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TULARÆMIA Francis 1921.1

IV. TRANSMISSION OF TULARÆMIA BY THE BEDBUG, CIMEX LECTULARIUS.

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The experiments here reported show that the bedbug Cimex lectularius, which commonly infests beds and bites human beings, is capable of transmitting tularæmia from an infected to a healthy white mouse. Two distinctly different methods of transmission were successful. In one method we followed the usual procedure of first allowing the insects to feed on an infected animal and then to feed on a healthy animal. In the other method we followed the first half of the usual procedure, but the second half was reversed; i. e., the mouse ate the infected bug instead of the infected bug biting the mouse. the usual method, transmission was successful in 10 experiments in which the intervals which elapsed between biting the infected mice and biting the healthy mice were a few seconds, 18 hours, 7 days, 15 days, and 71 days. Following the unusual method, transmission was successful in 55 experiments in which the intervals which elapsed between the bug's biting the infected mouse and the mouse's eating the infected bugs varied uniformly from 6 hours to 100 days; this latter limit promises to be still further extended and to keep pace with the natural length of life of a bedbug.

METHODS EMPLOYED.

White mice and guinea pigs were the experimental animals used. Infection was due to strain (S) of Bacterium tularense isolated in 1920 from a human case of tularæmia in Utah. The bedbugs were collected from cracks in the wooden cages in which the laboratory stock of guinea pigs and white mice are bred. The bugs were kept on hand from one to three weeks before attempting to infect them. They were confined at all times to test tubes. They sucked an infecting meal of blood either from the tail of an inoculated white

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mouse, the tail being poked into the tube through a hole in the gauze which covered the mouth of the tube, or they sucked through the gauze, the tube being inverted and held on the infected mouse's abdomen.

The white mice which were used for infecting the bugs were inoculated subcutaneously on the back with the heart's blood of a dead infected mouse.

Bugs were applied to a sick mouse between 72 and 96 hours after inoculation, since four days was the maximum life of a white mouse inoculated subcutaneously.

The tubes containing the bugs were always kept at room temperature, which during the summer was practically that of the outside air, but after cool weather arrived in October it was that of the ordinary steam-heated laboratory, which averaged 22° C.

A tube contained not over 40 bugs and was supplied with a strip of filter paper on which the bugs rested and deposited their feces. The white mice were kept in glass battery jars, one mouse to each jar.

TRANSMISSION FOLLOWING BEDBUG BITES.

Ten white mice died with typical lesions of tularæmia following bedbug bites. Their histories are summarized at the head of the table and given more in detail in their appropriate places in the body of the table. Experimental transmission by biting is largely dependent upon interrupted feeding. An engorged bug will not bite. A partially engorged bug will bite. Forced interruption of a bug's meal of blood on the infected animal conduces to a completion of that meal on the healthy animal. The shorter the interruption the greater the likelihood of transmission. The lengths of interruption in our experiments were (1) a few seconds, (2) 18 hours, (3) 48 hours, (4) no interruption.

(1) Interruption of a few seconds.—The bugs were allowed to feed 2½ minutes on the infected mouse and then, after only a second's interval, they were allowed to feed for 2½ minutes on the healthy mouse; they fed in this manner alternately on the infected and healthy mice during 15 minutes, thus permitting of three insertions of the proboscis in each of the two animals. It is not probable that all of the bugs made six insertions, but certainly each bug made multiple insertions. In this method transmission was probably accomplished purely by a grossly contaminated proboscis.

During an experiment the infected mouse and the healthy mouse were tied to a board, abdomens upward. The abdomen of the infected mouse was shaved three days previously, but the abdomen of the healthy mouse was unshaved and unclipped. The bugs were contained in a glass tube, the mouth of which was covered with gauze. The mouth of the tube was held alternately in contact with

the two abdomens, the bugs biting through the gauze. No evidences of defecation were noted on the hair of the healthy mice.

Results: Five sets of bugs, averaging 25 bugs to each set, bit four infected mice, and then, after an interval of only a few seconds, bit 5 healthy mice according to the above-mentioned interrupted method of feeding. Transmission occurred in all instances. The intervals which elapsed between biting the healthy mice and their deaths from tularæmia were $3\frac{1}{2}$, $4\frac{1}{2}$, $4\frac{1}{2}$, $5\frac{1}{2}$, and $5\frac{1}{2}$ days, the average being $4\frac{1}{2}$ days. There were no unsuccessful attempts at transmission when the period of interruption was only a few seconds.

