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OBSERVATIONS ON THE USE OF "PHENOL" LARVICIDES FOR MOSQUITO CONTROL

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For many years commercial larvicides, containing cresylic acid in sulfonated oil, and generally referred to as "phenol" or "phenolic" larvicides, have been reported to be effective in antimosquito work in heavily polluted waters, stagnant pools under buildings, and similar places where oil larvicides in usual quantity have been unsatisfactory. Within the past 3 or 4 years, commercial larvicides of this type have been used in some areas for general control of mosquito larvae. For this work, a compound having a phenol coefficient of 10 to 14 is diluted with water in the ratio of 1 to 30 and applied to the water surface as a fine spray; it is claimed that if the spray is sufficiently fine the larvicide will remain on the surface of the water sufficiently long to kill mosquito larvae. As these commercial larvicides may be diluted with any available water in the field, their use in mosquito control would obviate the transportation of large quantities of material from the headquarters, if effective at the dilution mentioned, as only a small volume of the larvicide need be transported to the location of the water supply.

A number of tests, in both the laboratory and the field, undertaken to demonstrate the value of these larvicides, are described in this report.

EFFICIENCY IN KILLING MOSQUITO LARVAE

Laboratory tests.—The phenol larvicide diluted 1 to 30 with water was tested in the laboratory by applying it in a fine spray from an atomizer to the surface of water in enamel pans in which mosquito larvae had been placed. The rate of application of the larvicide was determined by using exact quantities measured in graduated cylinders

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and expressed as gallons per acre; the number of mosquito larvae was determined by actual count. In testing the phenol larvicide, two types of controls were used: (1) larvae in pans similar to those treated with larvicide, allowed to remain completely untreated, and (2) larvae in similar pans treated with a kerosene spray.

TABLE 1.—Percentage mortality of Anopheles quadrimaculatus 4th stage larvae, untreated and treated with kerosene and various amounts of phenol larvicide. Larvicide applied to pans, each pan containing 30 A. quadrimaculatus 4th stage larvae in tap water

		Per	cent mortality		
Time after treatment	Ph	enol larvicide l	1-30	Kerosene	
	10 gal. per acre, dilution 1–315,000	15 gal. per acre, dilution 1–141,000	25 gal. per acre, dilution 1–156,000	10 gal. per acre	Untreated
1 hour	Nil 7	3 10	20 . 47	90 100	Nil Nil

The results of these tests are shown in tables 1 and 2. It appears from table 1 that phenol larvicide (1 to 30) applied at the rate of 25 gallons of the diluted spray per acre killed only 47 percent of anopheline larvae, whereas kerosene applied at the rate of only 10 gallons per acre killed 100 percent of larvae. In table 2 it is shown that phenol larvicide (1 to 30) applied at the rate of 25 gallons per acre killed only 7 percent of culicine larvae, as compared with 99 percent in the case of kerosene. Even when the phenol larvicide (1 to 30) was applied at the rate of 100 gallons per acre, only 42 percent mortality resulted.

TABLE 2.—Percentage mortality of Culex quinquefasciatus 3d and 4th stage larvae, untreated and treated with kerosene and various amounts of phenol larvicide. Larvicide applied to pans, each pan containing 20 Culex quinquefasciatus 3d and 4th stage larvae in pond water

		Per	cent mortality		
Time after treatment	Ph	enol larvicide l	1-30	Kerosene	
	25 gal. per acre, dilution 1–60,000	50 gal. per acre, dilution 1–30,000	100 gal. per acre, dilution 1–15,000	25 gal. per acre	Untreated
1 hour	47	11 16	36 42	95 99	Nil Nil

Field tests.—The unusual drought current at the time of the study (May 13-June 6, 1941) limited the selection of areas suitable for testing larvicides.

The general procedure in all field tests was first to sample the area by dipping with a one-pint dipper. The kind and number of larvae

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for each dip were recorded and the larvae carefully replaced. The measured area was then treated by spraying with a measured amount of larvicide with an ordinary orchard sprayer. The nozzle was so adjusted that the spray was as fine as could be projected a distance of 6 feet.

Approximately 2 hours after treatment the area was again dipped and the kind and number of larvae recorded for each dip. The dipping determinations are shown in table 3 as the average number of larvae per dip, together with the number of dips.

						Phe	nol la	arvic	ide 1-3	0				:	Kero	sene			
		Nu ber	m- of	per	luare	Ano lir	phe- tes	C	'uli- ines	Т	otal	per	uare	Ano lir	phe- nes	Cı cir	ıli- nes	To	tal
Number of test	Type of area	Before Hreatment	After 65	Application rate, gal. acre	Size of area treated in sq	Average number pcr dip before treatment	Percent mortality	Average number per dip before treatment	Percent mortality	Average number per dip before treatment	Percent mortality	Application rate, gal. acre	Size of area treated in sq feet	A versge number per dip before treatment	Percent mortality	Average number per dip before treatment	Percent mortality	Average number per dip before treatment	Percent mortality
1 1 2 3 4	Ditch Pond Pond Lagoon	60 54 50	60 54 50	15 35 95 20	1, 500 400 150 540	3. 30 0. 96 6. 20	28 65 25	2.92 .24 4.50	60 62 \$25-50 21	6. 22 1. 20 10. 7	43 65 225-50 22	15 35 20	1, 500 150 540	*1.0 8.14	88 	³ 1.3 6.66	68 3 98 59	³ 2.3 14.8	77 9 98 61

 TABLE 3.—Comparative action of phenol larvicide and kerosene on mosquito larvae

 2 hours after treatment in the field

¹ Ditch was clear of vegetation and covered with a heavy scum.

³ Estimate. ³ Only 50 dips.

Effects on vegetation and fish were noted at approximately 2 hours and 24 hours after treatment, respectively.

The area used in field test No. 4 was covered with a heavy mat of water milfoil (*Myriophyllum*). The results for the phenol and kerosene in this area are comparable and show the relative toxicity of the two larvicides, phenol 22 percent mortality and kerosene 61 percent.

Field test No. 3 was performed on a small "pond" of about 150 square feet in area and clear of all vegetation and flotage. No attempt was made to count the larvae, *Culex quinquefasciatus*, in this area because of the large number present. An estimate of the number of larvae present after treatment was only approximate. However, after treatment with the phenol larvicide (1 to 30) the live larvae present were still too numerous to count. Twenty hours after the application of the phenol larvicide and with innumerable larvae still present, kerosene was applied. An hour after the application of kerosene, several portions of the water area were carefully searched for live larvae. These observations form the basis for our estimate of a 98 percent kill for kerosene. The area in field test No. 2 was another pond which contained a number of minnows, but very few larvae. The choice of this pond was dictated by the presence of minnows; the number of larvae present was so small and the resultant errors of sampling so large that the value given for the percentage kill, 65 percent, is also subject to a large error.

Field test No. 1 was conducted on a small bayou from 3 to 12 feet wide. The area in which the phenol larvicide was applied was clear of vegetation and flotage, while the area used for kerosene was clear of vegetation but was covered with a heavy scum. The percentage killed, 77 percent for kerosene, in this area would probably have been greater if the water surface had been cleared of this scum.

OTHER EFFECTS OF PHENOL LARVICIDES

Action on fish.—Both laboratory and field observations revealed that the phenol larvicide tested had a harmful effect on fish. In table 4 it is shown that the larvicide applied at the rate of 25 gallons per acre to aquarium tanks killed 29 percent of goldfish and 25 percent of "shiners." This same rate of application gave only 47 percent kill of anopheline larvae and 7 percent of culicine larvae (tables 1 and 2, respectively) under laboratory conditions. When the larvicide was applied at the rate of 50 gallons per acre, a comparison of the percentage mortality for fish and for larvae is more striking—16 percent mortality for the culicine larvae as compared to 100 percent mortality for the goldfish.

		·	Percent m	ortality 24 h	ours after tre	atment	
	Number of fish		Phenol lar	vicide 1-30		Kerosene	
-Type of fish	used in each test	10 gal. per acre, dilution 1-800,000	15 gal. per acre, dilution 1–535,000	25 gal. per acre, dilution 1–320,000	50 gal. per acre, dilution 1–160,000	25 gal. per acre	Untreated
Goldfish Shiner minnows	14 12	Nil 33	16	29 25	100	Nil Nil	Nil Nil

 TABLE 4.—Percentage mortality of fish in aquaria, untreated and treated with kerosene and various amounts of phenol larvicide

Also, evidence of damage to fish was obtained in the field. In table 3, test No. 2, it is noted that application of the phenol larvicide at the rate of 35 gallons per acre killed 65 percent of total larvae. When this test was conducted, 10 minnows were recovered from the water after treatment by means of a hand dipper; 4 of these were dead, and the degree to which the other 6 were adversely affected by the larvicide was evident from the ease with which they were picked up in the dipper. In test No. 3, table 3, larvicide was applied at the rate of 95 gallons to the acre, giving a kill estimated at 25-50 percent of larvae at the end of an hour. Larvae in this pond were far too numerous to count. Two small catfish, each about 4 inches long, were found dead when the pool was inspected 20 hours after application of the larvicide.

Action on vegetation.—No studies designed to determine the effect of the larvicide on vegetation, either gross or microscopic, were undertaken. However, observations made while studying the effects on larvae and fish indicated that slight burning of leaves of a number of plants occurred following the application of the phenol larvicide. Such action has also been reported following the use of kerosene, although in both cases recovery of the affected vegetation appears to occur rapidly.

SUMMARY AND CONCLUSIONS

(1) Under the conditions of these experiments, phenol larvicide (diluted 1 to 30) applied at rates varying from 10 to 95 gallons per acre was less effective than kerosene.

(2) Phenol larvicide, as tested in these studies, was harmful to fish. In the laboratory, the larvicide applied at the rate of 50 gallons per acre killed 100 percent of fish but only 16 percent of larvae.

(3) The phenol larvicide as used in this study, because of its low toxicity for larvae and detrimental effect on fish, does not appear to be a desirable larvicide for general mosquito control.

THE DEPOSITION AND REMOVAL OF LEAD IN THE SOFT TISSUES (LIVER, KIDNEYS, AND SPLEEN)¹

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When lead is absorbed into the body there occurs a general flooding of all the tissues by the lead stream, which the system as a whole tends to correct either by increased excretion or by deposition of the lead in the bone tissue. The removal of lead by the latter method is a more or less temporary means of immobilizing the lead, but it continues, as Minot (1, 2, 3) has shown, after absorption has ceased, the lead being drawn from the softer tissues. Minot found that in animals killed several months after the last dose of lead by mouth the skeletal lead amounted to 97.0 to 98.5 percent of the total lead in the body.

In spite of the known transitory nature of the lead in the softer tissues, a great deal of data has been accumulated giving the concentration or amounts of lead in these tissues. In these cases information is usually lacking with respect to the time when exposure or

¹ From the Division of Industrial Hygiene, National Institute of Health.

absorption has ceased, except in the case of human autopsy material. This point is important if the significance of the amount of lead in the softer tissues is to be considered in relation to lead poisoning. In recent experiments in this laboratory it appeared that the interval of changing lead content of the soft tissues may not only be a matter of months or even weeks after the ingestion of lead has ceased, but it may be a matter only of days.

