## Synthesis and Preliminary Evaluations of Amide, Lactic Acid, and Terpenoid Derivatives of Substituted Phenoxycarboxylic Acids

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These investigations were undertaken for the purpose of utilizing materials of agricultural origin in the synthesis of conjugated biologically active compounds. All of the 32 new compounds prepared, amides, terpenoids, and lactic acid derivatives, induced plant-growth responses. In general, the magnitude of these responses was less than that induced by the parent regulating compounds. The data presented indicate that some derivatives may possess limited growth-regulating properties of interest inducing a localized response. In a few cases the coupling of compounds with halogenated phenoxy acids produced derivatives possessing increased growth-regulating properties.

THE SYNTHESIS and preliminary evalu-L ation of biologically active compounds have been reported in a series of papers (7-9, 13, 14). The broad purpose of the work is to create new and useful, synthetically prepared compounds from naturally occurring and derived source materials of agricultural origin. These investigations are motivated by the desire to obtain fundamental knowledge on the mode of action of growth regulators in plant metabolism and the effects of modification of chemical structure on the biological activity of plant growth substances. The immediate objective is the preparation of plant

<sup>1</sup> Present address, Courtaulds, Inc., Mobile, Ala. growth-regulating compounds possessing significantly different growth-regulating properties from those of parent substances holding prominence in the "growth regulator" field. Some of the desirable attributes sought are increased or decreased selectivity and translocatibility, modified phytotoxicity, and new growth-regulating properties.

This publication is a report on the preparation and some plant-regulating effects obtained with *p*-toluenesulfonamide (included because of chemical similarity to several naturally occurring amino acids), *p*-aminobenzoic acid, *p*aminosalicylic acid, and several lactic acid derivatives of halogenated phenoxycarboxylic acids; it also includes results of a preliminary evaluation of new derivatives of the terpenoids: nopol 1-[6,6-dimethylbicyclo-(1,1,3)-hept-2-en-2-yl]-ethan-2-ol (1), hydronopol, and Terposol No. 8 (chiefly terpinyl ethylene glycol ether).

The part reported here on amide derivatives is an extension of previous studies (7-9, 14) which have demonstrated that amino acid coupling can have a marked effect upon the growthregulating properties of a compound. This type of coupling has also resulted in an increase in the selectivity of the phenoxy-type growth regulator. Work on other amino acid, peptide, and protein hydrolyzate derivatives is in progress. Need for extended work covering a wide variety of terpenoids is indicated by the results obtained here and by others in the field of weed control (2, 4). New inexpensive terpenoids used as starting ma-