(2) Interruption of 18 hours.—The bugs were allowed to feed on the abdomen of the infected mouse for only 2½ minutes, after which they were set aside for 18 hours, at the end of which time they were applied for 12½ minutes in contact with the tail of a healthy mouse and allowed to feed to full engorgement. For this purpose the tail of the healthy mouse was poked through a hole in the gauze which covered the mouth of the tubes. No evidences of defectaion were noted on the tails of the healthy mice while being bitten by the bugs.

Results: Three sets of bugs, averaging 70 bugs to each set, fed for $2\frac{1}{2}$ minutes on three infected mice, and then after an interval of 18 hours, fed to engorgement on three healthy mice. Transmission occurred in two instances, the intervals which elapsed between biting the healthy mice and their deaths from tularemia being $4\frac{1}{2}$ and $5\frac{1}{2}$ days. The mouse bitten by the third set of bugs remained well. (See table, Lots 262, 263, and 264.)

- (3) Interruption of 48 hours.—Five sets of bugs, averaging 54 bugs to each set, fed for 2½ minutes on 5 infected mice and then, after an interval of 48 hours, fed to full engorgement on 5 healthy mice. Transmission occurred in no instance. (See table, Lots 272 to 284.)
- (4) Noninterrupted feeding.—This method of transmission varied from the foregoing methods in that the bugs' first meal was a full meal. They were allowed to feed on the abdomen or tail of the infected mouse to full engorgement without interruption.

Having sucked one full meal of infected blood, all subsequent feedings took place on healthy mice and were for the purpose of determining whether infection of healthy mice would follow the bites of infected bugs.

The second, third, and fourth feedings took place approximately on the third, sixth, and tenth days after the infecting meal. Subsequent feedings occurred approximately every 10 days throughout the life of the bugs.

While feeding on healthy mice, the bugs were applied in contact with the tails of these mice for periods of one hour, thus not only permitting them to feed to engorgement, but encouraging them to defecate on the tails. For this purpose the tail of the healthy mouse was kept poked for one hour into a slender glass tube containing the bugs. The bugs avoid resting on the tail of the mouse if possible; they even prefer glass to a mouse's tail as a resting place. In order to compel them to rest on the tail, the caliber of the glass tube was just enough larger than the diameter of the tail to permit the passage of the bugs along the tail, and, moreover, the tube was held in a vertical position so that the bugs were forced to rest on the tail of the mouse. The result was that the bugs deposited their feces principally on the tail so that the points of biting must have become overlaid with bug feces in some instances.

Results: Ten lots of bugs (Nos. 230 to 258), varying from 16 to 140 bugs, but averaging 88 bugs to the lot, fed to engorgement on 10 infected mice, and then, after intervals of from 3 to 110 days, fed to engorgement during one hour on the tails of 23 healthy mice, feeding an average of five times on each healthy mouse. Transmission occurred in 3 out of 23 mice. The intervals which elapsed between biting the infected mice and biting the 3 healthy mice were 7, 15, and 71 days, respectively, and the intervals between biting the 3 healthy mice and their deaths from tularæmia were 6, 5, and 7 days, respectively; the number of bugs employed in the three transmissions were 28, 24, and 14, respectively.

TRANSMISSION OF TULARÆMIA DUE TO EATING INFECTED BEDBUGS.

Transmission experiments with bedbugs and white mice which do not take into account the mouse's habit of eating bedbugs will be full of errors. White mice readily attack and eat living bedbugs; they eat dead bedbugs with equal readiness. If the bugs are infected with Bacterium tularense, the mouse is almost certain to die of tularæmia. One of our white mice confined with 45 infected living bugs within a small glass jar free from bedding or other hiding places for the bugs, ate all the bugs in less than an hour, being at the rate of one bug a minute. A mouse ate 15 living infected bugs over night, the mouse and bugs being loose in a glass battery jar containing bedding composed of coarse screenings of sawdust. Two healthy mice, dropped into a large jar containing cut hay and small wooden boxes and a plentiful supply of bugs which had been infected 21 days previously, died of tularæmia 31 and 51 days after entering the jar. Presumably the mice contracted the infection from eating the bugs rather than from the bites of the bugs.

Experiments in other fields of research designed to demonstrate transmission by fleas often have permitted the unrestricted presence of rodents and fleas in the same container. Such a procedure in bug-mouse experiments would leave the experimenter totally in the dark on the question of whether transmission resulted from the bedbug's biting the mouse or the mouse's eating the bedbug.