Frequently the metal content of a given tissue is related to its pathology and because of the probable variable shift in amount of the deposited material within a relatively short time the following experiments were undertaken to verify this and to find out the extent to which this shift occurred with lead during a reasonable interval.

As a change in the soft tissue content of lead had been noted in similar groups of animals killed within less than a week after lead ingestion had ceased, it was felt that a 14-day rest period would be sufficient to indicate a definite difference. The diet of a colony of 60 white rats was so arranged that each rat ingested approximately 15 mg. of lead carbonate per day (4). At the end of 52 days half of the surviving animals were killed. The remaining animals were continued for 14 days on a similar diet from which lead carbonate had been omitted and were then killed.

At death, the livers, kidneys, spleens, and bones were removed for lead analysis, and sufficient blood was taken for a calcium determination. The results thus obtained are given in table 1. The analytical results are based upon wet tissue weights. Since the spleens were very small and a loss was entailed in pathological examination, the amounts of lead determinable were very low and therefore less reliable than the lead content of liver, kidney, or bone.

Inspection of the values thus obtained indicates a distinctly greater quantity of lead in the livers and kidneys immediately following the lead ingestion period as compared with the values obtained following The average content in the livers of the former group the rest period. was 0.020 mg. per liver, or a concentration of 0.032 mg. per 10 gm. of The average content in the livers of the group following the tissue. rest period was 0.011 mg., or 0.016 mg. per 10 gm. of liver. Similarly. the average content in the kidneys of the first group was 0.055 mg., or 0.427 mg. per 10 gm. of kidney, while in the second group the average lead content was 0.031 mg. per kidney, or 0.217 mg. per 10 gm. The average lead content in the spleens of the first group of tissue. was 0.003 mg., as compared with 0.004 mg. in the second group, but, as stated previously, these quantities are so low that comparisons are scarcely reliable. The lead content of the livers and kidneys of the animals following a rest period of 14 days amounted to only 50 percent of that in those animals killed directly at the end of the lead-feeding period.

	ţ		Liver			K idne;	y		Spleer	1		Bone		100 cc.
Rat No.	Weight at dea	Weight, gm.	Total Pb, mg.	Pb/10 g., mg.	Weight, gm.	Total Pb, mg.	Pb/10 g., mg.	Weight, gm.	Total Pb, mg.	Pb/10 g., mg.	Weight, gm.	Total Pb, mg.	Pb/10 g., mg.	Blood Ca, mg/
344	$\begin{array}{c} 170\\ 170\\ 170\\ 144\\ 143\\ 150\\ 160\\ 142\\ 150\\ 185\\ 160\\ 185\\ 160\\ 172\\ 144\\ 165\\ 133\\ 135\\ 163\\ 160\\ 150\\ 150\\ 160\\ 160\\ \end{array}$	$\begin{array}{c} \textbf{7.556} \\ \textbf{6.567} \\ \textbf{6.567} \\ \textbf{6.5680} \\ \textbf{5.6651} \\ \textbf{6.5651} \\ \textbf{6.5651} \\ \textbf{6.680} \\ \textbf{5.6651} \\ \textbf{6.680} \\ \textbf{5.6651} \\ \textbf{6.68275} \\ \textbf{5.6651} \\ \textbf{5.6651} \\ \textbf{5.064} \\ \textbf{5.064} \\ \textbf{5.064} \\ \textbf{5.064} \\ \textbf{5.266} \\ 5.2$	$\begin{array}{c} 0.\ 022\\ .\ 021\\ .\ 023\\ .\ 026\\ .\ 016\\ .\ 016\\ .\ 019\\ .\ 020\\ .\ 020\\ .\ 019\\ .\ 023\\ .\ 020\\ .\ 023\\ .\ 020\\ .\ 020\\ .\ 025\\ .\ 015\\ .\ 027\\ .\ 016\\ .\ 021\\ \end{array}$	0.029 027 038 034 028 035 030 025 027 034 034 032 034 032 034 032 041 043 040 043 040 050 043 040 043 040 043 040 022 040 043 040 043 040 043 040 043 040 043 040 041 025 034 034 025 034 034 035 034 035 035 034 035 035 034 035 035 035 034 035 034 035 035 034 035 035 035 035 035 035 035 034 035 035 035 035 035 035 034 035 035 035 034 035 035 035 035 035 035 035 035 035 035	$\begin{array}{c} 1.13\\ 1.48\\ 1.16\\ 1.11\\ 1.33\\ 1.06\\ 1.45\\ 1.30\\ 1.37\\ 1.20\\ 1.37\\ 1.20\\ 1.37\\ 1.20\\ 1.44\\ 1.32\\ 1.26\\ 1.125\\ 1.07\\ 1.07\\ 1.27\\ 2.08\\ 1.25\\ 1.25\\ 1.25\\ 1.25\\ 1.25\\ 1.26$	$\begin{array}{c} 0.\ 053\\ .\ 050\\ .\ 042\\ .\ 037\\ .\ 054\\ .\ 054\\ .\ 054\\ .\ 054\\ .\ 054\\ .\ 054\\ .\ 056\\ .\ 081\\ .\ 056\\ .\ 084\\ .\ 042\\ .\ 056\\ .\ 084\\ .\ 042\\ .\ 056\\ .\ 064\\ .\ 057\\ .\ 030\\ .\ 064\\ .\ 055\\ .\ 069\\ .\ 069\\ .\ 091\\ \end{array}$	0. 465 .337 .365 .333 .287 .404 .320 .591 .421 .476 .582 .312 .373 .400 .294 .476 .555 .500 .720	$\begin{array}{c} \textbf{0.56}\\ \textbf{.82}\\ \textbf{.63}\\ \textbf{.94}\\ \textbf{.85}\\ \textbf{.72}\\ \textbf{.72}\\ \textbf{.72}\\ \textbf{.86}\\ \textbf{.72}\\ \textbf{.86}\\ \textbf{.72}\\ \textbf{.86}\\ \textbf{.72}\\ \textbf{.86}\\ \textbf{.76}\\ \textbf{.76}\\ \textbf{.76}\\ \textbf{.76}\\ \textbf{.89}\\ \textbf{.89}\\ \textbf{.89}\\ \textbf{.87}\\ \textbf{.893}\\ \textbf{.87}\\ \textbf{.894}\\ \textbf{.87}\\ \textbf{.894}\\ \textbf{.87}\\ \textbf{.895}\\ \textbf{.87}\\ \textbf{.895}\\ \textbf{.87}\\ \textbf{.895}\\ \textbf{.87}\\ \textbf{.895}\\ \textbf{.87}\\ $	0.003 .004 .000 .002 .000 .003 .003 .003 .003 .003 .003 .003 .003 .003 .003 .003 .003 .003 .003 .004 .005 .003 .005 .005 .003 .005 .003 .005 .003 .005	$\begin{array}{c} 0.\ 048\\ .\ 049\\ .\ 049\\ .\ 000\\ .\ 045\\ .\ 035\\ .\ 037\\ .\ 000\\ .\ 058\\ .\ 035\\ .\ 035\\ .\ 024\\ .\ 038\\ .\ 025\\ .\ 024\\ .\ 058\\ .\ 025\\ .\ 031\\ .\ 000\\ .\ 032\\ .\ 051\\ .\ 060\\ .\ 050\\ \end{array}$	$\begin{array}{c} \textbf{2.106} \\ \textbf{2.366} \\ \textbf{1.912} \\ \textbf{2.90} \\ \textbf{2.906} \\ \textbf{2.300} \\ \textbf{2.617} \\ \textbf{2.700} \\ \textbf{3.40} \\ \textbf{2.617} \\ \textbf{2.715} \\ \textbf{2.61} \\ \textbf{2.761} \\ \textbf{2.761} \\ \textbf{2.761} \\ \textbf{3.002} \\ 3.002$	0.66 32 75 41 1.42 .39 1.47 1.49 1.02 .96 .37 .57 1.22 	2.24 1.80 3.93 1.63 4.90 1.43 4.90 4.38 3.50 4.46 1.34 2.60 5.08 	9.7 10.4 11.0 10.2 8.9 9.7 10.6 9.7 12.5 8.2 9.7 12.5 8.2 9.7 9.9 9.9 9.9 9.9 9.9 9.9 10.4 1.0 1.0 1.0 1.0 1.0 1.0 1.0 1.0
Average			. 020	. 032		. 055	. 427		(. 003)	(. 035)		. 93	3. 30	10.0

TABLE	1Group	I.	Lead content	of	organs	of	' animals	killed	at	end	of	experiment
TUDDE	II aroup .			~,	0.94.00	~,	annand	1000000		0.000	~	

