mercapto-(9)10-dibutylphosphatostearoyl]morpholine.

EXAMPLE 1

N-methyl-N-butyleamide.—100 grams (0.33 mole) of oleoyl chloride was added dropwise and with stirring to a mixture of 29 grams (0.33 mole) of N-methyl-N-butylamine and 27 grams (0.33 mole) of pyridine. Stirring was continued until the heat of reaction subsided. The solid pyridine hydrochloride was filtered off and the filtrate was washed successively with aqueous HCl and water until acid free. It was dried, stripped, percolated through an activated alumina column, and the product removed from the percolate by stripping off the solvent. Analysis of the product: Percent C, 78.87 (theory 78.63); percent H, 13.02 (theory 12.82); percent N, 3.96 (theory 3.99).

EXAMPLE 2

N,N-dibutyleamide.—This material was prepared by the procedure of Example 1 from 100 grams (0.33 mole) of oleoyl chloride 42.5 grams (0.33 mole) of di-n-butylamine and 27 grams (0.33 mole) pyridine. Analysis of the product: Percent C, 78.94 (theory 79.25); percent H, 13.16 (theory 13.06); percent N, 3.44 (theory 3.56) confirm the preparation.

EXAMPLE 3

Ethoxyethoxyethoxy propyleamide.—This material was prepared by the procedure of Example 1 from 100 grams (0.33 mole) of oleoyl chloride, 73.0 grams (0.33 mole) ethoxyethoxyethoxypropylamine and 27.0 grams (0.33 mole) pyridine. Analysis of percent C, 71.84 (theory 72.04); H, 11.80 (theory 11.80), and N, 2.96 (theory 2.90) confirm the preparation.

EXAMPLE 4

Ethoxyethoxypropyleamide.—This material was prepared by the procedure of Example 1 from 100 grams (0.33 mole) of oleoyl chloride, 58.3 grams (0.33 mole) ethoxyethoxypropylamine and 27.0 grams (0.33 mole) of pyridine. Analyses of percent C, 71.45 (theory 73.80); percent H, 11.87 (theory 12.07); and percent N, 2.92 (theory 3.18) confirm the preparation.

EXAMPLE 5

Methoxyisopropyleamide.—This material was prepared by the procedure of Example 1 from 100 grams (0.33 mole) of oleoyl chloride, 29.3 grams (0.33 mole) methoxyisopropylamine and 27.0 grams (0.33 mole) pyridine. Analysis of percent C, 73.15 (theory 74.99); percent H, 12.01 (theory 11.93); and percent N, 3.67 (theory 3.97) confirm the preparation.

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EXAMPLE 6

Methoxyethyloleamide.—This material was prepared by the procedure of Example 1 from 100 grams (0.33 mole) of oleoyl chloride, 24.4 grams (0.33 mole) of methoxyethylamine and 27.0 grams (0.33 mole) pyridine. Analysis of percent C, 73.08 (theory 74.55); percent H, 11.98 (theory 11.83); and percent N, 3.31 (theory 4.14) confirm the preparation.

EXAMPLE 7

Oleoylmorpholine.—This material was prepared by the procedure of Example 1 from 100 grams (0.33 mole) of oleoyl chloride, 28.7 grams (0.33 mole) of morpholine and 27.0 grams (0.33 mole) of pyridine.

EXAMPLE 8

N,N-dibutyloleamide.—This product was prepared following the exact procedure of Example 2 except for the substitution of oleoyl chloride by linoleoyl chloride.

EXAMPLE 9

N,N-dibutyl-9,10-epithiostearamide.—120 grams (0.30 mole) of N,N-dibutyloleamide was epoxidized by the addition with stirring of 62.0 grams (0.36 mole) of meta-chloroperbenzoic acid in 630 ml. of chloroform. Reaction was continued for 1 hour beyond final addition and the excess peracid destroyed by a small amount of a 10% solution of Na_2SO_3 . The metachlorobenzoic acid was removed by a NaHCO_3 wash, followed by water washing. The CHCl_3 solution was then dried with anhydrous Na_2SO_4 and the CHCl_3 stripped off to recover the product N,N-dibutyl-9,10-epoxystearamide, oxirane content 3.45% (theory 3.91%). 100 grams (0.24 mole) of this product was added to a well stirred slurry of 55.7 g. (0.75 mole) of thiourea and 89.5 g. (0.72 mole) of benzoic acid in acetone. Stirring was continued for 3 hours beyond the terminal addition whereupon the benzoic acid was washed out with 38.8 g. (0.36 mole) of Na_2CO_3 and the organic product extracted with hexane. The organic phase extract was dried and stripped to recover the product N,N-dibutyl-9,10-epithiostearamide containing 6.41% sulfur (theory 6.67%).