### Table I. Physical Properties, Yields, and Analytical Data for Halogenated Phenoxy Acyl Chlorides

		Boiling	Melting Reint					Analys	ses, %		
		° C.	° C.		Yield.	Car	bon	Hydro	ogen	Chic	rine
Name	Formula	(Uncor.)	(Uncor.)	$n_{\rm D}^{2.5}$	%	Caled.	Found	Caled.	Found	Caled.	Found
4-Chlorophenoxyacetyl chloride <sup>a</sup>	$\mathrm{C}_{\delta}\mathrm{H}_{6}\mathrm{Cl}_{2}\mathrm{O}_{2}$	92.0/1.0 mm.	22.0-24.0	1.5463	85.0	46.86	47.07	2,95	3.04	34.85	35.40
oxyacetyl chloride	$C_{\vartheta}H_8Cl_2O_2$	70.0/0.007 mm.	28.5-30.0	1.5400%	93.5	49.34	50.04	3.68	4.17	32.37	32.56
acetyl chloride <sup>c</sup> 2,4,5-Trichlorophenoxy	$\mathrm{C_8H_5Cl_3O_2}$	102-105/0.3 mm.	50.3-52.3 <sup>d</sup>	1.5622	87,8	40.13	40.22	2.11	1.90	44.42	44.30
acetyl chloride <sup>e</sup> 2-(2,4-Dichlorophen- oxy)propionyl chlo-	$C_8H_4Cl_4O_2$	128–130/2.0 mm.	84.5-85.0 <i>d</i>	1.5590/	74.2	35.08	35.25	1.47	1.81	51.76	51.68
ride <sup>g</sup> 2-(2,4,5-Trichlorophen-	$\mathrm{C_9H_7Cl_3O_2}$	76.0/0.1 mm.	<sup>h</sup>	1.5440	90.4	42.59	43.25	2.74	3.23	41.96	41.97
ride 4-(2.4-Dichlorophen-	$C_{\vartheta}H_6Cl_4O_2$	111-117/0.3 mm.	<sup>h</sup>	1.5601 <i>i</i>	73.0	37.54	37.73	2.10	2.23	49.25	48.96
oxy)butyryl chloride 4-(2-Methyl-4-chloro- phenoxy)butyryl	$\mathrm{C}_{10}\mathrm{H}_9\mathrm{Cl}_3\mathrm{O}_2$	136-140/0.2 mm.	17.0-22.0	1.5465	89.1	44,89	44.93	3,39	3.38	39.76	39.43
chloride	$\mathrm{C_{11}H_{12}Cl_2O_2}$	115-122/0.2 mm.	5.0-9.0	1.5340	73.5	53.46	53.45	4.90	5.19	28,70	28.62
<sup>a</sup> B.p. 142°/17 mm., <sup>b</sup> At 32.8° C. <sup>c</sup> B.p. 155–157°/22 m <sup>d</sup> Cor. <sup>e</sup> B.p. 165–167°/6 mm	m.p. 18.8, yiel nm., yield 84–9 n., m.p. 78–79	ld 90% ( <i>10</i> ). 90% ( <i>14</i> ). 9°, yield 80% ( <i>6</i> ).		<sup>f</sup> At 94° <sup>g</sup> B.p. 13 <sup>h</sup> Liquid <sup>i</sup> At 26.	°C. 37–139° at 5° 0°C.	°/9 mm. C.	$n_{\rm D}^{20} =$	1.5475, y	ield 84 $\%$	(5).	