TABLE 1.—Group I	. Lead	l content of	organs of	' animals	after	14-day	lead-free	diet
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			Liver			Kidne	y		Spleer	n		Bone		
Rat No.	Weight at death	Weight, gm.	Total Pb, mg.	Pb/10 g., mg.	Weight, gm.	Total Pb, mg.	Pb/10 g., mg.	Weight, gm.	Total Pb, mg.	Pb/10 g., mg.	Weight, gm.	Total Pb, mg.	Pb/10 g., mg.	Blood Ca, mg/10
369 369 370 371 371 373 373 374 374 375 376 376 377 378 379 383 384 383 384 384 385 386 389 389 390 391 392 383 394 385 394 392 393 394 396 396	2000 1800 1800 1577 1600 1577 1600 1700 1600 1705 1655 1655 1655 1651 1600 1500 1600 1900 1605 1622 1855 1855 201	$\begin{array}{c} 7.\ 08\\ 6.\ 00\\ 5.\ 98\\ 7.\ 42\\ 5.\ 37\\ 6.\ 21\\$	0.010 .011 .018 .002 .004 .007 .002 .034 .003 .004 .004 .009 .013 .008 .018 .018 .018 .018 .018 .009 .004 .009 .004 .009 .008 .018 .012 .009 .004 .009 .008 .013 .008 .018 .009 .008 .013 .008 .018 .009 .008 .013 .008 .018 .009 .008 .008 .018 .009 .008 .008 .018 .008 .008 .018 .008 .008 .018 .008 .018 .008 .008 .018 .008 .018 .008 .018 .008 .018 .008 .018 .008 .018 .008 .018 .008 .018 .008 .018 .018 .008 .018 .018 .008 .014 .014 .018 .018 .018 .014 .014 .018 .018 .018 .014 .014 .009 .018 .018 .014 .014 .009 .008 .018 .016 .014 .014 .009 .008 .018 .016 .008 .016 .008 .016 .008 .016 .008 .016 .008 .016 .008 .016 .008 .016 .008 .016 .008 .016 .008 .016 .004 .009 .004 .005 .004 .004 .005 .004 .005 .004 .006 .004 .005 .004 .006 .004 .006 .004 .006 .004 .006	0.014 0.018 0.016 0.000 0.006 0.011 0.042 0.005 0.007 0.019 0.011 0.013 0.017 0.014 0.020 0.020 0.020 0.014 0.0210000000000	$\begin{array}{c} 1.\ 61\\ 1.\ 32\\ 1.\ 50\\ 1.\ 32\\ 1.\ 30\\ 1.\ 32\ 1.\ 32\ 1.\ 32\ 1.\ 32\$	0.049 .031 .019 .046 .020 .026 .020 .032 .035 .050 .017 .027 .044 .020 .038 .025 .050 .017 .027 .044 .020 .038 .025 .038 .025 .036 .021 .046 .020 .032 .046 .021 .032 .035 .046 .021 .035 .035 .046 .021 .035 .046 .021 .035 .035 .046 .035 .046 .035 .046 .035 .046 .035 .046 .046 .046 .046 .026 .035 .045 .045 .045 .045 .045 .045 .045 .04	0.304 -233 -125 -350 -154 -200 -154 -223 -254 -150 -1254 -223 -254 -150 -167 -178 -167 -178 -271 -271 -271 -271 -272 -212 -215 -215 -200 -154 -201 -201 -201 -201 -201 -201 -201 -201	0.71 .81 .66 .67 .53 .59 .53 .59 .53 .59 .53 .59 .84 .80 .87 .77 .68 .67 .68 .67 .53 .59 .59 .84 .80 .65 .53 .59 .84 .80 .67 .77 .68 .67 .77 .68 .67 .70 .84 .80 .84 .80 .80 .80 .80 .80 .80 .80 .80 .80 .80	0.002 .004 .003 .002 .000 .001 .005 .000 .003 .001 .007 .002 .004 .001 .003 .004 .003 .004 .003 .004 .003 .005 .007 .007 .007 .007 .007 .007 .007	0. 035 048 052 040 027 000 025 088 000 040 040 036 058 017 166 050 056 056 056 056 074 074 074 071 098	$\begin{array}{c} \textbf{4.05}\\ \textbf{3.02}\\ \textbf{3.22}\\ \textbf{3.09}\\ \textbf{3.07}\\ \textbf{4.165}\\ \textbf{2.752}\\ \textbf{3.266}\\ \textbf{3.13}\\ \textbf{2.802}\\ \textbf{3.05}\\ \textbf{3.13}\\ \textbf{2.802}\\ \textbf{3.055}\\ \textbf{3.05}\\ \textbf{3.45}\\ \textbf{3.666}\\ \textbf{3.280}\\ \textbf{3.566}\\ \textbf{3.666}\\ \textbf{3.666}\\$	1.55 1.17 1.21 1.17 1.18 .53 .87 1.13 1.36 .80 1.00 1.22 .53 1.07 .55 1.78 .07 1.21 1.39 .69	$\begin{array}{c} 3.83\\ 3.79\\ 3.76\\ 3.79\\ 3.85\\ 2.00\\ 3.22\\ 4.01\\ 4.49\\ 3.58\\ 3.58\\ 4.86\\ 2.65\\ 3.28\\ 3.54\\ 4.86\\ 2.08\\ 3.54\\ 4.86\\ 3.28\\ 3.54\\ 4.86\\ 3.28\\ 3.54\\ 4.86\\ 4.86\\ 3.28\\ 3.54\\ 4.86\\ 6.60\\ 2.34\\ 4.86\\ 4.92\\ 3.36\\ 3.40\\ 4.86\\ 4.92\\ 3.36\\ 3.40\\ 4.92\\ 3.36\\ 3.40\\ 4.92\\ 3.36\\ 3.40\\ 4.92\\ 3.36\\ 3.40\\ 4.92\\ 3.36\\ 3.40\\ 4.92\\ 3.36\\ 3.40\\ 4.92\\ 3.36\\ 3.40\\ 4.92\\ 3.36\\ 3.40\\ 4.92\\ 3.36\\ 3.40\\ 4.92\\ 3.36\\ 3.40\\ 4.92\\ 3.36\\ 3.40\\ 4.92\\ 3.36\\ 3.40\\ 4.92\\ 3.36\\ 3.40\\ 4.92\\ 3.36\\ 3.40\\ 4.92\\ 3.36\\ 3.40\\ 4.92\\ 3.36\\ 3.40\\ 3.40\\ 4.92\\ 3.36\\ 3.40\\ 4.92\\ 3.40\\ 4.92\\$	$\begin{array}{c} 11.\ 0\\ 11.\ 0\\ 8\\ 12.\ 5\\ 9.\ 7\\ 10.\ 1\\ 10.\ 8\\ 10.\ 8\\ 10.\ 8\\ 10.\ 8\\ 10.\ 8\\ 10.\ 8\\ 10.\ 8\\ 11.\ 9\\ 11.\ 9\\ 10.\ 8\\ 11.\ 9\\ 11.\ 9\\ 10.\ 8\\ 11.\ 9\\ 11.\ 9\\ 11.\ 9\\ 12.\ 5\\ 11.\ 0\\ 12.\ 1\\ 12.\ 5\\ 11.\ 0\\ 12.\ 12.\ 12.\ 12.\ 12.\ 12.\ 12.\ 12.\$
Average			. 011	. 016		. 031	. 217		(. 004)	(. 049)		1.14	3. 65	11.4

In contrast with the lead content of the softer tissues, the bones show a slight increase in lead content in the second group, although these animals received no more lead than the first group and furthermore had an opportunity of excreting part of the lead during the 14day rest period. The average amount of lead present in the bones of the animals of the first group was 0.93 mg. per animal, or 3.30 mg. per 10 gm. of bone tissue, while that of the second group was 1.14 mg. per animal, or 3.65 mg. per 10 gm. of bone tissue. Thus, there was a slight but unmistakable increase in bone lead which must have occurred at the expense of the lead contained in the softer tissues (fig. 1).



The blood calcium values of the two groups show individual variations about the normal figure. However, in common with other values obtained in lead-poisoned animals, there appears to be a slight rise in the average blood calcium value on return to a normal diet. The average blood calcium value of the first group was 10.0 mg. per 100 cc. as compared with 11.4 mg. following the rest period. Further experimental work is in progress with respect to the possible effect of lead absorption upon blood calcium.

PATHOLOGY

Paraffin sections were made from the liver, kidneys, spleen, pancreas, heart, stomach, duodenum, jejunum, ileum, large intestine, and mesenteric lymph nodes and were routinely stained by Lillie's eosinpolychrome methylene blue method (δ). Sections from all of the spleens and a representative number of kidneys were treated with acidulated ferrocyanide to demonstrate the presence or absence of iron-bearing pigment. A random selection of kidney sections was stained with hematoxylin and eosin as a check on the eosin-methylene blue stain for the detection of oxyphil intranuclear inclusions. A total of 1,000 sections from 53 rats was studied.

Kidneys.—Sections from the kidneys of 22 rats killed immediately after the 6-week feeding period were available for study. Enlarged cells, with large vesicular nuclei and no radial striations, were observed in the proximal convoluted tubules of the 22 animals and in both the proximal and distal convoluted tubules in 17 rats. Round, brown, cytoplasmic pigment granules were noted in the swollen tubular cells of only 3 rats, and traces of minute, brown pigment particles were present in the same region in 6 other animals.

Neither the brown granules nor the pigment particles reacted to a test for iron. Small interstitial collections of lymphocytes were noted in 9 rats. In two instances they were accompanied by casts in the straight collecting tubules and by a few small areas of tubular cells with basophilic cytoplasm, indicative of very early retrograde change. Lymphocytes, isolated or in very small foci, were also observed in the renal interstitial tissue in 6 animals. The glomerular tufts and interstitial capillaries were not congested.

Swollen convoluted tubule cells were much less evident in the kidneys of 24 rats fed lead carbonate for 6 weeks and returned to a normal diet for 2 weeks than they were in the preceding series. In the preceding series the typical finding was a three plus or marked degree of prevalence of swollen convoluted tubule cells; in this series the typical finding was a one plus degree of prevalence. As before, this phenomenon was principally localized in the proximal tubules. It was present in the distal tubules of all but 9 rats. Small numbers of brown pigment particles were almost invariably present in a few swollen cells in the proximal convoluted tubules. This was a more regular finding in this series than in the preceding series.

A few oxyphil intranuclear inclusion bodies were present in the swollen cells of the proximal convoluted tubules in only one animal. A few lymphocytes were found in the interstitial tissue of the kidney in one rat. Casts were absent in all of the animals. Two of the rats showed a slight congestion of the capillaries but this appeared to be of no significance. Spleen.—The splenic corpuscles were generally large and well defined and were surrounded by fairly large zones of pale-staining cells. The cavernous veins were usually filled with blood. The degree of relative perifollicular anemia varied inversely with the amount of blood in the cavernous veins. Splenic myelosis with accompanying megakaryocytes was present in a degree approximating that found in normal rats. The marked decrease in number of myeloid cells observed in rats fed lead carbonate for 1 and 2 years (4) was not noted in these 6-week feeding experiments. Diffuse iron reaction of cells of the pulp, however, was similar in degree to that found in animals fed for longer periods. A small number of brown pigmented particles which did not react for iron were also present. Lymphocytic infiltration of the muscular trabeculae was slight to moderately marked and occurred in all of the animals. Nuclear fragments were found in the follicles in 20 of the 24 animals examined.

In the rats allowed the normal diet for 2 weeks following 6 weeks of lead ingestion the splenic corpuscles were usually fairly small and well defined with a surrounding zone of paler-staining cells. This zone of relative anemia and the amount of blood in the cavernous veins were identical in degree with those found in the animals examined immediately after the lead ingestion period. The amount of splenic myelosis and the number of megakaryocytes were somewhat reduced both in degree and in the number of animals in which they were found. Myeloid hyperplasia, generally present in rats, was absent in 2 and very scant in 4 of the animals of this series. It was not as prominently decreased as was noted in rats fed lead carbonate for 1 and 2 years.

The amount of iron-bearing pigment in the pulp was likewise somewhat diminished in degree and was absent or very scant in 8 of the 24 rats examined. Follicular phagocytosis, indicated by nuclear fragments in the follicles, was absent in 8 animals and was decreased in amount to half that noted in the animals killed immediately after feeding. Lymphocytic infiltration of the trabeculae was essentially the same as observed in the animals examined immediately after the lead feeding period.

Liver.—The cytoplasm of the liver cells was finely granular and usually fairly dense. The nuclei showed no unusual variation in size and no oxyphil inclusions were noted. The histological picture was essentially the same following return to the normal diet.

Lungs.—Subacute bronchopneumonia, in one case with bronchial epithelium proliferation, occurred in 4 of 23 rats examined in the first series and in 4 of 28 rats examined in the second group. In 2 others of this latter group the larger bronchi were filled with polymorphonuclear leucocytes, granular debris, and nuclear remains. As in the first group of animals these changes appeared to be unrelated to the lead unless they might be interpreted as an indication of general debility. The heart, pancreas, stomach, duodenum, jejunum, ileum, large intestine, and mesenteric lymph nodes showed no changes of note.

DISCUSSION

The most significant change observed in the kidney was the presence of swollen cells, with large vesicular nuclei and granular cytoplasm, in the convoluted tubules. They occurred in the proximal convoluted tubules in all of the animals but were less numerous and frequently absent in the distal convoluted tubules in both series of tests. These cells were noted in isolated tubules or in groups of tubules. The number of these cells was appreciably reduced in the animals returned to a normal diet. The oxyphil intranuclear inclusions described by Blackman (6) and consistently found in the nuclei of rats fed lead carbonate for 1 and 2 years (4) were not present except in one animal which had been returned to a normal diet for 2 weeks. In this instance they were few in number. Brown pigment granules were found in the swollen cells of 3 rats and traces of brown pigment in 6 others studied at the end of the 2-week lead-ingestion period. Traces of this pigment were found in 23 of the 24 rats following return to a normal diet for 2 weeks. Like the swollen cells, the pigment occurred most frequently in the proximal convoluted tubules.