EXAMPLE 10

N-(9,10-epithiostearoyl)morpholine.—This product was prepared by the exact procedure described in Example 9 except for the substitution of oleoylmorpholine for the N,N-dibutyloleamide of Example 9. Analysis of product showed percent C, 71.22 (theory 69.34); percent H, 11.22 (theory 10.77); percent N, 3.66 (theory 3.68); percent S, 3.44 (theory 7.80).

EXAMPLE 11

N,N-dibutyl-9,10-12,13-diepithiostearamide.—This product was prepared following the procedure described in Example 9 except for the replacement of the N,N-dibutyloleamide of Example 9 by N,N-dibutyl linoleamide, and a doubling of the molar ratios of all reagents. The product analyzed as follows: percent C, 69.64 (theory 68.57); percent H, 10.93 (theory 10.76); percent N, 3.10 (theory 3.07); percent S, 11.7 (theory 14.06).

EXAMPLE 12

N-ethoxyethoxyethoxypropyl-9,10-epithiostearamide.—To 166 g. of N-ethoxyethoxyethoxypropyl oleamide, Example 3, was added dropwise and with stirring 73.8 g. of m-chloroperbenzoic acid in 800 ml. of CHCl_3 . The spent m-chloroperbenzoic acid was removed with 10% aqueous NaHCO_3 and the epoxide recovered by drying and stripping off the CHCl_3 . 171 g. of this epoxide was added dropwise to a slurry of 56 g. thiourea in 1000 ml. of acetone and the concurrent addition of 43.4 g. of glacial acetic acid and the reaction mixture stirred for 3 hours. The glacial acetic acid was neutralized with Na_2CO_3 and the episulfide, N-ethoxyethoxyethoxypropyl-

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9,10-epithiostearamide, extracted with Skellysolve B and washed, dried, and stripped. Its elemental analysis was C=67.1 (66.52), H=11.42 (10.88), N=2.67 (2.87), S=2.29 (6.57).

EXAMPLE 13

N-ethoxyethoxyethoxypropyl-9(10)mercaptostearamide.—17 g. of N-ethoxyethoxyethoxypropyloleamide in CCl_4 was treated in the cold with a CCl_4 solution containing 60.9 g. of bromine. Any unreacted bromine was removed by a thiosulfate wash, 14 g. of the resulting dibromo compound was added dropwise to a well stirred solution of 72 g. of $\text{Na}_2\text{S}\cdot 9\text{H}_2\text{O}$ in dimethylsulfoxide maintained at 80° C. Stirring was continued at 80° C. for 3 hours. Water was added and the reaction product was extracted with hexane; washed, dried, and stripped. Elemental analysis of the product was C=66.3 (66.5), H=11.1 (10.9), N=2.7 (2.9), S=4.4 (6.8).

EXAMPLE 14

N,N-dibutyl-9(10)[dibutylphosphono]stearamide.—100 g. of N,N-dibutyloleamide, Example 2, 148 g. of dibutyl phosphite, and 1.29 g. of benzoyl peroxide catalyst were heated at 115° C. for 3½ hours. Additional units of 1.29 g. of benzoyl peroxide was added after the first and second hours of reaction time. The excess dibutyl phosphite was then removed by distillation at reduced pressure, 0.45 mm. Hg. The stillpot contents showed strong adsorption bands at 8, 9.3, and 9.7 microns characteristics of the phosphonate group. The elemental analysis was C=obs 71.46 (69.50), H=obs 12.42 (11.92), N=obs 2.20 (2.38), P=obs 4.24 (5.28) percent respectively.