### Table II. Physical Properties, Yields, and Analytical Data for Amide Derivatives of Halogenated Phenoxy Acids

		14 P		Yie	ld			Analy	ses, %	
		°C.ª	Cru	de	Refi	ned	Chlo	rine	Nitr	ogen
Name	Formula	(Cor.)	G	%	G	%	Caled.	Found	Calcd.	Found
<i>p</i> -Toluenesulfonamide										
N-(4-chlorophenoxyacetyl)-	C <sub>15</sub> H <sub>14</sub> ClNO <sub>4</sub> S	169-171	4.37	64.2	1.72	25.3	10.43	10.54	4.12	4.02
N-(2-methyl-4-chlorophenoxyacetyl)-	C <sub>16</sub> H <sub>16</sub> ClNO <sub>4</sub> S	204-206	5.34	75.3	2.20	28.5	10.02	9.93	3.96	3.96
N-(2,4-dichlorophenoxyacetyl)-	$C_{15}H_{13}Cl_2NO_4S$	170-171	3.96	52.7	2.75	36.6	18,96	18.99	3.75	3.75
N-(2,4,5-trichlorophenoxyacetyl)-	$C_{15}H_{12}Cl_3NO_4S$	177-179	6.48	79.4	3.35	41.0	26.03	25.74	3.43	3.45
N-[DL-2-(2,4-dichlorophenoxy)propionyl]-	$C_{16}H_{15}Cl_2NO_4S$	123-125	4.78	63.8	1.35	18.0	18.29	18.12	3.61	3.51
Aminobenzoic acid										
N-(4-chlorophenoxyacetyl)-	$C_{15}H_{12}ClNO_4$	237-239	4.10	66.9	3.85	62.8	11.59	11.92	4.58	4.63
N-(2-methyl-4-chlorophenoxyacetyl)-	$C_{16}H_{14}ClNO_4$	260–263 <sup>b</sup>	3.30	51.6	2.78	43.5	11.09	11.17	4.38	4.39
N-(2,4-dichlorophenoxyacetyl)-	$C_{15}H_{11}Cl_2NO_4$	272-275	3.02	83.6	2.30	64.1	20.55	20.70	4.11	4.13
N-(2,4,5-trichlorophenoxyacetyl)-	$C_{15}H_{10}Cl_3NO_4$	310-315°			1.13ª	15.1	28.39	27.99	3.74	3.72
N-[DL-2-(2,4-dichlorophenoxy)propionyl]-	$C_{16}H_{13}Cl_2NO_4$	208-2095	14.18	100.0	6.13	44.5	20.02	20.02	3.96	3.99
<i>p</i> -Aminosalicylic acid										
N-(4-chlorophenoxyacetyl)-	$C_{15}H_{12}CINO_5$	225-227	4.32	67.0	1.95	30.2	11.02	11.12	4.37	4.36
N-(2-methyl-4-chlorophenoxyacetyl)-	$C_{16}H_{14}CINO_{5}$	223–227¢			1.68	25.0	10.54	10.56	4.21	4.17
N-(2,4-dichlorophenoxyacetyl)-	$C_{15}H_{11}Cl_2NO_5$	256-258	4.92	68.9	1.55	21.7	19.90	19.40	3.92	3.84
N-(2,4,5-trichlorophenoxyacetyl)-	C15H10Cl3NO5	281–284 <sup>b</sup>			2.07	26.5	27.23	27.02	3.59	3.54
N-[pl-2-(2,4-dichlorophenoxy)propiony]]-	$C_{16}H_{13}Cl_2NO_5$	215-221	3.81	51.4	2.91	39.3	19,16	19.07	3.78	3.79

<sup>a</sup> Recrystallized once or more from ethyl alcohol water and ethyl acetate-petroleum ether unless otherwise indicated.

<sup>b</sup> Recrystallized from ethyl alcohol water only,

Recrystallized from ethyl alcohol water and ethylene glycol monoethyl ether.

<sup>d</sup> A portion of sample lost by accident.

e Recrystallized twice from methanol water.

# Table III. Physical Properties, Yields, and Analytical Data for Lactic Acid Derivatives of Halogenated Phenoxy Acids

		Boiling Banae			Yield,	%			Analys	es, %		
		° Č.				Re-	Car	bon	Hydr	ogen	Chic	orine
Name	Formula	(Uncor.)	<b>n</b> $^{25}_{ m D}$	d425	Crude	fined	Caled.	Found	Caled.	Found	Caled.	Found
DL-2-[DL-2-(2,4,5-tri- chlorophenoxy)pro- pionoxy] propionic acid methyl ester <sup>a</sup> DL-2-[DL-2-(2,4,5-tri- chlorophenoxy) pro-	$C_{13}H_{13}Cl_3O_{\delta}$	170-90/0.2mm.	1.5232	1.3649	78	66	43.91	44.41	3,68	3.82	29.91	29.59
pionoxy] propionic acid <i>n</i> -butyl ester DL-2-(2-methyl-4- chlorophenoxy ace-	$\mathrm{C}_{16}H_{19}\mathrm{Cl}_{3}\mathrm{O}_{5}$	187–93/0.2 mm.	1.5094	1.2679		75	48.32	48.53	4.82	4.92	26.75	26.30
toxy)propionic acid methyl ester DL-2-(2-methyl-4- chlorophenoxy ace-	$C_{13}H_{1\delta}ClO_{\delta}$	167–75/0.2 mm.	1.5077	1.2340		65	54.46	54.38	5.27	5.16	12.37	12.69
toxy)propionic acid n-butyl ester	$\mathrm{C}_{16}\mathrm{H}_{21}\mathrm{ClO}_{5}$	190-205/0.2 mm.	1.4954	1.1520	69	53	58.45	58.00	6.44 Nite	6.05	10.78	10.99
DL-2-(2,4-dichloro- phenoxyacetoxy)pro pionic acid <i>N</i> -ethyl- amide	- C <sub>13</sub> H <sub>15</sub> Cl <sub>2</sub> NO <sub>4</sub>	108.5–9.5 (m.p. cor.)			21	12			4.37	4.47	22.15	22.18
<sup>a</sup> Solidified to soft w	axy crystals.											