It is interesting to note that the proximal convoluted tubule cells show the most change as indicated by the hyper-regenerative cells and the cytoplasmic brown pigment. Marshall and Grafflin (?) have shown that the proximal convoluted segment of the renal tubule of the glomerular fish, sculpin, can both secrete and reabsorb. This would suggest that the cells of the proximal convoluted tubules show the greatest damage because of their greater selective exposure. In a previous study (4) it was suggested that lead has a damaging action on the kidney and these findings tend to confirm those experiments. The significant reduction in the pathological findings, both in degree and in number of animals affected, following return to a normal diet, seems to indicate that the toxic effects on the kidney, with due consideration to amount and duration of exposure, are susceptible of repair after the intake of lead has ceased. This is in keeping with Beintker's (8) opinion that "temporary degenerative changes" occur at the beginning of lead intoxication.

Hemosiderin was present in the spleens of all of the animals examined immediately after the lead-ingestion period. It was slight to moderately marked in degree. Hemosiderosis was essentially of the same degree as that found in animals fed lead carbonate for 1 and 2 years. In the animals examined 2 weeks after cessation of the lead diet, hemosiderin was absent in 2 and only minute traces were found in 6 rats. The average degree was also about one-third less than that found in the animals killed at the end of the lead-feeding period. Thus it appears that blood destruction in the spleen is a relatively early finding and that it tends to diminish when lead is withheld from the diet.

Splenic myelosis or erythroleucopoietic activity, on the other hand, did not vary appreciably from that usually found in normal rats. It



FIGURE 2.—Number and percentage of experimental animals found to have specified amounts of pathologi changes. Percentage of animals with specified reaction: 3 +, marked; 2 +, moderately marked; 1 + moderate; \pm , slight; blank, no deviation from normal or very slight amount.

was, however, slightly reduced in the animals examined 2 weeks after return to normal diet. The marked decrease of splenic myelosis seen in rats fed lead carbonate for 1 and 2 years was not present in those animals used in the 6-week feeding experiments. Seiffert and Arnold (9) have observed an increased destruction of blood cells in the spleen, particularly the erythroblasts. Such a destruction of blood would not only account for the presence of the hemosiderin but also would suggest that the decrease in myeloid cells, prominent in the 1- and 2year experiments, is due to the destruction of these young forms. Thus, it appears that significant interference with blood formation in the spleen takes place some time after 6 weeks and before 1 year. Nuclear fragments in the follicles were frequently found in the animals examined at the end of the lead carbonate feeding period. Their occurrence, both as to number of animals involved and average degree, was appreciably reduced after a return to normal diet for 2 weeks. Recent cell destruction, as indicated by this follicular phagocytosis, also appears to regress when the ratis returned to normal diet. Other changes in the spleen, such as the perifollicular zones of anemia and the amount of blood in the cavernous veins, seem to be of minor importance.

Changes in the other organs examined were not significant.

It appears from these pathological findings that injury produced by lead in the kidney and spleen of the rat occurs relatively early, is of a temporary nature, and is subject to repair (fig. 2).

CONCLUSIONS

It has been shown that lead deposited in the softer tissues in rats fed approximately 15 mg. of lead carbonate a day for 6 weeks is transitory in nature and may be diminished by 50 percent within 2 weeks merely by restoration to a normal diet. Coincidently there is a slight rise in the lead content of the bone tissue. A slight depression in blood calcium was noted in animals fed the lead diet as compared with those restored to the normal diet.

The ingestion by rats of 15 mg. of lead carbonate per day causes rather severe injury to the cells of the renal convoluted tubules, particularly those of the proximal group, in 6 weeks. This damage is markedly reduced following return to a normal diet for 2 weeks, and appears to be temporary in nature. Significant pathologic changes in the spleen are reduced to some extent after the leaded diet is withdrawn, but the approach to normal is not as marked as in the kidney.

The reduction of lead in the soft tissues observed chemically parallels the repair of injury produced by the lead.

The chemical and pathologic changes associated with the softer tissues of rats following the ingestion of lead carbonate in the quantity used in this study would appear to be of a transient nature.

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WEIL'S DISEASE¹

A REPORT OF 51 CASES OCCURRING IN PUERTO RICO AND THE UNITED STATES

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Weil's disease has been commonly reported in Europe and Asia, but there are relatively few reports in the literature concerning the disease in this country.

The National Institute of Health from time to time receives specimens from various parts of the country for examination as an aid in the diagnosis of Weil's disease. In many cases physicians also submit clinical data where the diagnosis of Weil's disease is confirmed. The material presented in this paper and based upon this type of data indicates that leptospirosis is by no means a rare disease in this country and that the infrequency with which it is reported is due either to failure to recognize the disease, or to infrequent reporting of the condition.

It is not the purpose of this paper to discuss the clinical aspects of leptospirosis since this belongs in the province of the physician who treats the patient. It is to be hoped that these physicians will describe their cases in adequate clinical notes in order to further recognition of the disease. With the same objective of stimulating interest in this condition, it is proposed in this paper to summarize briefly the American literature to date and to present additional cases diagnosed with the aid of tests performed in this laboratory.

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In 1905 Stimson (1) observed an organism, which he called Spirochaeta interrogans, in the tissue of a patient dying during an outbreak of yellow fever in New Orleans. This is probably the first instance in which Weil's disease was encountered in this country (2, 3), although it was then not recognized as such. Wadsworth et al. (4) reported a case in a laboratory worker in 1922. Following this a number of other cases were reported (5, 6, 7, 8, 9, 10, 11), and by 1935 Jeghers, Houghton, and Foley (12) were able to add the 11th case to the American literature, exclusive of the one noted by Stimson.

In considering the incidence of Weil's disease in this country it might be well to separate the proved cases from those merely alluded to in the literature, and from those in which adequate laboratory data were not obtained. Thus, 10 of the cases cited above are proved, while 2 others (6, 11) are not fully confirmed. It is difficult to evaluate the 7 cases reported by Gaines and Johnson (13) and there is little data upon which to form conclusions concerning reports from Texas (14) and Rhode Island (15).

Molner and Meyer (16) state that 7 cases of Weil's disease occurred in Detroit, and Meyer et al. (17) have evidence concerning at least 11 cases in California. Syverton, Stiles, and Berry (18) call attention to a case in Rochester, N. Y., and Goldberg and Davens (19) refer to another in Baltimore, Md. These seem to be founded on adequate evidence, but further description is desirable.

The 5 instances of leptospirosis cited by Mulholland (20), Martmer (21), Glotzer (22), Elton (23), and Haschec and Tobey (24) are fully described and based on sufficient laboratory evidence to establish their identity. In addition, 12 other cases which have been reported recently fulfill all the requirements necessary for their acceptance as proved cases of Weil's disease (19, 25, 26, 27, 28). It would appear that there are at least 27 fully described cases and 20 presumptive cases of Weil's disease reported in the American literature at the present time.

Most of these cases have occurred among adult males, but 3 children and 2 adult females are included in the group of 47. In most instances the occupation of the individual accounted for sufficient exposure to infection. One case occurred in a laboratory worker and a number of cases have been traced to swimming in water to which wild rats have easy access. Cooks, gardeners, meat-handlers, dairymen, veterinarians, poultry dressers, quarry workers, sewer workers, and fish cutters have been involved. Geographically the cases are widespread, occurring in 11 States and the District of Columbia (table 1).

	Cases previou	usly reported	Cases not	t previously	y reported	
State or area	Proved	Presump- tive	Icteric	Anicteric	No clini- cal data	Total
Alabama			10	2		12
California	1 (11)	11 (17)	1			13
Connecticut	1 (25)		1		1	3
District of Columbia	1 (9)			1		2
Georgia			1			1
Louisiana						1
Maryland	1 (19)	1 (19)	2	1		8
Massachusetts	1 (12)	7 (16)	-			10
Michigan	1 (01)	1 (10)	1		1	13
New Vork	5 (4.5.7. 22)	1 (18)	•		-	5
Ohio	8 (27, 28)	- (,	1			ä
Pennsylvania	4 (6, 8, 27)		3			7
Virginia	2 (10, 20)		2	3		7
Wisconsin			3			3
Puerto Rico			2		3	5
Total	27	20	34	7	10	98

TABLE 1.—Geographic distribution of cases of Weil's disease

NOTE.-Figures in parentheses refer to articles in bibliography in which cases were reported.

DATA ON NEW CASES OF WEIL'S DISEASE

The data presented in this paper include 51 cases of Weil's disease previously unreported from the United States and Puerto Rico. Among them are 33 cases having a more or less classical picture of the condition. One additional patient from Alabama was jaundiced about 2 years prior to serological examination and was not hospitalized, although he was unable to work for a month after onset of illness. The data also contain material concerning 7 individuals possessing antibodies against Leptospira icterohaemorrhagiae who had never had an illness in which jaundice was an apparent symptom. These were observed during the course of routine serological examinations of groups considered to be exposed to danger of infection. Agglutination and mouse protection tests (29) have been used to identify these cases. In addition, agglutination tests have revealed agglutinins in diagnostic titers against this organism in 10 human serums derived from Michigan, New Jersey, Puerto Rico, and Connecticut. No clinical material has been made available in this group of cases. The cases are summarized in table 2.

The geographic distribution of the cases are shown in table 1, together with the distribution of previously reported cases. Thus, 98 cases of Weil's disease have been observed in Puerto Rico and in 14 States and the District of Columbia in the United States. Alabama, Georgia, Wisconsin, and Puerto Rico have been added to the area from which cases had previously been studied. It is a matter of singular interest that in any given locality the reporting of cases increases with the discovery of a single case. In Wisconsin 2 cases were reported soon after the diagnosis had been established in one patient. Four cases were recognized in Puerto Rico following the identification of the original case. It is difficult to believe that Weil's disease is present only in certain localized areas when conditions necessary for its occurrence in human beings are so widespread. From experience in the various areas it would appear that the spotty distribution of cases is probably due to failure to recognize the disease.

The distribution according to sex of the cases in this series is essentially the same as given in previous reports. Thirty-nine of the patients were males and 2 were females. One frank infection was noted in a colored female between 50 and 60 years of age and an inapparent infection occurred in a female aged 22 years. There were 29 cases in white and 11 in colored individuals. The patients ranged from 18 to 58 years of age, with an average of 33.9 years.

It has been stated (27) that the majority of cases reported in this country occurred during the summer months. In our series 18.5 percent of the cases occurred from December through February, 22.2 percent from March through May, 40.7 percent from June through August, and 18.5 percent from September through November.