EXAMPLE 15

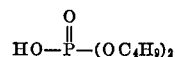
N-9,10-12,13-diepithiostearoylmorpholine.—This product was prepared by the same procedure described in Example 9 except for the substitution of linoleoyl morpholine for the N,N-dibutyloleamide of Example 8 and the doubling of molar proportions of the m-chloroperbenzoic acid in the epoxidation step and the thiourea and benzoic acid in the "epithioation" step. The elemental analysis was C=obs 68.89 (63.92), H=obs 10.26 (9.44), N=obs 3.29 (3.38), S=obs 4.00 (15.49) percent respectively.

EXAMPLE 16

N,N-Dibutyl-[9(10)-hydroxy-(9)10-dibutylphosphato]stearamide.—44.7 g. of dibutylhydrogen phosphate was added to 182.2 g. of N,N-dibutyl-9,10-epoxystearamide at 90–95° C. with stirring. Reaction conditions were continued for 3½ hours after termination of the addition. The reaction product was dissolved in commercial hexane and the unreacted dibutyl hydrogen phosphate neutralized and washed out with portions of aqueous 10% NaHCO_3 followed by several water washes. Acidulation of the hexane phase with diluted HCl followed by several water washes, drying and stripping, yielded the product. Elemental analysis was, C=obs 66.56 (65.91), H=obs 11.04 (11.30), N=obs 2.90 (2.26), P=obs 3.04 (4.20).

EXAMPLE 17

N-[9(10)mercapto-(9)10-dibutylphosphatostearoyl]morpholine.—12 g. of dibutylhydrogen phosphate was added dropwise with stirring to 51.0 g. of (9,10-epithiostearoyl)morpholine, the preparation of Example 10, at 85–90° C. and the heating and stirring continued for 3 hours beyond the terminal addition. The reaction product was dissolved in commercial hexane and any excess or unreacted



was neutralized and washed out with aqueous 10% NaHCO_3 . The hexane extract was washed several times with water, dried and stripped. The elemental analysis of the product was C=obs 66.74 (69.36), H=obs 10.69

(11.05), N=obs 2.89 (3.32), S=obs 3.27 (2.17), P=obs 1.68 (2.10) percent respectively.

EXAMPLE 18

The compounds prepared in accordance to the procedures given in Example 1 through 7 were evaluated as

lubricants or additives in a Shell 4-ball extreme pressure tester at 1440 p.p.m. following ASTM Procedure D 2596-67T. Loads were increased in increments of 20 kg. to weld point and the test ran for one minute or to weld whichever occurred first. Commercial hypoid fluid SAE #90 was employed as the control E.P. lubricant.