terials are being supplied by the Naval Stores Section of the Agricultural Research Service's Southern Regional Research Laboratory. The chemistry on the preparation and purification of the terpenoid derivatives used in the present experiments has been reserved for publication with extended studies now in progress on volatility and herbicidal properties.

The development of "lactoprene" rubbers at the Eastern Regional Research Laboratory made available a wide variety of lactic acid derivatives for synthetic purposes. Attempts in 1953 to couple lactic acid and some of its derivatives with halogenated phenoxycarbox-

ylic acids were unsuccessful. However, a single compound, DL-2-(2,4-dichlorophenoxyacetoxy)propionic acid N-ethylamide, combining DL-N-ethyllactamide with 2,4-dichlorophenoxyacetic acid, was made and preliminary tests showed that it possessed growth-regulating properties; also it was quick-acting on the test plants used. A discovery made about this time (11) that leaves of test plants (corn was an exception) were able to translocate plant regulating carbamates more readily when the compounds contain the lactic acid group kept alive the desire to incorporate lactic acid into the phenoxy type regulator. Four more lactic acid derivatives have now been made and a preliminary evaluation of all five compounds is reported here.

#### Experimental

Compounds used for synthetic purposes were the best obtainable from commercial sources, utilized without further purification. Where new derivatives prepared were solids. no attempts were made to improve yields by working up mother liquors.

For the preparation of the amide derivatives of halogenated phenoxycarboxylic acids obtained by coupling ptoluenesulfonamide,p-aminosalicylic acid, and p-aminobenzoic acid Schotten-Bau-

mann techniques were used (7-9). When it was necessary, halogenated phenoxycarboxylic acids were first converted to their acyl chlorides by the use of thionyl chloride employing the method of Freed (3). Informative data concerning the properties of all the chlorides used are summarized in Table I. Values obtained by other investigators are also presented. Data on 4-(2,4-dichlorophenoxy)butyryl and 4-(2-methyl-4chlorophenoxy)butyryl chlorides are included, because these compounds have been used to prepare amino acid derivatives which are being evaluated as herbicides.

The preparation of 2,4,5-trichlorophenoxyacetyl-p-toluenesulfonamide is illustrative of that used for the other sulfonamides and for the derivatives of p-aminobenzoic and p-aminosalicylic acids. Melting points, yields, and analytical data for these compounds are recorded in Table II.

2,4,5-Trichlorophenoxyacetyl-p-toluenesulfonamide. To p-toluenesulfonamide (3.42 grams, 0.02M) dissolved in 60 ml. of 1N sodium hydroxide chilled to 5° C, was added, dropwise with continuous stirring, a cold solution containing 2,4,5-trichlorophenoxyacetyl chloride (5.46 grams, 0.02M) dissolved in 30 ml. of benzene. The reaction mixture was stirred for 3 hours as it warmed to room temperature. Three 20-ml. portions of ether were used for extractions made in a separatory funnel. The combined ether extracts were washed with 20 ml. of distilled water. The washing was returned to the alkaline solution, which was then acidified with 1N hydrochloric acid using Congo red paper as an indicator. The acidified solution was placed in the refrigerator overnight. The white crystalline product was filtered off, slurried three times with water, and finally dried in a vacuum desiccator by continuous overnight pumping. The product was ground and washed three times with small portions of warm petroleum ether (boiling range  $63^{\circ}$  to  $70^{\circ}$  C.). The crude yield was 6.48 grams (79.4%), melting point 165-70° C. Following two recrystallizations by dissolving in hot ethyl acetate and precipitating with petroleum ether the melting point was 174-9° C.