There are no unusual features about the occupations of the individuals concerned in this report. Among the anicteric or inapparent cases were a waitress, a carpenter, a market employee, 2 slaughterhouse employees, and 2 coal miners, while among the cases with typical Weil's disease were sewer workers, merchants, coal miners, a barber, a plumber, a seamen, a farmer, fishcutters, poultry dressers, a bricklayer, and slaughterhouse workers. Two patients had handled rats prior to the onset of illness.

SUMMARY

A series of 51 previously unreported cases of Weil's disease occurring in Puerto Rico and the United States are discussed. Thirty-three of these are based on adequate clinical and laboratory data and 7 upon laboratory data alone. Another group of 7 anicteric and inapparent cases is also included.

TABLE	2.—Clinical	data	concerning	previously	unreported

					а	tion	omet	onset
Numbe	Name	Age	Ser	Color	Locatio	Occupa	Date of	Type of
123	JC TH SB	35 23 23	M M M	w w w	Roxbury, Mass Atlanta, Ga Cambridge, Mass	Storekeeper Picked up rat Slaughterhouse worker.	June 14, 1940 July 25, 1940 Feb. 3, 1941	Sudden do do
4 5 6	MT CL CS	46 47 27	M M M	W W W	Los Angeles, Calif New Britain, Conn Milwaukee, Wis	Plumber Merchant Slaughterhouse worker.		
7 8 9	CB EW	48 51 30	M M M	W W W	dodo Secaucus, N. J	Sewer worker do Slaughterhouse worker.	May 27, 1939 1941	Sudden Sudden
10 11 12	RT JL GL	+ 48 40	M M M	W W W	Chelsea, Massdo Cincinnati, Ohio	Fish cutter do Unemployed	Jan. 4, 1941 Sept. 16, 1940 June 1940	
13 14	JM	18	M M	w	Philadelphia, Pa	Dicked up rat Aug. 12, 1940.	Aug. 17, 1940	
15 16 17 18	FDA JA EDR GM	28 41 39 25	M M M	W W C	Puerto Ricodo Richmond, Va	Barber Seaman Farmer Poultry picker	June 12, 1940 Nov. 30, 1940 Nov. 4, 1940 Dec. 1, 1940	Sudden
19 20 21 22	RM RP LT RHC Jr	28 35 21	M M M		Birmingham, Alado do Lewisburg, Ala	Coal miner dodo	Mar. 14, 1938 June 23, 1938 Apr. 7, 1940	Suaden do do
23 24 25 26 27	RS GM RS	39 24 27 22	M M M	₩₩ ₩₩	Lewisburg, Ala.	do	Aug. 1, 1938 Oct. 21, 1938 Mar. 27, 1939 Nov. 28, 1939	
28 29 30	CF PE SE EP	39 50-60 58 57	M F M	o¥c¥c	Gardendale, Ala Baltimore, Md dodo.	dodo Poultry dresser .	June 19, 1940 Aug. 19, 1940 Dec. 10, 1940 Feb. 22, 1941	· · · · · · · · · · · · · · · · · · ·
32 33 34 35	WT GY RDC IM	28 42	M M M M	Č ₩ ₩	do do Lewisburg, Ala Seraucus, N. J	Sewer worker Coal miner	Apr. 16, 1941 May 9, 1941	
36 37 38 39	AG CP		M M M		Puerto Rico do do Connecticut			
40 41 42 43	JE VC FH SW		M M		Detroit, Mich do do do do			
44 45 46 47	HB WF	57	M M M	W W C	do Baltimore, Md Richmond, Vado	Carpenter Marketer Slaughterer		
48 49 50 51	AB LH DF	22 	M F M M	W W W	Washington, D. C. Lewisburg, Alado	Waitress Coal miner		

¹ Died. Typical lesions of Weil's disease noted at post-mortem examination.

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Chill	Гетег , ° Г .	Headache	Nausea and/or vom	Pain in chest, legs, o domen	Liver enlargement	Conjunctivitis	Hemorrhages	Oliguria	Jaundice	Ictaric Index	N. P. N. Mg. (perce	B. P.	Hb. (percent)	R. B. C.	W. B. C.	P. M. N. (percent)	Casts	W.B.C.	R. B. C.	Albumin	Bile	Agglutination test v ktero. Results
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cases of Weil's disease in the United States and Puerto Rico

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SOME SPECIAL EPIDEMIOLOGICAL AND CLINICAL FEA-TURES OF PLAGUE IN NORTHEASTERN BRAZIL¹

By ATILIO MACCHIAVELLO, Epidemiologist, Pan American Sanitary Bureau

An epidemiological and bacteriological study of plague in northeastern Brazil extending from August 1939 to September 1940, made in collaboration with Brazilian specialists,² brought to light a number of interesting features regarding the disease in man, rats, fleas, and wild rodents in rural areas.³

The region studied includes an agricultural district subject to severe droughts, and semidesert areas which alternate with swamps and rocky ground. The following were found to be important factors in the epidemiology of plague in this region: Rains (when human plague increases, as in 1924–26, 1929, 1935, and 1939–40); droughts (when it decreases, owing to less cultivation, migration of local rodents in search of food, fewer fleas, and other conditions); the kind of agriculture (with mandioca, cotton, mamona, poroto (or peas), and particularly corn, the most attractive to rats, which migrate in search of the corn embryo, possibly for vitamin E); and the type of building construction.

The domestic black rat, *Rattus rattus*, was proved to be the most important factor in rural plague in northeastern Brazil, both human plague and plague among other rodents being accidental and secondary to plague in this rat. If true sylvatic plague does exist in Brazil, as has often been suggested, it has not, in the author's opinion, been conclusively demonstrated.

Xenopsylla cheopis and X. brasiliensis were the most common fleas on R. rattus and R. alexandrinus (which generally lives in the fields); Rhopalopsyllus and possibly Parapsyllus were found on field rats, prefix (Cavia aperea) and mocós (Heredon rupestris), though these rodents were rarely flea-infested. X. cheopis and a Chiastopsylla sp. were discovered on Monodelphis caught in rat nests. Some Pulex irritans, and, rarely, Echidnophaga gallinacea, were found on rattus and alexandrinus. However, X. cheopis was the only ectoparasite found plague infected. Even when X. brasiliensis were taken from a rat on which infected cheopis had been found, they failed to produce plague.

Plague infection was again demonstrated in the prea, Mus sylvaticus, and Mus minusculus, which previous investigators had found naturally

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¹ Résumé taken from a more extended paper appearing in Spanish in the Bulletin of the Pan American Sanitary Bureau, May 1941, p. 441. The complete report of these studies in Spanish may be obtained by applying to the Pan American Sanitary Bureau, Washington, D. C.

³ Brito, Garcia Rosa, Martins de Almeida, E. de Silva, Arcoverde, and Trigueros collaborated closely in the epidemiological studies, Paracampos in the bacteriological studies.

³ For a history of plague in Brazil see Bol. Of. San. Pan., Nov. 1940, p. 1081. (In English.)

infected; and it was reported for the first time in Brazil in Oryctolagus cuniculus (rabbit) and Lepus sp. (hare), and possibly Oryzomys intermedius.

During the studies, two factors were noted which explain many of the puzzling phenomena observed in connection with rural plague in In some rural areas, the high external temperatures (some-Brazil. times over 96.8° F.) prevent X. cheopis from living outside the nests of its rat host, in which the temperature is from 9° to 14.4° F. cooler than that outside. Under these circumstances, plague infection is found only in the nests, where the fleas remain, feeding usually on the young rats. A subterranean epizootic follows, slow in its progress, and with very little human repercussion. Should the weather change and the temperature drop, as happens during the rains, the fleas may leave their shelters, and the epizootic then comes to the surface. This explains the abrupt onset of plague following the rains. It may even spread to wild rodents, possibly through the intermediation of Monodelphis domesticus and certain field mice, especially M. sylvaticus and M. minusculus. An increase in the flea population and in the cheopis index of field rats and the presence of cheopis on wild rodents have been observed during cooler weather. The human cases, as might be expected, have occurred chiefly among individuals working around rat nests, or among persons whose beds are close to a wall in which rats have their holes.

The rat nests remain inhabited by infected fleas after a large part of the rat population has died. The remaining rats are immune to plague, and the disease would soon die out (the infection in fleas in the tropics, it was again demonstrated, becomes progressively more attenuated and disappears in a few weeks) were it not for the arrival of new susceptible animals, either as the result of breeding or through migration. It is thought that the decreasing virulence of plague in fleas may be responsible for the mildness of human cases at the end of epidemics.

It is believed that an area in which plague has died out because of a lack of susceptible material may be reinfected if the rodent population happens to become sufficiently great at the same time that an opportunity for reintroduction of the plague organism occurs, such as the migration of infected rats. If this happens, however, at a time when one of the periodic nonplague epizootics has practically wiped out the rodent population, reinfection will not occur, but may skip several years until another opportunity arrives. This would explain the reappearance of plague in 5- or 10-year cycles in certain areas. It may be mentioned that no evidence of chronic rat plague was discovered.

The other factor observed was that while the severe, periodic, extensive, and highly fatal epizootics among various wild animals are due to several nonplague organisms, and are a natural means of reducing the surplus population of a given species, plague epizootics may coexist with them (in the case of rodents). This circumstance and the plague-like characteristics of some of the nonplague epizootics have doubtless been the basis for the assumption that sylvatic plague exists in Brazil. Such epizootics had been observed, however, long before plague entered the country. Plague often coexists with these epizootics when they occur in migrating rodents, either because these rodents carry it with them or because they find it in the invaded regions. The nonplague epizootics are characterized by a different causative agent in each species (at times, however, in different epizootics, different causative agents may be found in the same species); by great severity and high mortality; by the conferring of a specific immunity lasting until the susceptible population is renewed; and by a form of disease usually septicemic in nature. Among the causative organisms discovered were Pasteurella lepiseptica, and possibly Past. nseudotuberculosis, in rabbits; Past. avicida in chickens and ducks; a colibacillus in mocos; and what seemed to be a new species of Pasteurella in Didelphis sp.⁴ The nonplague epizootics among rats were caused by Past. muricida, Loefflerella whitmori, Past. pseudotuberculosis rodentium, a Klebsiella which produced paralysis of the lower extremities. a paracolibacillus, and possibly Trypanosoma lewisi, with the roles of Listerella monocytogenes, Brucella bronchiseptica, Corynebacterium nseudotuberculosis murium, and Actinobacillus sp., undetermined, though these organisms were found.

It was observed that, when a nonplague epizootic attacks a plagueinfected rat community, there is a rapid increase in rat mortality, a rise in the flea index among the survivors, and an increase in the number of human cases of plague. The rapid increase in rat mortality, leading almost to complete extermination of the rat population, results in an abrupt decline in the number of cases of rat and human plague and cessation of the nonplague epizootic. After the epizootic ends, one or two cases of human plague may occur, possibly due to some remaining infected fleas. In wild rodents, the mixed infection makes it difficult to demonstrate plague, because, whereas plague requires at least 3 days to develop, the other disease produces an acute septicemia in less than 48 hours. During the investigations, five types of plague epizootics and three types of murine migrations were studied in detail.