We conducted 72 bug-eating experiments, 55 of which were successful. The infected bugs came from 10 lots (Nos. 230 to 258). These bugs had fed once to engorgement on infected mice and were fed subsequently every 3 to 10 days on healthy mice. These healthy mice served to test the infectiveness of the bug bites. Some of the bugs were found dead every morning, while others were dying and consequently almost motionless. These infected dead or dying bugs were fed to mice, thereby infecting the mice. The same lots of bugs, therefore, were used in the bug-biting and bug-eating experiments.

Bugs which were to be eaten were dropped, together with a white mouse, into a bottomless glass cylinder 4 inches in diameter which rested on a blotter; the purpose of the blotter was to absorb the urine. The mouse remained until he had eaten the bugs, which in some instances was overnight. The mouse was then transferred to a glass battery jar, commonly used as a mouse cage, each mouse occupying a separate jar.

Summary.—Seventy-two white mice ate dying or dead bugs from each of 10 lots of infected bugs. Of this number, 55 died from tularæmia and 17 remained well. The number of bugs offered to and eaten by each mouse varied from 1 to 10, the average being 3. The average length of time which elapsed between eating infected bugs and death of the mice from tularæmia was $4\frac{1}{2}$ days. Of 20 mice which each ate one infected bug, 14 died acutely from tularæmia. The average length of time from the date of infection of the 14 bugs until they were eaten was 65 days. Three white mice which each ate a bug infected 100 days previously died five, four, and five days later from tularæmia.

Bacterium tularense suffers no apparent diminution of virulence by reason of long residence in the bedbug.

It is the intention to continue these eating experiments throughout the life of the bugs, which promises to be long.

INFECTIVITY OF FRESH BEDBUG FECES.

While transmission experiments with our 10 lots of infected bugs were in progress, the infectivity of the bug feces was frequently tested. The strips of filter paper on which the bugs habitually rested served also for the deposition of feces. When the strips were renewed, the old ones were soaked in saline solution, rubbed in a mortar, strained through gauze, and the suspension was either injected subcutaneously into a guinea pig or mixed with corn meal and fed to a white mouse. From each of ten lots of infected bugs two samples of feces were collected about 10 days apart and between 17 and 58 days after the date of infection of the bugs. Both samples from a given lot of bugs were fed at the time of collection to the same mouse. Ten mice

which were thus fed with fresh feces from 10 lots of infected bugs remained well.

Forty-one samples of feces, representing about three samples from each of the 10 lots of infected bugs, were injected, while fresh, subcutaneously into 41 guinea pigs. At least one sample in each of the 10 lots was collected 35 days or more after the date on which bugs were infected, and in one instance the sample was collected 120 days after the date of infection. The 41 guinea pigs all died acutely from tularæmia.

The two sets of experiments show in marked contrast the invariable susceptibility of the guinea pig's subcutaneous tissue to the infected bedbug feces in question, and the invariable resistance of the mouse's stomach to the same material.

Summary.—The fresh feces of bedbugs which were infected with Becterium tularense by sucking the blood of infected white mice and which were fed every 10 days thereafter on the blood of healthy white mice, contained virulent organisms of this infection at all times and did so up to 120 days after the date of infection of the bugs, and probably will do so throughout the life of the bugs.

INFECTIVITY OF DRIED FECES OF BEDBUGS.

Feces deposited by our 10 lots of infected bugs on filter papers, between 46 and 75 days after the dates of infection of the bugs, were subsequently set aside and allowed to remain on those filter papers in a dried condition at an average room temperature of 20° C. unexposed to direct light. At the end of 20 days of drying, the filter papers were soaked in saline solution, ground in a mortar, and the pooled suspension of feces was injected subcutaneously into two guinea pigs, causing their deaths within six days with typical lesions of tularæmia.

Feces deposited on filter papers by 8 lots of infected bugs (see table, Lots 262 to 284) between 18 and 70 days after the dates of infection of the bugs, were subsequently kept under the conditions described above for 25 days, at the end of which time the pooled suspension was injected subcutaneously into a guinea pig, causing its death 8 days later with typical lesions of tularæmia.

TRANSMISSION TO GUINEA PIGS.

The foregoing experiments relate to transmission to white mice. A few attempts were made on transmission to guinea pigs by infected bedbugs. Bugs from 6 of our 10 lots of infected bugs bit 6 guinea pigs, respectively. The guinea pigs were exposed to the bugs under the following conditions.