TABLE I
Antiwear lubricant properties of fatty acid amides

Compound	Avg. wear scar, mm.		Additive
	Without additive	With 5% additive	
1..... N, N-bis(2-ethoxyethyl)oleamide.....	0.597	0.735	N, N-di-butyl-9, 10-epithiostearamide.
2..... N, N-di-n-butyl-9, 10-epithiostearamide.....	0.842	None	
3..... Oleoylmorpholine.....	0.623	0.658	Do.
4..... N, N-dimethyloleamide.....	0.797	0.987	Do.
5..... N, N-di-n-propyloleamide.....	0.908	0.885	Do.
6..... N, N-di-n-hexyloleamide.....	0.893	0.898	Do.
7..... N, N-di-n-butyl-9, 10-epithiostearamide.....	0.758	0.798	Do.
8..... N, N-di-n-butyl amide of sel. hydrogenated cottonseed fatty acids.....	0.793	0.953	Do.
9..... N, N-bis(2-methoxyethyl)oleamide.....	0.626	0.725	Do.
10..... N, N-dibutyloleamide.....	0.710		
11..... N-oleyl-4-propylpiperidine.....	0.847		
12..... Morpholide of sel. hydrogenated cottonseed fatty acids.....	0.642		
13..... N-methyl-N-butyl-9, 10-epithiostearamide.....	0.607		
14..... N-ethoxyethoxyethoxypropyloleamide.....	0.503		
15..... N-ethoxyethoxypropyloleamide.....	0.526		
16..... N-methoxyisopropyloleamide.....	0.705		
17..... N-methoxyethyl-9, 10-epithiostearamide.....	0.420		
18..... 102 sec paraffin oil.....		0.902	N-methoxyethyl-9, 10-epithiostearamide.
19..... D.O.S.....		0.906	Do.
20..... N, N-dibutyl-9(10)-carboboxyoctadecanamide.....	0.610		
21..... 102 sec paraffin oil.....		0.818	N, N-dibutyl-9(10)-carboboxyoctadecanamide.
22..... D.O.S.....		0.945	Do.
23..... N-[9(10)mercapto-(9)10-dibutylphosphatostearoyl]morpholine.....	0.615		
24..... 102 sec paraffin oil.....		0.552	N-[9(10)mercapto(9)10-dibutylphosphatostearoyl]morpholine.
25..... D.O.S.....		0.535	Do.
26..... N, N-dibutyl-9(10)-hydroxy-(9)10-dibutylphosphatostearamide.....	0.498		
27..... 102 sec paraffin oil.....		0.498	N, N-dibutyl-9(10)-hydroxy-(9)10-dibutylphosphatostearamide.
28..... D.O.S.....		0.498	Do.
29..... N, N-dibutyl-9(10)-[dibutylphosphono]stearamide.....	0.530		
30..... N-ethoxyethoxyethoxypropyl-9, 10-epithiostearamide.....	0.707		
31..... N-ethoxyethoxyethoxypropyl-9(10)mercaptostearamide.....	0.690		
32..... N-(9, 10-12, 13-diepithiostearoyl)morpholine.....	0.665		
33..... Bis(2-ethylhexyl)sebacate.....	0.872		
34..... 100 sec paraffin oil.....		0.803	
35..... Aero Shell (Mil-7808).....		0.587	
36..... Gulf Pride, Single G, MS, multiviscosity.....		0.447	
37..... 100 sec paraffin oil.....		0.813	N, N-dibutyl-9, 10-12, 13-diepithiostearamide.
38..... Bis(2-ethylhexyl)sebacate.....		0.836	Do.
39..... 100 sec paraffin oil.....		0.723	N-9, 10-epithiostearoyl morpholine.
40..... Bis(2-ethylhexyl)sebacate.....		0.915	Do.
41..... 100 sec paraffin oil.....		0.917	N, N-dibutyl-9, 10-dichlorostearamide.
42..... Bis(2-ethylhexyl)sebacate.....		0.848	Do.

base lubricants in the Shell Four-Ball Wear tests following a modified procedure of ASTM D2266-67. The resulting scar diameter on the balls after running at 600 r.p.m. for 1 hour at 120° C. and 50 kg. load were compared with the scars obtained using commercial lubricants such as Aeroshell Mil-L-70D (di-2-ethylhexyl sebacate and additive), Gulfpride, single G, MS, multiviscosity, and 100 sec. paraffin oil. The results of these tests are reported in Table I.

It can be seen from the data presented that the N-alkoxyalkyl and N,N-di(alkoxyalkyl)oleamides, without the assistance of additives, exhibited wear characteristics (i.e., low scar diameters) comparable to the commercial controls Aeroshell and Gulfpride, which do contain additives to improve performance. N,N-dibutyl-[9(10)-hydroxy-(9)10-dibutylphosphato]stearamide and N-[9(10)mercapto-(9)10-dibutylphosphatostearoyl]morpholine not only exhibit low wear characteristics in themselves but also impart as additives such characteristics to paraffin oil and DOS base lubricants.

EXAMPLE 19

The various amides were evaluated as extreme pressure

The performance of the amides tested is given in Tables II, III, and IV.

TABLE II
Extreme pressure tests (ASTM D2596-67T)