With regard to human plague, the most common type was bubonic. There was, however, a lower mortality, and a lesser tendency of the buboes to suppurate, than in classic plague. No pneumonic plague, either primary or secondary, was observed. There were 20 cases of

⁴ Should this indeed prove a new type, the author suggests that it be named *Past. longi*, in recognition of the many years of plague work of Dr. John D. Long.

primary septicemic plague, some of which exhibited gastrointestinal and pulmonary symptoms, and jaundice.

The investigators were able to present the first bacteriologic proof that ingua de frio is actually plague. This disease, also known as febre de caroço, though the term was originally applied to ordinary bubonic plague, appears generally in children under 15, and is characterized by mild and transitory symptoms, monoglandular swellings without much inflammation or pain, and a tendency of the gland swelling to become ligneous, and to reoccur, or to be reabsorbed. It appears sporadically where plague is endemic and tends to disappear when epidemic. There is no special relationship between cases, though they occasionally appear in small foci. In some of these foci the first cases are severe and even fatal, the later ones becoming increasingly milder. There is a history of previous plague in rattus; and it is possible that this type of plague may have some relation to an attenuated virus in fleas.

A new plague syndrome was observed-a multiglandular fever. This was found in 9 out of 263 cases, and followed a prolonged course with a severe effect on the patient. It was characterized by septicemic fever, wasting, sometimes cachexia, pronounced anemia, multiple successive buboes about the size of an orange, with a tendency to suppurate, alopecia, frequent gastrointestinal and (or) urinary symptoms, and low mortality. The average case had more than 3 buboes (41.3 percent cervical, 37.9 percent inguinal, 13.8 percent cephalic, and 6.9 percent axillary), which usually appeared after the onset of the general symptoms. The clinical picture was that of attentuated plague (with P. pestis in the buboes but not in the blood), with septicemia. The latter was due in 3 cases to a paracolibacillus, in 3 to Brucella bronchiseptica, in 1 case to Klebsiella sp., and in 1 to Past. pseudotuberculosis rodentium (believed to be the sixth or seventh reported case of human infection with this organism, and the first of a mixed infection of it and plague). These cases appeared on the border areas of plague-infected and plague-free zones where conditions might be considered more or less unfavorable to plague.

The strains of the plague organism from *ingua de frío* studied in the laboratory were quite fragile and seemed to have lost their invasive power while retaining the power to produce a potent endotoxin. Those of the multiglandular fever were typical of, though less virulent than, ordinary strains.

The extensive epizootics seen among domestic cats in Brazil were found to be due, in some cases at least, to a filterable virus, and the disease was given the descriptive name "adeno-myelo-enterosis" by the author and his collaborator, Bezerra Coutinho. It was felt that there may be a definite relationship between this disease and migrating rats, since the rats can harbor the causative virus in the brain Some 40 plague strains of human, murine, preá, rabbit, flea, and other origin were studied, including a murine strain from rats from Recife, where there had been no human plague. Among the observations may be mentioned the finding that the morphology and appearance of plague cultures depended on the organ from which the strain was taken and on the culture medium, rather than, for instance, on virulence, although the smooth form, when present in flat colonies, or those with a depressed center, was usually the least virulent. Survival studies of P. pestis showed tropical conditions to be very unfavorable to it. It was observed that gram-negative bacilli, such as may cause some nonplague epizootics, were antagonistic to the development of plague organisms. Plague cultures from bone marrow for diagnostic purposes were not as successful as they have been reported elsewhere.

It is suggested that further investigation include, among other things, the study of the possible influence of cold weather and of feeding on the blood of susceptible animals in reviving the virulence of plague in fleas; the possibility of reviving the discredited biologic prophylaxis in view of nonplague epizootics in rats; and measures to destroy fleas in rat nests. (Flame-throwers and cyanogas are among the measures now being used.)

PREVALENCE OF ENCEPHALITIS IN THE UNITED STATES

A serious outbreak of encephalitis has been reported in some of the West North Central States since the first of July. On July 14, Dr. Maysil M. Williams, State Health Officer of North Dakota, reported the occurrence of 25 cases in that State from July 1 to 12.¹ Ten cases had been reported in the State during the first half of the year. Up to August 9, a total of 363 cases, with 41 deaths, had been reported in North Dakota.

Recent reports show an increasing number of cases of the disease in both South Dakota and Minnesota. In the latter State a high incidence has been reported in the western area adjacent to North Dakota.

The type of the disease has not yet been definitely determined, although there is some evidence that it is the western equine type. Medical officers and entomologists of the United States Public Health Service are collaborating with the State health departments in an

¹ Pub. Health Rep., July 18, 1941, p. 1849.

investigation of the outbreak, with particular reference to the type of the disease and the method of spread. Especial consideration is being given to the possibility of insect vectors and to the existence of some natural reservoir of the disease.

The numbers of cases reported in the 3 States to August 9 this year are as follows:

State	JanJune	Week ended-										
	1941	July 5	July 12	July 19	July 19 July 26		Aug. 9					
North Dakota South Dakota	10 0		² <u>2</u> 5	31	65	54 19	178 61					
Minnesota	2	3	1	11	39	35	65					

² July 1-12.

Incomplete figures show approximately 950 cases of encephalitis reported in the United States for the year to date, i. e., up to August 9, 1941, as compared with 541 cases for the period January to August, inclusive, in 1940. Of 108 cases reported in Washington State last year, 86 were stated to be the equine type of disease, and of 3 cases reported in Nevada, 1 case was recorded as equine encephalitis. No other States designated the type of the disease in their reports for 1940.

DEATHS DURING WEEK ENDED AUGUST 2, 1941

[From the Weekly Mortality Index, issued by the Bureau of the Census, Department of Commerce]

	Week ended Aug. 2, 1941	Correspond- ing week, 1940
Data from 88 large cities of the United States: Total deaths. A verage for 3 prior years. Total deaths, first 31 weeks of year. Deaths per 1,000 population, first 31 weeks of year, annual rate. Deaths under 1 year of age. A verage for 3 prior years. Deaths under 1 year of age, first 31 weeks of year. Deaths under 1 year of age, first 31 weeks of year. Deaths under 1 year of age, first 31 weeks of year. Deaths under 1 year of age, first 31 weeks of year. Deaths in force. Policies in force. Number of death claims. Death claims per 1,000 policies in force, annual rate. Death claims per 1,000 policies, first 31 weeks of year, annual rate.	8, 516 7, 714 270, 113 12. 2 585 502 16, 352 64, 399, 236 10, 739 8, 7 9, 9	8, 763 271, 522 12. 2 562 15, 690 65, 006, 071 11, 753 9. 5 10. 0

PREVALENCE OF DISEASE

No health department, State or local, can effectively prevent or control diseases without knowledge of when, where, and under what conditions cases are occurring

UNITED STATES

REPORTS FROM STATES FOR WEEK ENDED AUGUST 9, 1941 Summary

A total of 422 cases of poliomyelitis was reported during the current week as compared with 326 for the preceding week. The number of cases reported currently and the total number reported to date (first 32 weeks) are above the figures for corresponding periods of any year since 1937.

The South Atlantic and East South Central States reported 261, or 62 percent of the total cases for the current week. States reporting 10 or more cases for the current week (with last week's figures in parentheses) are as follows: Alabama 80 (49); Georgia 71 (71); Tennessee 31 (13); New York 30 (12); Ohio 27 (16); Pennsylvania 17 (15); South Carolina 16 (5); New Jersey 13 (5); Kentucky 13 (7); Florida 13 (27); Minnesota 12 (3); Indiana 12 (5); Maryland 11 (14); Michigan 10 (8); North Carolina 10 (0); and Mississippi 10 (9).

North Dakota reported 178 cases of encephalitis, Minnesota 65, and South Dakota 61. A total of 363 cases, with 41 deaths, has been reported in North Dakota up to August 9, and recent reports show increasing incidence in South Dakota, and in Minnesota, especially in areas near the North Dakota border.¹

Of 22 cases of Rocky Mountain spotted fever reported, only 3 occurred in the Mountain States. Four cases were reported in Maryland and 3 cases each in Illinois and North Carolina. Of 86 cases of endemic typhus fever, 34 cases occurred in Texas, 20 in Georgia, and 14 in Florida.

The death rate for the current week in 88 large cities is 10.6 per 1,000 population as compared with 11.9 for the preceding week and with a 3-year (1938-40) average of 10.1.

¹ See page 1661.

Telegraphic morbidity reports from State health officers for the week ended August 9, 1941, and comparison with corresponding week of 1940 and 5-year median

In these tables a zero indicates a definite report, while leaders imply that, although none were reported, cases may have occurred.

			_										
	1	Diphth	eria		Influenza			Measle	8	M me	Meningitis, meningococcus		
Division and State	Wee	k ended	1 Me-	Week	ended	Me-	Week	ended	Me-	Weel	r ende	d Me-	
	Aug. 9, 1941	Aug. 10, 1940	dian 1936- 40	Aug. 9, 1941	Aug. 10, 1940	dian 1936- 40	Aug. 9, 1941	Aug. 10, 1940	dian 1936– 40	Aug. 9, 1941	Aug. 10, 1940	dian 1936- 40	
NEW ENG.													
Maine	-		S S	8			. 19	2 3					
Vermont		i g		2			. 14	1 10	3 2			ğ ğ	
Rhode Island	: i	i č	j d	j				2 1	1 1			ŏ	
Connecticut	· 1	u c	ייו	l		•	. 34		7 18	5 1		0 0	
MID. ATL.	1					1 .	206	2 28	8 909				
New Jersey 3				2 2		2 3	66	12	50	i	i	i i	
Pennsylvania	. 8	8 10		8		-	. 156	5 11	s 118	2		4	
E. NO. CEN.	,						77	10	10	2			
Indiana		7		i i		2 3	i ii		3 6	ī	ģ		
Michigan 4				2		4	88	153	S 30 S 60	Ö			
Wisconsin	0	0		5	1	9 14	159	141	61	0	0	Ó	
W. NO. CEN.] .			1.							
Iowa ³		2	2			: 1	24	30		Ö			
Missouri 3		07	5	15		. 14	23					9	
South Dakota	5	İ	ī				3	i	2	Ŏ	ģ	Ō	
Kansas	1	6	3	1			25	14	6	ŏ		ö	
80. ATL.													
Delaware	0	1	0				2	l g	0	0	ļ	ļ	
Dist. of Col.	1	1	2			4	11	i	5	ő		ō	
Virginia ¹		9	10	74	46	4	48	37	31		0		
North Carolina 3	15	6	22	3	110		62	14	14	1	Ō	Ĩ	
Georgia 1 8	15	2	10	16	5		59	6		Ó	Ő	Ö	
Florida 1	1		4	4		1 1	9	4	2	. 0	1	0	
Kentucky	1	3	10	1		1 1	14	27	14	2	1	Ι.	
Tennessee 1	ļ	4	7	10	6	9	41	6	6	2	Ô	2	
Mississippi 1 4	2	ő	10	10	J		8	20	4	2			
W. SO. CEN.													
Arkansas	6	3	7	15	4	7	32	0	0	0	0	0	
Oklahoma	Ó	3	4	7	36	11	15	1	2	Ó	2	Ŏ	
Texas	25	15	20	348	151	39	106	52	33	U	2	3	
Montana 3	1	0	0		3		1	8	8	0	0	0	
Idaho	Ō	Ŏ	Ŏ				Ô	Ŏ	2	ŏ	ŏ	ŏ	
Colorado	6	4	4	····ii	6		23	2 4	8	0	0	Ŭ	
New Mexico	0	0	0	18	6		75	9	5	0	0	0	
Utah 34	Ő	ō	Ō	ĩ		·	10	12	12	ò	ŏ	ŏ	
PACIFIC	0						°			Ű			
Washington	0	2	1				1	4	12	0	0	0	
Oregon California	2	0 10	0 17	4	1	1	6 101	25 62	11 87	2	Ó	02	
Total	173	153	297	717	451	279	1, 821	1, 539	1, 111	32	20	42	
2 weeks	7, 460	8, 688	13, 402	599, 256	168, 789	151, 299	829, 197	227. 464	269. 437	1, 391	1, 138	2,114	
	,		.,	,	,			,		-, + !			