A healthy guinea pig and infected living bugs were dropped into a bottomless glass cylinder, 6 inches in diameter, which rested on a blotter; the purpose of the blotter was to absorb the urine. Guinea pig and bugs remained in the cylinder overnight. The guinea pigs never showed any tendency to eat the bugs. The bugs were plainly seen to feed to engorgement on the feet of the guinea pigs. They never crawled over the guinea pig's feet but approached only close enough to insert their outstretched proboscides.

The bugs were counted into the cylinder in the evening and counted out in the morning. The evening and morning counts always tallied, with one exception. The guinea pigs all remained well, with the exception of one (see Lot 258). This one was the guinea pig in whose cylinder we failed to recover one bug one morning, 47 having been counted in and only 46 having been counted out. In this case the blotter had been chewed to small bits. The presumption is that the guinea pig unintentionally swallowed one infected bug in chewing its blotter, because it died 7 days later from tularæmia and showed five typical cervical buboes and a chain of typical lymph glands in the mesentery, in addition to the ordinary typical lesions of the spleen and liver.

In order to test the susceptibility of guinea pigs to infection by eating, five lots of 2 bugs each were fed to engorgement on 5 infected mice and 24 hours later were fed to five guinea pigs, respectively, each pair of bugs being ground in a mortar with normal saline solution and applied to a small piece of bread which the guinea pigs readily ate. One of the five guinea pigs died seven days after eating the two infected bugs and showed the typical lesions of tularæma; the other four remained well. This is taken as strong evidence that the one guinea pig which died from tularæmia after confinement with 46 infected bedbugs contracted its infection from the accidental ingestion of the one missing bug rather than from the bites of the bugs.

In another experiment pooled fresh feces from our 10 lots of infected bugs were fed on bread to a guniea pig with negative results.

Summary.—An average of 35 infected bedbugs from each of 6 of our 10 lots of infected bugs were exposed with 6 healthy guinea pigs, respectively, during approximately the third, sixth, and tenth nights after the date of infection of the bugs. Although freely bitten by the bugs, only 1 of the guinea pigs contracted tularemia, and in this instance infection is believed to have taken place from the ingestion of a missing infected bug (see Lot No. 258).

Acknowledgment: The determination of specimens of Cimex lectularius were made by Dr. H. E. Ewing, of the Burcau of Entomology, Department of Agriculture.

GENERAL SUMMARY.

The common bedbug *Cimex lectularius* transmitted tularæmia from infected to healthy mice in 10 instances, in which the intervals which elapsed between biting the infected and biting the healthy mice were a few seconds, 18 hours, 7 days, 15 days, and 71 days. The exact parts played by bites and by feces in the 10 transmissions are impossible of determination.

White mice readily eat living and dead bugs.

White mice which eat infected bugs usually contract tularæmia. Of 20 white mice which each ate one infected bug, 14 died acutely from tularæmia. The average length of time from the date of infection of the 14 bugs until they were eaten was 65 days. Three white mice which each ate a bug infected 100 days previously died 5, 4, and 5 days later from tularæmia.

Guinea pigs apparently do not eat bugs intentionally.

Guinea pigs bitten by infected bugs failed to contract tularæmia, with one exception; in the latter instance the guinea pig probably ate one infected bug unintentionally and thereby contracted the infection.

The fresh feces of bedbugs which were infected with *Bacterium tularense* by sucking the blood of infected white mice, and which were fed every 10 days thereafter on the blood of healthy white mice, contained virulent organisms of this infection at all times and did so up to 120 days after the date of infection of the bugs.

Feces of infected bedbugs deposited on filter papers at least 46 days after the dates of infection of the bugs and subsequently dried for 20 days contained virulent organisms of *Bacterium tylarense* at the end of that time.

In spite of the last two preceding paragraphs, the fresh feces of infected bugs have always failed to infect white mice or guinea pigs which ate those feces.

Bacterium tularense suffered no apparent diminution of virulence by reason of long residence in bedbugs. This virulence was manifested by acute death from tularæmia within seven, five, five, and six days, respectively, in cases of—

- (1) A mouse which was bitten by bugs infected 71 days previously;
- (2) A mouse which ate a bug infected 100 days previously;
- (3) A guinea pig which was infected with fresh feces of bugs infected 120 days previously; and
- (4) A guinea pig injected with bug feces which had been deposited on filter paper at least 46 days after the date of infection of the bugs and which had subsequently dried on filter paper at an average room temperature of 20° C. for 20 days.