A third recrystallization performed by dissolving in hot ethyl alcohol and precipitating with water produced a product with a satisfactory melting point, 177 to 179° C. The final yield was 3.35 grams (41.0%) (Table II).

The lactic acid derivatives were prepared by two varying techniques. First the n-ethylamide of DL-2-(2,4-dichlorophenoxyacetoxy) propionic acid was made by the azeotropic technique.

DL - 2-(2,4-Dichlorophenoxyacetoxy) propionic Acid N-Ethylamide. DL-N-Ethyllactamide (12) (78.0 grams,

120

Table IV. Plant Growth-Regulating Activ	rity of An	nide,	, Laci Aov	ic Ac er (S	, Cu	nd To cumb	erpen er (C	), Ba	erivo rley (	ıtives B), α	nd Co	alog rrn ((	enate (n)°	Чdр	enox	y Aci	ids o	n Bla	х Х	alent	ine B	ean	(SB),	Sun	
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Compounds Tested	VB	S	U	٧B	s	υ	VB VB	s	U	٧B	S	U	VB	s	υ I	<b>V</b> 8	s	ΰ	٧B	s	U	•	ວົ	B	E,5
Parent acids 4-Chlorophenoxyacetic 2-Methyl-4-chlorophenoxyacetic 2,4-Dichlorophenoxyacetic 2,4,5-Trichlorophenoxyacetic	0 <del>-</del> 2 <del>-</del> 0	$5^{p}$ $\frac{1}{2}$ $3^{p}$ $0^{p}$	ตั <i>ต</i> ์ ดัง ดั	ભ જેં છ જે	ົດ ຕິດ ຕັ	ัก ทั้ ก ทั้	0000	5° - 1° 3° 0°	ั <i>สิ</i> ที่ ค ร้	3 0 0 <del>1</del>	3565	3503	`n 0 n 5	0000	0000	3 <b>*</b> 3 °	10°0°	రింరి	ົກ ຕົ້ ຕ ຕົ້	÷ô30	0-100	0101	0 ~ ~ ~ ~	<i>w00w</i>	0-0-
DL-2-(2,4-Dichlorophenoxy) propionic DL-2-(2,4,5-Trichlorophenoxy) propionic	3° 2°	5° 1	3° Ô	Å Å	3° 1°	ñ ĥ	0°0	°0≁	Å Ô	00	10	15 IS	ŝĉ	ô ô	င်စိ	3° 0°	16	စ်စ်	ñ %	16	60	7	-	ŝ	<del></del>
<ul> <li><i>b</i>-Tolucnesulfonamides</li> <li><i>N</i>-(4-chlorophenoxyacetyl)-</li> <li><i>N</i>-(2,methyl-4-chlorophenoxyacetyl)-</li> <li><i>N</i>-(2,4,5-trichlorophenoxyacetyl)-</li> <li><i>N</i>-(2,4,5-trichlorophenoxyacetyl)-</li> <li><i>N</i>-[DL-2-(2,4-dichlorophenoxy) propionyl]-</li> </ul>	-000-	~~~~ <sup>0</sup>	~~~~	<i>~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~</i>	(1) (2) (2) (3) (3) (4) (5) (6) (6) (7)	\$\$\$\$\$\$	00000	0 $0$ $0$ $0$ $3$	000-0	0,-12,-3,	0 - 5 - 5	ŝ−77-3	$\omega \odot \omega \omega \omega$	00000	00000	00,000	$-\alpha \phi - \alpha$	- n 0 n n	<i>იი იი იი ი</i> ი	$-\frac{1}{2}$	n n n n n n	∞00−−	000	00000	00
<i>p</i> -Aminobenzoic acids <i>N</i> -(4-chlorophenoxyacctyl)- <i>N</i> -(2-methyl-4-chlorophenoxyacctyl)- <i>N</i> -(2,4-dichlorophenoxyacctyl)- <i>N</i> -(2,4,5-trichlorophenoxyacctyl)- <i>N</i> -[DL-2-(2,4-dichlorophenoxy) propionyl]-	-0000	-000-	01 3 3	00503	00707	0-7075	00000	00000	0-000	11600	00-	00100	00003	00000	00000	-0000		03000	005-3	0 0 3 1 5	00000	-00-10	00	-0888	07107
<i>P</i> -Aminosalicylic acids <i>N</i> -(4-chlorophenoxyacctyl)- <i>N</i> -(2-methyl-4-chlorophenoxyacctyl)- <i>N</i> -(2,4,5-trichlorophenoxyacctyl)- <i>N</i> -(2,4,5-trichlorophenoxyacctyl)- <i>N</i> -[DL-2-(2,4-dichlorophenoxy) propionyl]-	00	11210	∞ ∞ ∞ ∞ −	0 0 10 0 0	~~~~~		00000	00000	00-00	0°0000	0535	<b>ဝ</b> ဝက်က်က်	~ 0 0 0 0	<i>кк</i> -00	00000	00000	00000	- ~ 0	с с с с с с с с с с с с с с с с с с с	66933		00	00		01101