See footnotes at end of table.

Telegraphic morbidit	y reports from Sto	ste health officers	for the week of and 5-year n	nded August 9,
1941, and compari	son with correspon	ding week of 194		redian—Con.
		•		

	Po	liomye	litis	Sc	arlet fo	ver		Smallpo	ox Typhoid and typhoid fe		d para- ever	
Division and State	Week	ended	Me-	Week	ended	Me	Week	ended	Me-	Weel	ended	Me-
	Aug. 9, 1941	Aug. 10, 1940	dian 1936- 40	Aug. 9, 1941	Aug. 10, 1940	dian 1936- 40	Aug. 9, 1941	Aug. 10, 1940	dian 1936- 40	Aug. 9, 1941	Aug. 10, 1940	dian 1936- 40
NEW ENG.								· ·	Ì			
Maine New Hampshire Vermont. Massachusetts. Rhode Island Connecticut.	0 1 4 0 1	1 0 0 0 2	0 0 1 2 0 2	0 5 1 59 3 6	1 2 3 21 0 4	2 1 0 21 3 4			0 0 0 0 0	0 1 0 3 0 0 0		3 0 2 2 2
MID. ATL. New York ¹ New Jersey ³	30 13	71	9	61 19	65 18	71 14	0	0	0	30 4	16 10	16 6
Pennsylvania ³	17	1	3	41	57	57	0	0	.0	28	15	22
Ohio Indiana Illinois ³ Michigan ⁴ Wisconsin	27 12 8 10 1	16 41 4 31 0	9 1 8 24 2	50 8 35 35 34	33 11 53 46 35	61 23 64 76 31	0 1 1 0 0	0 0 0 0	0 0 2 1 1	10 3 16 2 0	8 6 7 6 0	16 7 35 12 2
w. NO. CEN. Minnesota Iowa ³ Missouri ³ North Dakota South Dakota Nebraska	12 0 0 0 0	1 19 12 0 2 1	4 2 3 0 1	9 5 12 0 6 5	12 6 13 3 4 5	19 9 15 5 5 5	0 0 1 0 1	5 0 0 3 1	3 2 1 0 1 1	1 5 9 1 1 0	3 5 19 5 0 1	1 5 19 0 0
Kansas	1	26	6	12	16	24	0	. 0	0	7	5	5
Delaware	0 11 2 3 1 10 16 71 13	0 1 20 2 0 0 0	0 1 0 2 1 2 2 0 2	0 9 3 15 7 18 1 6 2	0 10 1 9 11 16 1 7 4	1 10 2 9 12 25 1 6 2	0 0 0 0 0 0 0 0 0	0 0 0 0 1 0 0	0 0 0 0 0 0 0 0	0 11 0 9 5 13 8 27 6	0 3 8 6 3 10 23 4	1 12 23 22 13 15 26 3
E. SO. CEN.				1								
Kentucky Tennessee ¹ Alabama ¹ Mississippi ¹⁴	13 31 80 10	16 3 1 0	4 1 2 2	8 12 14 1	13 7 4 6	17 8 8 3	00000	0 0 0 0	0 0 0 0	17 12 13 15	16 11 14 14	43 30 18 13
W. SO. CEN. Arkansas Louisiana ¹ Oklahoma Texas ¹	3 2 1 4	1 5 5 12	0 1 2 1	2 0 9 7	6 1 4 9	6 5 6 17	0 0 0 0	0 0 1 0	0 0 0 0	15 8 4 28	30 14 19 72	23 19 21 87
MOUNTAIN Montana I	0	s	,	5	5	5	,	0	0	,	1	1
Idaho Wyoming ⁸ Colorado New Mexico Arizona	0 0 1 0 0	0 1 1 1 0	0 0 1 1 0	1 0 4 1 5	2 3 8 0	2 2 11 5 1	00000	0 0 12 0 0	0 0 0 0 0	0 0 5 2 2	0 1 1 2 0	1 1 2 6 1
Utah 3 4 Nevada	2 0	1	0	1 0.	5	5	0	U 		0	3 	1
PACIFIC Washington Oregon California	3 0 7	16 2 12	0 1 12	8 5 35	1 6 35	8 7 39	0 0 0	0 0 1	0 0 3	0 3 4	0 0 15	2 3 15
Total	422	278	261	585	582	798	5	25	25	330	384	608
32 weeks	2, 273	1, 681	1, 681	91, 371 1	18, 285	136, 453	1, 178	1, 952	7, 914	4, 143	4, 592	6, 602

See footnotes at end of table.

Telegraphic morbidity reports	from Stat	e health officers	for th	e week ended	August 9,
1941, and comparison	with corre	esponding week	of 19.	40—Continu	ed

	Whoopi	ng cough		Whoopi	ng cough	
Division and State	Week	ended	Division and State	Week ended		
	Aug. 9, 1941	Aug. 10, 1940		Aug. 9, 1941	Aug, 10, 1940	
NEW ENG. Maine New Hampshire Vermont Massachusetts Rhode Island Connecticut MID. ATL.	20 0 3 171 25 49	38 0 14 147 1 25	BO. ATL.—continued South Carolina 1 Georgia 1 3. Florida 1 E. SO. CEN. Kentucky Tonnessee 1	124 37 19 60 27	21 20 6 58 54	
New York ¹ New Jersey ³ Pennsylvania ³	272 99 216	843 91 42 6	Alabama ¹ Mississippi ¹ ⁴ W. 80. CEN.	4	9	
E. NO. CEN. Ohio Indiana. Illinois 3 Michigan 4 Wisconsin	435 21 204 309 233	243 18 152 287 83	Arkansas. Louisiana ¹ Oklahoma Texas ¹ MOUNTAIN	4 15 22 178	18 27 22 223	
W. NO. CEN. Minnesota Iowa ³ Missouri ³ North Dakota Nebraska Kansas	58 48 9 2 4 13 101	44 42 31 13 8 5 50	Montana ^a Idaho Wyoming ^a Colorado New Mexico Arizona Utah ^a 4 Nevada PACIFIC	26 46 10 110 4 14 33 4	2 7 8 9 8 5 5 5 5	
SO. ATL. Delaware	0 74 21 81 21	6 118 6 36 50	Washington Oregon California Total	81 14 293 3, 772	40 30 260 3, 302	
Delaware	0 74 21 81 21 158	6 118 6 36 50 146	Washington Oregon California Total 32 weeks	3, 77 143, 39	14	

¹ Typhus fever, week ended August 9, 1941, 86 cases as follows: New York, 1; Virginia, 1; South Carolina, 2; Georgia, 20; Florida, 14; Tennessee, 1; Alabama, 6; Mississippi, 2; Louisiana, 5; Texas, 34.
² New York City only.
³ Rocky Mountain spotted fever, week ended August 9, 1941, 22 cases as follows: New Jersey, 1; Pennsylvania, 1; Illinois, 3; Iowa, 1; Missouri, 2; Maryland, 4; District of Columbia, 2; West Virginia, 1; North Carolina, 3; Georgia, 1; Montana, 1; Wyoming, 1; Utah, 1.
⁴ Period ended earlier than Saturday.

WEEKLY REPORTS FROM CITIES

City reports for week ended July 26, 1941

This table summarizes the reports received weekly from a selected list of 90 cities for the purpose of showing a cross section of the current urban incidence of the communicable diseases listed in the table.

State and city	Diph- theria cases	Infl Cases	uenza Deaths	Mea- sles cases	Pneu- monia deaths	Scar- let fever cases	Small- pox cases	Tuber- culosis deaths	Ty- phoid fever cases	Whoop- ing cough cases	Deaths, all causes
Data for 90 cities: 5-year average Current week	74 49	25 29	10 6	725 70 8	280 228	284 223	4	845 307	57 31	1, 408 1, 285	
Maine: Portland	0		0	0	1	0	0	0	0	17	20
New Hampshire: Concord	0		0	0	0	0	0	0	0	0	4
Vermont: Barre	0		0	0	0	0	0	0	0	0	6
Massachusetts: Boston	2		0	40	9	16	0	4	0	34	174
Fall River	Ō		1	0	1	5	Ó	0	0	13	28
Worcester	ŏ		ŏ	1	3	5	ŏ	4	ŏ	3	54
Rhode Island: Providence	3		0	10	3	1	0	2	0	44	. 35
Connecticut: Bridgenort	0		0	8	0	0	0	0	0	1	27
Hartford	Ŏ		Ŏ	3	0	Ŏ	Ŏ	i	Ŏ	1	40
New Haven	v		Ů	.4	1	v			v	Ů	
New York: Buffalo	0		1	. 7	8	6	0	4	0	3	94
New York	5	4	1	52 11	35	25		68 0	5	115	1, 258
Syracuse	ŏ		ŏ	19	ō	Ž	Ŏ	ĭ	Ŏ	17	35
New Jersey: Camden	0		0	0	1	0	0	1	0	9	.21
Nowark Trenton	0			7	4	6 1		3	0	17	76 34
Pennsylvania:	ů	9		7	19	0		16	3	36	304
Pittsburgh	1	^	Ő	39	5	5	Ŏ	4	Ő	67	134
Reading	0		0	2	0	0	0	1	0	. 1	14
Ohio:				19	6			7	9	5	114
Cleveland	Ŏ	l i	Ŏ	5	4	13	Ŏ	7	Ő	81	187
Columbus Indiana:	0		0	9	Ů	1	0	5	U	16	74
Fort Wayne	0		0	1	1 9	0		0	0	4	19 101
South Bend	Ő		ŏ	2	õ	ŏ	ŏ	ŏ	ŏ	Ő	20
Terre Haute Illinois:	0		0	U	0	U	.0	1	. "	U	11
Chicago	10	1	1	16 14	93	25 1	0	46	0	66 0	620 17
Michigan:						10				75	054
Detroit Flint	0		U O	32	8 1	18	0	0	ŏ	15 9	200
Grand Rapids	0		0	11	1	. 3	0	0	0	5	32
Kenosha	0		0	4	0 1	1	0	0	0	0	8
Racine	0		0	10	ō	1	Ő	ő	0	8	12
Superior	0		0	2	1	1	0	0	0	8	6
Minnesota:	<u>م</u>			9		A	n	0	6	13	18
Minneapolis	Ő		Ŏ	1	1	2	Ŏ	1	ŏ	6	97
St. Paul Missouri:	0		0	3	7	1	0	U	U	14	73
Kansas City	0		0	8 0	37	3 0	0	4	0	13 0	102 29
St. Louis	2	2	ŏ	10	6	4	ŏ	ğ	ŏ	24	173
North Dakota: Fargo	0		0	0	0	0	0	1	0	5	14
Nebraska:	0		6	1	6	2	0	1	0	1	48
Kansas:										26	19
Wichita	ŏ		l öl	0	6	Ó	ŏ	il	ŏl	6	40