Transmission of tularamia by beddugs of the species Cimex lectularius.

	Remarks.	86 Lot No. 246. 86 Lot No. 246. 86 Lot No. 246. 86 Lot No. 220. 86 Lot No. 220. 86 Lot No. 220. 86 Lot No. 230. 86 Lot No. 246. 86 Lot No. 246. 86 Lot No. 246.	Reported above.	Reported above.
species comes iccumulas.	Transmission following bug blee: Longth of time after date of infection of bugs when bugs were allowed to bite a healthy white mouse or guinea pig. "Posttive" means death from tularsemia.	Few seconds; 25 bugs bit abdomen of mouse. Positive. Few seconds; 30 bugs bit abdomen of mouse. Positive. Few seconds; 32 bugs bit abdomen of mouse. Positive. Few seconds; 30 bugs bit abdomen of mouse. Positive. Few seconds; 30 bugs bit abdomen of mouse. Positive. 18 hours; 62 bugs bit tail of mouse. Positive. 18 hours; 62 bugs bit tail of mouse. Positive. 7 days; 28 bugs bit tail of mouse. Positive. 11 days; 34 bugs bit tail of mouse. Positive. 11 days; 44 bugs bit tail of mouse. Positive.	Few seconds; 25 burs bit abdomon of mouse 60. Positive. 8 days; 16 bugs bit tail of mouse 5. Negative. 15 days; 30 bugs bit tail of mouse 5. 11 days; 30 bugs bit tail of mouse 5. 18 days; 28 bugs bit tail of mouse 5. 18 days; 30 bugs bit tail of mouse 6. 18 days; 30 bugs bit tail of mouse 6. 27 days; 30 bugs bit tail of mouse 6. 27 days; 18 bugs bit tail of mouse 6. 36 days; 18 bugs bit tail of mouse 6. 36 days; 14 bugs bit tail of mouse 6. 34 days; 14 bugs bit tail of mouse 6. 34 days; 3 bugs bit tail of mouse 6. 34 days; 3 bugs bit tail of mouse 6. 34 days; 3 bugs bit tail of mouse 6. 35 days; 3 bugs bit tail of mouse 6. 36 days; 3 bugs bit tail of mouse 6.	Few seconds; 30 bugs bit abdomen of mouse 8. Positive. 2 days; 20 bugs bit tail of mouse 14. Negative. Positive. 2 days; 20 bugs bit tail of mouse 14. Negative. B days; 30 bugs bit tail of mouse 14. 14 days; 35 bugs bit tail of mouse 14. 14 days; 25 bugs bit tail of mouse 14. 18 days; 22 bugs bit tail of mouse 14. 18 days; 22 bugs bit tail of mouse 14. 18 days; 30 bugs bit tail of mouse 15. 18 days; 30 bugs bit tail of mouse 15. 18 days; 30 bugs bit tail of mouse 15. 18 days; 30 bugs bit tail of mouse 15. 18 days; 30 bugs bit tail of mouse 15. 18 days; 34 bugs bit feet of mouse 15. 18 days; 45 bugs bit feet of guinea pig 2. Negative. 18 days; 43 bugs bit feet of guinea pig 2. 18 days; 43 bugs bit feet of guinea pig 2.
Theremoseon of them while of occupated the species Cines returning	Transmission due to eating injected bugs. Length of time after date of infection of bugs when dying or dead bugs were eaten by a white mouse. "Positive" means death from tularæmis.	res.' following bedbug bites are assembled in brief at the right or ready reference.)	6 hours; fed 2 dying bugs to mouse 82. Positive. 5. 6 hours; fed 2 dying bugs to mouse 83. Positive. 2 days; fed 6 dead bugs to mouse 84. Positive. 4. 6 days; fed 5 dying bugs to mouse 88. Positive. 4. 6 days; fed 5 dead bugs to mouse 88. Positive. 55 days; fed 5 dead bugs to mouse 87. Positive. 67. 67 days; fed 2 dead bugs to mouse 87. Positive. 74 days; fed 1 dead bug to mouse 99. Positive.	6 hours; fed 2 dying bugs to mouse 10. Positive. 6 hours; fed 2 dying bugs to mouse 11. Negative. 6 hours; fed 2 dying bugs to mouse 12. Negative. 6 hours; fed 2 dying bugs to mouse 13. Negative. 13 days; fed 9 dead bugs to mouse 14. Negative. 22 days; fed 10 dying bugs to mouse 15. Negative. 37 days; fed 10 dying bugs to mouse 16. Negative. 84 days; fed 6 dead bugs to mouse 10. Negative.
	Number of days Number of days Number of days after date of infec- tion of bugs when their fresh feess were (I) infected into a guinea pig or (2) fed to a white mouse. "Positive" means death from tulatre- mia.	(The 10" Positives" following bed of the table for ready reference.)	(1) Feces injected: 13 days. Positive. 23 days. Positive. 43 days. Positive. (2) Feces fed: 22 days. Negative. 35 days. Negative.	(1) Feces injected: 20days. Positive. 30days. Positive. (12) Feces fed: 22 days. Negative. 35 days. Negative.
	Various lots of bugs, each lot was infected by biting a separate mouse.		Lot No. 244. 80 bugs infected Sept. 15, 1921.	Lot No. 248. 140 b u g s infected Sept. 19; 1921.