Compound	Weld point	
	No additive	With 5% #2
1..... N, N-bis(2-ethoxyethyl)oleamide.....	120	170
2..... N, N-di-n-butyl-9, 10-epithiostearamide.....	300	
3..... Oleoylmorpholine.....	120	150
4..... N, N-dimethyloleamide.....	120	140
5..... N, N-di-n-propyloleamide.....	120	150
6..... N, N-di-n-hexyloleamide.....	120	180
7..... N, N-di-n-butyl-9, 10-epithiostearamide.....	120	200
8..... N, N-di-n-butyl amide of hydrogenated cottonseed fatty acids.....	120	170
9..... N, N-bis(2-methoxyethyl)oleamide.....	120	160
10..... N-methyl-N-butyl-9, 10-epithiostearamide.....	100	
11..... N, N-dibutyl-9, 10-epithiostearamide.....	300	
12..... N, N-dibutyl-9, 10-12, 13-diepithiostearamide.....	440	
13..... N-ethoxyethoxyethoxypropyloleamide.....	120	
14..... N-ethoxyethoxypropyloleamide.....	120	
15..... N-methoxyisopropyloleamide.....	100	
16..... N-methoxyethyl-9, 10-epithiostearamide.....	100	
17..... N-9, 10-epithiostearoylmorpholine.....	380	
18..... Bis(2-ethylhexyl)sebacate.....	120	
19..... 100 sec paraffin oil.....	100	
20..... SAE #90 commercial hypoid fluid.....	280	

TABLE III

Evaluation of N,N-dibutyl-9,10-12,13-di-epithiostearamide as an additive and as a base oil in extreme pressure tests. (ASTM D2596-67T)

Base oil	Percent additive	Load, kg.	Wear scar, mm.
100 sec paraffin oil	(5%)	140	1.90
Do	(5%)	200	2.35
Do	(5%)	220	2.75
Do	(5%)	240	Weld
Bis(2-ethylhexyl)sebacate	(5%)	120	1.51
Do	(5%)	180	1.59
Do	(5%)	190	2.47
Do	(5%)	200	Weld
Do	(10%)	240	2.88
Do	(10%)	260	3.55
Do	(10%)	270	Weld
N,N-dibutyl-9,10-12,13-di-epithiostearamide	None	120	1.38
Do	None	160	1.61
Do	None	200	1.89
Do	None	280	2.13
Do	None	320	2.47
Do	None	360	2.70
Do	None	400	2.88
Do	None	440	Weld

Extreme pressure tests on N-ethoxyethoxyethoxypropyl-9,10-epithiostearamide (PC-O-No. 7682)

Base oil	Additive	Applied load, kg.	Avg. wear scar, mm.
7682	None	120	1.05
7682	None	160	1.75
7682	None	240	2.35
7682	None	300	2.19
7682	None	340	2.55
7682	None	360	2.62
7682	None	380	Weld
102 sec paraffin oil	7682 (5%)	120	2.28
Do	7682 (5%)	140	2.46
Do	7682 (5%)	150	Weld
D.O.S.	7682 (5%)	120	2.23
D.O.S.	7682 (5%)	140	2.44
D.O.S.	7682 (5%)	150	Weld

Extreme pressure tests on N-ethoxyethoxyethoxypropyl-9(10)-mercaptostearamide (PC-O-No. 7683)

7683	None	200	1.54
7683	None	300	2.26
7683	None	400	2.25
7683	None	450	2.15
7683	None	500	2.15
102 sec paraffin oil	7683 (5%)	120	1.97
Do	7683 (5%)	140	2.35
Do	7683 (5%)	150	2.54
Do	7683 (5%)	160	Weld
D.O.S.	7683 (5%)	120	1.68
D.O.S.	7683 (5%)	140	2.13
D.O.S.	7683 (5%)	160	2.59
D.O.S.	7683 (5%)	170	2.72
D.O.S.	7683 (5%)	180	Weld

Extreme pressure tests on N-(9,10-12,13-di-epithiostearoyl)morpholine (PC-O-No. 7684)

7684	None	200	2.21
7684	None	260	2.38
7684	None	340	2.43
7684	None	400	2.75
7684	None	460	2.90
7684	None	500	2.98
102 sec paraffin oil	7684 (5%)	100	2.23
Do	7684 (5%)	120	2.61
Do	7684 (5%)	130	2.74
Do	7684 (5%)	140	Weld
D.O.S.	7684 (5%)	120	2.54
D.O.S.	7684 (5%)	140	2.58
D.O.S.	7684 (5%)	150	Weld
N-methoxyethyloleamide	7684 (5%)	140	2.53
Do	7684 (5%)	160	2.77
Do	7684 (5%)	170	Weld

Extreme pressure tests on N,N-dibutyl-9(10)-(dibutylphosphono)stearamide (PC-O-No. 7685)