Lactic acid derivatives n2-[2-n(2,4,5-trichlorophenoxy) propionoxy  - monionic acid, methyl seter	~	<del>.</del>	ĸ			-	<del>.</del>	,	άÇ	<del>.</del>	-	ç	c	-	ç	-	c	ç	<del>.</del>	c	с с	<del>, -</del>	-
DL-2-(2-4,5-tricki) and DL-2-(2-4,5-tricki) and DL-2-(2-01-(2,4,5-tricki)) propionoxy ]- propionic acid, <i>n</i> -butyl ester	n 0	- 0	, <del></del>		, ,	> 0	- 0	— ر	0	- 7	0	4 0	• •	0	2 V	- 0		1 m	- 0	> —		• •	~ ~
DL-2-(2-methyl-4-chlorophenoxyacetoxy) propionic acid, methyl ester	3	3	$2^{h}$			0 4	3	34	34	7	2¢	-	0	90	5	0	$0^{p}$	3	c	<b>0</b>	0	-	2
DI-2-(Z-methyl-4-chlorrophenoxyacctoxy) propionic acid, n-butyl ester DL-2-(2,4-dichlorophenoxyacctoxy) propionic acid	3	3	3	ŝ	-	0	3	3	φŊ	2	2	0	0	0	<del></del>	-	-	2	-	-	0	2	2
N-ethylamide	2	3	3	~	-	0	0	3	0	-	2	0	c	2	2		<del>.                                    </del>	3	1	-	3	-	0
Parent terpenoid compounds Hydronopol Nopol	00	00	00	-0		00	00	00	00	00	00	00	00	00	00	00	00	00	00	00	00	00	Ċ O
Hydronopyl- 4-chlorophenoxyacetate 2-mcthyl-4-chlorophenoxyacctate 2,4-dichlorophenoxyacctate 2,4,5-trichlorophenoxyacctate	00%+0	00000	(i)	005		-0000	00000	00000	~0000	00000	ຈິ <i>ເ</i> ຈີ ທີ່ ທີ່	00000	00000	66666	00-00	000-0	00000	0000-	00000	00000	00000	1 0 0 1	000
Nopyl- 4-chlorophenoxyacetate		0	n m	- ~			0	• •	3	0 °0	ŝ î	0	o °o	- °0	• •	• •	0	- ~~					- 4
2-methyl-4-chlorophenoxyacetate 2,4-dichlorophenoxyacetate	00	• •	ς Ω	- 0		00	0 O	00		00	ÅÅ	0 0	రీరే	°0	00	c	00	2 5	00	<u> </u>	 	70	0 0
2,4,5-trichlorophenoxyacetate DL-2-(2,4,dichlorophenoxy) propionate	- 0	00	ς Ω	~~	01.01	00	¢ ¢	00	00	0 0	3¢ 3¢	- 0	°0°0	6°0	- 0	с о	00	77	00	00	0 0	- 0	- 0
DL-2-(2,4,5-trichlorophenoxy) propionate 2-(a-Terpinyloxyethyl)-p-chlorophenoxyacctate	<i></i> со	00	ŝ	20		00	00	00	0 0	00	3; 3;	ς Ω	ôô	ÓÔ	77	00	00	ς	00	00	00		0 0
<sup>a</sup> Responses observed on 14th day after treatment. 0, <sup>b</sup> Evaluated at 0.1 concentration (equivalent to about <sup>c</sup> Evaluated at 0.01 concentration.	no effec 15γof	t; 1, s test co	light cl mpour	fect; : id per	2, moc plant;	lerate e initia	sffect; 1 conce	3, ma entrati	rked el on, 15	ffect. 0γpen	. plant	, see (	9) for	metho	ds).								