Transmission of tular mia by bedbugs of the species Cimex lectularius—Continued.

Remarks.	Reported above.	Reported above.	Reported above.
Transmission following bug bites: Length of time after date of infection of bugs when bugs were allowed to bite a healthy white mouse or guines pig. "Positive" means death from tularsamia.	Few seconds; 31 burs bit abdomen of mouse 17. Positive. 3 days; 45 bugs bit feet of ginnea pig 3. Negative. 6 days; 43 bugs bit feet of guinea pig 3. 14 days; 41 bugs bit feet of guinea pig 3. 14 days; 35 bugs bit tail of mouse 22. 17 days; 38 bugs bit tail of mouse 22. 26 days; 14 bugs bit tail of mouse 22. 46 days; 10 bugs bit tail of mouse 25. 46 days; 10 bugs bit tail of mouse 25. 46 days; 10 bugs bit tail of mouse 25. 46 days; 4 bugs bit tail of mouse 25. 46 days; 4 bugs bit tail of mouse 25. 46 days; 4 bugs bit tail of mouse 25. 46 days; 4 bugs bit tail of mouse 25. 46 days; 4 bugs bit tail of mouse 25. 46 days; 4 bugs bit tail of mouse 25.	Few seconds: 30 bugs bit abdomen of mouse 33. Positive. 5 days; 45 bugs bit fail of mouse 34. Negative. 7 days; 41 bugs bit fail of mouse 34. Negative. 10 days; 80 bugs bit fail of mouse 34. 16 days; 82 bugs bit fail of mouse 35. Negative. 20 days; 28 bugs bit fail of mouse 35. Negative. 27 days; 28 bugs bit fail of mouse 36. Negative. 37 days; 17 bugs bit fail of mouse 38. 37 days; 19 bugs bit fail of mouse 38. 57 days; 9 bugs bit fail of mouse 36. 77 days; 9 bugs bit fail of mouse 36. 77 days; 9 bugs bit fail of mouse 36. 77 days; 7 bugs bit fail of mouse 36. 77 days; 7 bugs bit fail of mouse 36.	36 hours; 47 bugs bit feet of guinea pig 6. Positive. 3 days, 40 bugs bit feet of guinea pig. 6. Tositive. 7 days; 28 bugs bit tall of mouse 6s. Positive. 16 days; 20 bugs bit tall of mouse 6s. Positive. 19 days; 20 bugs bit tall of mouse 6s. Roguive. 26 days; 20 bugs bit tall of mouse 6s. Negative. 40 days; 18 bugs bit tall of mouse 6s. 15 days; 38 bugs bit tall of mouse 6s.
Transmission due to eating infected bugs. Length of time after date of infection of bugs when dying or dead bugs wereasten by a white mouse. "Positive" means death from tularæmia.	nours; fed 2 dying bugs to mouse 18. Positive. nours; fed 2 dying bugs to mouse 30. Positive. nours; fed 2 dying bugs to mouse 20. Positive. days; fod 8 dead bugs to mouse 21. Positive. days; led 5 dying bugs to mouse 22. Positive. days; fed 5 dead bugs to mouse 23. Positive. days; fed 5 dead bugs to mouse 24. Positive. days; fed 1 dead bug to mouse 88. Positive. days; fed 1 dead bug to mouse 102. Negative. days; fed 1 dead bug to mouse 102. Negative.	1 day; fed 2 dying bugs to mouse 28. Positive, I day; fed 2 dying bugs to mouse 27. Positive, 1 day; fed 2 dying bugs to mouse 28. Positive, 27 days; fed 4 dead bugs to mouse 29. Positive, 30 days; fed 5 dying bugs to mouse 30. Positive, 35 days; fed 6 dead bugs to mouse 31. Positive, 44 days; fed 6 dead bugs to mouse 31. Positive, 47 days; fed 1 dead bugs to mouse 32. Positive, 47 days; fed 1 dead bug to mouse 38. Positive, 77 days; fed 1 dying bug to mouse 98. Positive.	6 hours; fed 2 dying bugs to mouse 49. Positive. 12 hours; fed 2 dying bugs to mouse 50. Positive. 27 days; fed 2 dying bugs to mouse 51. Positive. 28 days; fed 8 dead bugs to mouse 52. Positive. 28 days; fed 6 dead bugs to mouse 53. Positive. 23 days; fed 2 dead bugs to mouse 54. Positive. 31 days; fed 2 dead bugs to mouse 55. Positive. 38 days; fed 1 dying bug to mouse 56. Positive.
Infectivity of bug fees: Number of days after date of infec- tion of bugs when their fresh fees were (1) Injected into a guinea pig or (2) fed to a white mouse. "Positive" means death from tulatæ- mia.	(1) Focos injected: 61 I7 days. Positive. 61 26 days. Positive. 63 days. Positive. 63 days. Positive. 32 I7 days. Negative. 33 days. Negative. 33 days. Negative. 56 67	(1) Foces injected: 16 days. Positive 27 days. Positive 37 days. Positive (2) Feter ici- 17 days. Negative 27 days. Negative	(1) Feces injected: 7 days. Fostitve 24 days. Positive 41 days. Positive (2) Feces fed: 19 days. Negative 29 days. Negative
Various lots of bugs; each lot was infected by biting a separate mouse.	No. 250. 90 u g s infected pt. 23, 1921.	Lot No. 252. 86 bugs infected Sept. 22, 1921.	Lot No. 238. 130 bugs infected Sopt. 20, 1921.