7685	None	120	3.05
7685	None	140	4.50
7685	None	150	Weld

TABLE III—Continued

Extreme pressure tests on N,N-dibutyl-9(10)-hydroxy-9(10)-dibutylphosphato]stearamide (PC-O-No. 7686)

5	Base oil	Additive	Applied load, kg.	Avg. wear scar, mm.
	7686	None	200	0.887
	7686	None	300	2.78
	7686	None	340	4.32
	7686	None	350	4.50
	7686	None	360	Weld
10	100 sec paraffin oil	7686 (5%)	120	2.28
	Do	7686 (5%)	140	2.80
	Do	7686 (5%)	150	3.01
	Do	7686 (5%)	160	Weld
	D.O.S.	7686 (5%)	120	1.95
	D.O.S.	7686 (5%)	140	2.85
	D.O.S.	7686 (5%)	150	3.03
	D.O.S.	7686 (5%)	160	Weld
15	Extreme pressure tests on N-[9(10)-mercapto-9(10)-dibutylphosphato]stearoyl]morpholine (OC-O-No. 7687)			
	7687	None	120	0.608
	7687	None	160	1.45
	7687	None	200	1.40
20	7687	None	240	1.70
	7687	None	280	3.36
25	Extreme pressure tests on N,N-dibutyl-9(10)-carbobutyloxyoctadecanamide (PC-O-No. 7688)			
	7688	None	80	1.92
	7688	None	90	2.18
	7688	None	100	2.45
	7688	None	120	Weld

Extreme pressure tests on N,N-dibutyl-9(10)-carbobutyloxyoctadecanamide (PC-O-No. 7688)

7688	None	80	1.92
7688	None	90	2.18
7688	None	100	2.45
7688	None	120	Weld

1 Incipient seizure.
 2 5 seconds.
 3 4 seconds.
 4 10 seconds.
 5 46 seconds.
 6 Test stopped because of extreme decomposition. Did not weld.
 7 Test stopped because of extreme decomposition and noxious fumes. Did not weld.

TABLE IV

Evaluation of N-9,10-epithiostearoyl morpholine as an additive and as a base oil in extreme pressure tests (ASTM D 2596-67T)

40	Base oil	Percent additive	Load, kg.	Wear scar, mm.
	100 sec paraffin oil	(5%)	120	2.00
	Do	(5%)	140	2.80
	Do	(5%)	160	3.25
	Do	(5%)	180	Weld
45	Bis(2-ethylhexyl)sebacate	(5%)	80	1.57
	Do	(5%)	120	2.03
	Do	(5%)	140	2.41
	Do	(5%)	160	2.94
	Do	(5%)	180	3.00
	Do	(5%)	200	Weld
50	N-9,10-epithiostearoylmorpholine	None	120	0.70
	Do	None	200	2.13
	Do	None	240	2.43
	Do	None	300	2.68
	Do	None	380	Weld

1 10 seconds. 2 5 seconds. 3 50 seconds.

We claim:

1. The compound N - (ethoxyethoxyethoxy)propyl-9(10)-mercaptostearamide.

References Cited

UNITED STATES PATENTS

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65 ELBERT L. ROBERTS, Primary Examiner

U.S. Cl. X.R.

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(5/69)

UNITED STATES PATENT OFFICE
• CERTIFICATE OF CORRECTION

Patent No. 3,801,609 Dated April 2, 1974

Inventor(s) Frank C. Magne, Robert R. Mod, Gene Sumrell and Winfred E. Parker

It is certified that error appears in the above-identified patent and that said Letters Patent are hereby corrected as shown below:

Column 1, line 44, change "phosphorous" to - phosphorus -.

Column 3, line 17, change "N,N-dibutyloleamide" to - N,N-dibutylinoleamide

line 35, change "0.75 moles" to - 0.73 mole -

line 35, change "0.72 moles" to - 0.73 mole -

Column 4, line 53, change "10°" to - 10% -

Column 6, line 2, change "p.p.m." to - RPM -

line 4, change "ran" to - run -

Signed and sealed this 21st day of January 1975.

(SEAL)
Attest:

McCOY M. GIBSON JR.
Attesting Officer

C. MARSHALL DANN
Commissioner of Patents