State and city	Diph-	Inf	luenza	Mea-	Pneu-	Scar- let	Small-	Tuber-	Ty- phoid	Whoop-	Deaths,
	Cases	Cases	Deaths	Cases	deaths	fever cases	cases	deaths	fever cases	cough cases	causes
Delaware: Wilmington	0		0	2	2	8	0	0	0	0	21
Maryland:		1 .		118	l ,	0		, ,	•	62	184
Cumberland	ŏ	1	ŏ	1	ó	ŏ	ŏ	6	ŏ	0 0	104
Frederick	Ő		Ō	1	Ó	Ō	Ó	Ó	Ŏ	Ō	4
District of Colum-											
Dia: Weshington	0		6	14	ĸ	8	0	18	0	12	166
Virginia:	·		, i		Ŭ		Ŭ				100
Lynchburg	1		0	11	0	0	0	0	0	4	10
Richmond			N N	2		8	0		3	U	39
West Virginia:	v		, v	-	ľ	v	v	Ň	v		. 0
Charleston	0		0	0	2	0	0	1	0	0	30
Wheeling	0		0	1	1	0	0	1	0	1	17
Raleigh	0		0	4	0	0	0	0	0	15	16
Wilmington	ŏ		ŏ	Ő	ŏ	ŏ	ŏ	ŏ	ŏ	24	13
Winston-Salem.	0		0	7	0	0	0	2	0	6	13
Cherleston	0	,	6	•	1	2	0	1			14
Georgia:	v	-	•	v	-	-	Ň	-	- 1	۰	14
Atlanta	0		0	0	2	1	0	5	1	0	77
Brunswick	0		0	.0	0	0	0	0	0	0	5
Savannan	v		۷I	10	z		•	٥	U I	2	27
Tampa	1		0	0	0	0	0	1	0	4	16
Tennessee:											
Memphis	0	5	0	4	2	0	0		2	19	88
Alabama:	U		U I	•	- 1	0	0	• •	2		49
Birmingham	0	1	0	3	1	1	0	8	2	8	57
Mobile	Ó		Ó	Ó	2	0	6	Ő	0	Ō	23
Astronom				· 1					- 1		
Little Rock	6		0	1	1	0	6	0	0	1	20
Louisiana:	ľ,		ľ ľ	-	•	Ľ ľ	Ů,	°	° I	-	
New Orleans	1		0	0	8	6	0	10	2	2	149
Shreveport	0		0	0	8	0	0	1	0	0	45
Dallas	0		0	8	0	3	0	1	3	1	65
Galveston	ŏ		ŏ	ŏ	i	ŏ	ŏ	ī	ŏ	ō	12
Houston	5		0	4	5	2	0	- 4 ·	3	2	70
San Antonio	0	1	U	1	10	1	0	5	0	4	67
Montana:											
Billings	0		0	0	0	0	0	0	0	0	11
Great Falls	0.		0	0	1	0	0	0	<u>o</u>	5	8
Missoula	Å.		N N			1					5
Idaho:	•		۳I	v I	° I	- 1	°	°	۳I	۳I	7
Boise	0 .		0	0	0	1	0	0	0	0	7
Colorado:				1.0							-
Pueblo	ŏ.		Ň	13	î	ŏ	- NI	8 I	il	2	70
Utah:	•		Ŭ,	•	-	, i	•	Ů	- 1		10
Salt Lake City_	0		0	2	0	1	0	0	1	32	24
Washington:											
Spokane	1-		N N		9	2	81		8	16	130
Tacoma	ŏŀ		ŏl	ŏĺ	1 l	í	ŏ	ŏI	ŏl	18	34 23
California:	- -		Ĩ	Ĭ	- 1	- 1	۳I	۳I	۳I	~	200
Los Angeles	2	7	0	23	6	9	0	14	0	. 69	331
Sacramento	1		N N	91	v		0	1	<u> </u>	4	28 160
Doll FIGHUSOU.	۳l	1	۳I	11	۳I	•	۲I	1	۳I	10	108
									_		

City reports for week ended July 26, 1941-Continued

State and city	Meni mening	ngitis, ;ococcus	Polio- mye-	State and city	Menin mening	Polio- mye-	
	Cases	Deaths	Cases		Cases	Deaths	Cases
Massachusetts: Springfield	0 4 1 0 0 0 0 0 0 1	0 1 0 0 0 0 0 0 0 0	1 4 0 1 8 2 4 1 1 1 4 1	District of Columbia: Washington	0 0 1 0 0 0 0 1 0 0	0 0 0 0 0 0 1 1 0 0	1 1 1 3 1 8 1 0 0 1 1 5
Baltimore	2	0	8		Ů	Ů	-

City reports for week ended July 26, 1941-Continued

Encephalitis, epidemic or lethargic.—Cases: Fargo, 17. Deaths: Fargo, 2; Seattle, 1. Pellagra.—Cases: Boston, 1; Charleston, S. C., 1; Atlanta, 1: Savannah, 4. Rabies in man.—Deaths: Nashville, 1. Typhus fever.—Cases: New York, 5; Tampa, 1; New Orleans, 1; Houston, 1; Los Angeles, 1.

PLAGUE INFECTION IN COLORADO, MONTANA, NORTH DAKOTA. AND WASHINGTON

Plague infection has been reported, under dates of July 29 and 30, and August 1, 1941, to have been found upon examination of specimens at the laboratory in San Francisco, Calif., as follows:

IN GROUND SQUIRRELS, MARMOTS, AND FLEAS IN SAN MIGUEL COUNTY, COLO.

In tissue from a ground squirrel, C. variegatus grammurus, shot July 16 on a ranch 3 miles west and 6 miles south of Placerville; in tissue from 2 marmots, Marmota flaviventris, shot July 17 at Horsefly Mesa, 27 miles south of Montrose; and in a pool of 30 fleas from 5 marmots of the same species, shot 3 miles west of Placerville on July 14.

IN FLEAS FROM GROUND SQUIRRELS IN BEAVERHEAD COUNTY, MONT.

In a pool of 77 fleas from 120 ground squirrels, C. columbianus, shot 3 miles west of Big Hole Battlefield.

IN FLEAS FROM GROUND SQUIRRELS IN DIVIDE COUNTY, N. DAK.

In 3 pools of fleas from ground squirrels, C. richardsonii, one a pool of 48 fleas from 44 ground squirrels shot July 11 on a ranch approximately 7 miles northeast of Crosby, another a pool of 54 fleas from 41 ground squirrels, shot 4½ miles north of Crosby on July 11, and the 402839°-41-----3

third a pool of 61 fleas from 49 ground squirrels shot July 12 on a ranch about 8 miles northeast of Crosby.

IN A GROUND SQUIRREL AND FLEAS FROM GROUND SQUIRRELS IN STEVENS COUNTY, WASH.

In tissue from a ground squirrel, *C. columbianus*, shot July 8 at a camp in Kauiksu National Forest, 15 miles northeast of Coleville, and in a pool of 175 fleas from 55 ground squirrels of the same species, shot July 12 along Twelve Mile Creek, 14 miles southeast of Coleville.

TERRITORIES AND POSSESSIONS

HAWAII TERRITORY

Plague (rodent).—Rats proved positive for plague infection have been reported from Kalopa Homesteads, Hamakua District, Island of Hawaii, T. H., as follows: 1 rat, July 8; and 2 rats, July 11.

FOREIGN REPORTS

CANADA

Provinces—Communicable diseases—Week ended July 5, 1941.— During the week ended July 5, 1941, cases of certain communicable diseases were reported by the Department of Pensions and National Health of Canada as follows:

Disease	Prince Edward Island	Nova Scotia	New Bruns- wick	Que- bec	Ori- tario	Mani- toba	Sas- katch- ewan	Al- berta	British Colum- bia	Total
Cerebrospinal meningitis. Chickenpox. Diphtheria.	2	3 17 9	1	5 55 18	6 159 1		2 55	3 45	37	22 427 33
Influenza		7			2		2		79	90
Measles		19	6	255 54	461 87	41	48 42	10 20	61 4	901 211
Pneumonia Poliomyelitis		3			1	1			4	8
Scarlet fever Tuberculosis Typhoid and paraty-	3	7 35	4	54 43	119 41	42	11 37	7 1	13 	219 170
phoid fever		1	1	23 275	1 119		2	3	22	25 422

Manitoba—Poliomyelitis.—Information received under date of August 8, 1941, states that 97 new cases of poliomyelitis were reported in the Province of Manitoba for the week ended August 8, making a total of 288 cases since July 1, 1941.

To date, the number of cases reported is more than half the total recorded during the epidemic of 1936, when there were 539 cases with 33 deaths.

No cases of poliomyelitis have been reported from the various military training camps in Manicoba.

SWITZERLAND

Communicable diseases—April 1941.—During the month of April 1941, cases of certain communicable diseases were reported in Switzerland as follows:

Disease	Cases	Disease	Cases
Cerebrospinal meningitis Chickenpoz. Diphtheria. German measles Influenza Measles Mumps Paratyphold fever	33 165 79 333 30 308 103 1	Poliomyelitis Scarlet fever Trachoma Tuberculosis Typhoid fever Undulant fever Whooping cough	2 291 2 304 4 16 190

REPORTS OF CHOLERA, PLAGUE, SMALLPOX, TYPHUS FEVER, AND YELLOW FEVER RECEIVED DURING THE CURRENT WEEK

NOTE.—Only those places are included which had not previously reported any of the above-named diseases, except yellow fever, during the current year. All reports of yellow fever are published currently. A cumulative table showing the reported prevalence of these diseases for the year to date is published in the PUBLIC HEALTH REPORTS for the last Friday of each month.

Cholera

India—Burma—Akyab.—A report dated July 2, 1941, stated that the town and port of Akyab, Burma, had been declared by the Government to be infected with cholera from June 27, 1941.

China.—During the period July 5 to 26, 1941, cases of cholera were reported in China as follows: Macao, 132; Shanghai, 27.

Plague

Palestine—Haifa.—Information dated July 29, 1941, reported the occurrence of 2 cases of human plague in Haifa during the week.

Indochina (French)—Cochinchina—Chaudoc.—During the period June 20-30, 1941, 17 fatal cases of plague were reported in Chaudoc.

Smallpox

The report of 1 case of smallpox in Santiago de Cuba, Cuba, for the week ended April 5, 1941 (Public Health Reports, May 9, 1941, p. 1038; May 30, 1941, p. 1188; June 27, 1941, p. 1348; and July 25, 1941, p. 1533) was an error. Later information states that this was a case of measles, and was erroneously recorded as smallpox.

Yellow Fever

Brazil.—Deaths from yellow fever have been reported in Brazil as follows: Para, June 2, 1; Bahia, June 12, 1; June 13, 1.

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