		Reported above.	
Negative. Negative.	Negative. Negative. Negative.	Positive. Negative.	1. Negative. 1. Signature. 1. Negative. Negative.
19 days; 47 bugs bit tail of mouse 66. 29 days; 35 bugs bit feet of mouse 66, 40 days; 28 bugs bit feet of mouse 66, 50 days; 25 bugs bit teld of mouse 66, 60 days; 16 bugs bit tail of mouse 97. 70 days; 16 bugs bit tail of mouse 97.	28 days; 38 bugs bit tall of mouse 1. 30 days; 38 bugs bit tall of mouse 1. 26 days; 38 bugs bit tall of mouse 2. 36 days; 28 bugs bit tall of mouse 2. 38 days; 38 bugs bit tall of mouse 2. 44 days; 38 bugs bit tall of mouse 3. 50 days; 37 bugs bit tall of mouse 3. 50 days; 17 bugs bit tall of mouse 3. 66 days; 16 bugs bit tall of mouse 3. 66 days; 16 bugs bit tall of mouse 3. 67 days; 12 bugs bit tall of mouse 3. 67 days; 12 bugs bit tall of mouse 3. 68 days; 12 bugs bit tall of mouse 3. 68 days; 12 bugs bit tall of mouse 3.	1 day; 66 bugs bit tail of mouse 4. 3 days; 34 bugs bit tail of mouse 4. 6 days; 35 bugs bit tail of mouse 4. 20 days; 27 bugs bit tail of mouse 4. 31 days; 25 bugs bit tail of mouse 4. 31 days; 25 bugs bit tail of mouse 4. 42 days; 21 bugs bit tail of mouse 4. 42 days; 12 bugs bit tail of mouse 4. 42 days; 13 bugs bit tail of mouse 4. 61 days; 18 bugs bit tail of mouse 4. 71 days; 18 bugs bit tail of mouse 4. 71 days; 14 bugs bit tail of mouse 4. 80 days; 14 bugs bit tail of mouse 5. 80 days; 14 bugs bit tail of mouse 5.	2 days; 14 bugs bit feet of guines pig 1. Negar days; 14 bugs bit feet of guines pig 1. 10 days; 13 bugs bit feet of guines pig 1. 13 days; 13 bugs bit feet of guines pig 1. 13 days; 13 bugs bit fiel of mouse 7. Negative. 21 days; 7 bugs bit tail of mouse 7. Negative. 40 days; 5 bugs bit tail of mouse 7. 50 days; 5 bugs bit tail of mouse 7. 50 days; 4 bugs bit tail of mouse 7. 70 days; 4 bugs bit tail of mouse 7. 70 days; 8 bugs bit tail of mouse 7. 70 days; 4 bugs bit tail of mouse 7. 70 days; 4 bugs bit tail of mouse 7. 70 days; 4 bugs bit tail of mouse 7.
36 days; fed 1 dying bug to mouse 57. Positive. 36 days; fed 1 dying bug to mouse 58. Positive. 37 days; fed 1 dead bug to mouse 59. Negative. 40 days; fed 3 dead bugs to mouse 61. Positive. 54 days; fed 1 dying bug to mouse 92. Positive.	16 days; fed 45 living bugs to mouse 78. Positive. 28 days; fed 20 living bugs to mouse 79. Positive. 35 days; et 4 dying bugs to mouse 80. Positive. 37 days; fed 5 dead bugs to mouse 81. Positive. 17 days; fed 6 dead bugs to mouse 82. Positive. 83 days; fed 1 dead bug to mouse 82. Positive. 100 days; fed 1 dying bug to mouse 93. Positive. 160 days; fed 1 dying bug to mouse 94. Positive. 163 days; fed 1 doad bug to mouse 105. Negative.	ays: fed 5 dylng bugs to mouse 77. Negative. ays: fed 1 dead bug to mouse 85. Negative. days: fed 1 dead bug to mouse 103. Positive. days; fed 1 dead bug to mouse 104. Positive.	15 days; fed 3 dead bugs to mouse 75. Positive. 51 days; fed 1 dead bug to mouse 76. Positive.
•	(1) Feess injected: 38 days. Positive, 38 days. Positive, 38 days. Positive, 44 days. Positive, 44 days. Positive, 71 days. Positive, 90 days. Positive, 100 days. Positive, 110 days. Positive, 120 days. Positive, 148 days. Negative, 48 days. Negative, 48 days. Negative, 48 days. Negative, 58 days. Negative,	cted Iday. Positive. 67 d days. Positive. 100 10 days. Positive. 100 10 days. Positive. 100 10 days. Positive. 20 days. Positive. 20 days. Positive. 60 days. Positive. 60 days. Positive. 50 days. Positive. 60 days. Nogative. 46 days. Nogative. 46 days. Nogative.	(1) Feces injected: 21 days. Positive. 30 days. Positive. 40 days. Positive. (2) Feces fed: 26 days. Negalive. 35 days. Negalive.
	Lot No. 230, 112 bugs infected Sept. 1, 1921.	Lot No. 234. 60 bugs infected Sept. 8, 1921.	Lot No. 247. 16 bugs in foc tod Sept. 19, 1921.