0.67M) and 2,4-dichlorophenoxyacetic acid (148 grams, 0.67M) were mixed in a flask to which were added 400 ml. of xylene and 2 grams of sulfuric acid. This reaction mixture was refluxed until no more water could be collected in a moisture tube (16 hours). The xylene was removed under reduced pressure yielding a residual heavy oil. Attempts to recrystallize this oil from ethyl acetate and ethyl alcohol were unsuccessful.

The product was effectively extracted batchwise from the oil by hot petroleum ether (boiling point  $35^{\circ}$  to  $59^{\circ}$  C.) from which it crystallized on cooling. The crude yield of combined extracts after one recrystallization from petroleum ether was 44.8 grams (20.9%). Two recrystallizations from a mixture of hot ethyl acetate and petroleum ether (boiling point  $63^{\circ}$  to  $70^{\circ}$  C.) gave 26.0 grams (yield 12.1%) of purified product melting at 108.5– $9.5^{\circ}$  C. (Table III).

The other four lactic acid derivatives, DL - 2 - (2 - methyl - 4 - chlorophenoxy)acetoxy) propionic acid methyl and *n*butyl esters, and DL-2-[DL-2-(2,4,5)trichlorophenoxy) propionoxy] propionic acid methyl and *n*-butyl esters, were prepared by a pyridine catalyzed esterification of the hydroxyl group of the appropriate lactic acid esters. The following example illustrates the procedure.

DL - 2 - [DL - 2 - (2,4,5 - Trichlorophenoxy)propionoxy]propionic Acid n-Butyl Ester. DL-Butyl lactate (52.6 grams, 0.36M) was diluted to 500 ml. with a 6 to 1 mixture of petroleum ether (boiling point 63° to 70° C.) and benzene, pyridine (34 ml., 0.42M) was added, and the mixture was stirred in an ice bath. The DL-2-(2,4,5-trichlorophenoxy)propionvl chloride (86.4 grams. 0.30M), diluted with petroleum etherbenzene mixture, was added dropwise over a 10-minute period. A rapid separation of pyridine hydrochloride indicated an immediate reaction in formation of the product. After 2 hours, the cold reaction mixture was allowed to warm to room temperature. The bulk of the pyridine hydrochloride was removed by filtration and the petroleum ether-benzene solution washed with water, dilute hydrochloric acid, aqueous sodium carbonate, methanol (50%), and water. Following two stages of adsorption treatment with a mixture of sodium sulfate, carbon, diatomaceous earth, and Florisil, the product was vacuum distilled at 187-93° C. at 0.2 mm. of mercury with a constant index of refraction  $(n_{D}^{21})$ = 1.5094). The specific gravity of the product was 1.2679 at 25° C. and the yield was 75%.

Data on the physical properties, yields, and analyses for all the lactic acid derivatives are recorded in Table III.

Methods Used in Plant Tests. The parent and derivatized halogenated phenoxy acids were evaluated for plant

growth regulating activity using the dicotyledonous plants, Pinto bean, large seeded sunflower, Arlington White Spine cucumber, and the monocotyledonous plants Wong barley, and U. S.-13 hybrid corn. The lanolin assay method was used on dicots and the coated sand assay method on monocots (9).

At intervals of 2, 4, 7, and 14 days following treatment, the degree of growth modification induced by the various compounds was estimated and scored according to the intensity of growth responses. Responses studied were: stem curvature, growth inhibition, epinasty, formative effects, and induced cell proliferation (gall formation). Table IV shows responses observed 14 days after treatment, to conserve space the other values have been omitted. These data are representative, although they do not show the rate of response or relative progressive effectiveness of the compounds tested. The experiments were designed to indicate growth regulating properties rather than the herbicidal potentialities of the compounds.

Results with Plant Tests. In general, coupling of halogenated phenoxy acids with p-toluenesulfonamide, p-aminobenzoic acid, and p-aminosalicylic acid and terpenoids resulted in compounds that induced plant-growth responses, but the magnitude of the different responses was less than that induced by the parent regulating compounds (Table IV). From the standpoint of utilization of these new regulating compounds, the parent phenoxy acids sometimes induce unwanted responses when used for a specific purpose-for example, to improve fruit set. The present data indicate that some derivatives used here may possess limited growth regulating properties of interest. For example, it is desirable to induce fruit set chemically without marked side effects such as modification and suppression of leaf growth. Regulators that induce a localized response are of interest. In the present tests, 2,4,5-trichlorophenoxyacetic acid induced marked responses (formative effects) at a distance from the site of application when applied to bean plants. The hydronopol derivative of this acid, on the other hand, induced a moderate response which was apparently localized within the treated region of the stem.

In a few cases, coupling of the phenoxy acids resulted in compounds with increased growth-regulating properties. When applied to cucumbers, for example, the lactic acid derivative, DL-2-(2 - methyl - 4 - chlorophenoxyacetoxy) propionic acid, methyl ester was more effective than its parent acid, 2-methyl-4-chlorophenoxyacetic acid.

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### FRUIT DROP CONTROL

# **Evaluation of Chlorine-Substituted Phenoxyacetic Acids and Amides for Retarding Abscission of Apple Fruits**

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The amide forms of chlorine-substituted phenoxyacetic plant regulators were more effective in retarding mature fruit abscission than the corresponding acids. When chlorine was substituted in all possible combinations on mono-, di-, tri-, tetra-, and penta- positions on the benzene nucleus, in all instances where the 6 position contained chlorine the compound was rated as relatively inactive. To a lesser degree chlorine substitution in the 3 position also reduced the effectiveness of a compound. The 17 acids and 17 amides were compared directly on three varieties of mature apple fruits; the complete series of 17 amides was applied to six varieties.

PLANT-REGULATING ACTIVITIES of chemicals can be compared or evaluated in various ways. Retardation of the abscission of mature fruits of apple was used as the response in the present study, because that type of plant reaction appears to be very different from the cell elongation or multiplication, formative

growth effects, parthenocarpy, modification of organs, and repression or stimulation of root or vegetative bud meristems or others (11, 15) commonly used as measures of response.

According to McCown (4) the harvest drop of mature apple fruits involves dissolution changes in the walls and

lamellae of old, lignified cells. The possible role of plant-regulating chemicals in controlling the metabolism of pectic substances through enzymatic processes has been suggested by Neeley et al. (10). Other workers have shown that chemical changes associated with the ripening of fruits can be stimulated or