Transmission of tularæmia by bedbugs of the species Cimex lectularius-Continued.

i	1	1	اؤ	l g	1
Romaris.		·	Reported above.	Reported above.	
gth of time after date o allowed to bite a "Positive" means	4. Nogative. 4. Nogative.	5. Negative. Negative. Negative.	Positive,	Positive.	
Transmission following bug bites: Longth of time after date of infection of bugs when bugs were allowed to bite a healthy white mouse or guines pig. "Positive" means death from tularemia.	3 days; 28 bugs bit feet of guines pig 4. 5 days; 24 bugs bit feet of guines pig 4. 8 days; 23 bugs bit feet of guines pig 4. 8 days; 30 bugs bit feet of formouse 41. 20 days; 14 bugs bit tail of mouse 41. 29 days; 9 bugs bit tail of mouse 41. 4 days; 8 bugs bit tail of mouse 41. 54 days; 6 bugs bit tail of mouse 41. 54 days; 6 bugs bit tail of mouse 41. 54 days; 6 bugs bit tail of mouse 41. 54 days; 6 bugs bit tail of mouse 41. 54 days; 6 bugs bit tail of mouse 41. 74 days; 6 bugs bit tail of mouse 41.	a days; 50 bugs bit feet of guinea pig 5. Negati 6 days; 46 bugs bit feet of guinea pig 5. Negative 5 days; 55 bugs bit feet of guinea pig 6. Negative 6 days; 50 bugs bit tail of mouse 4f. 12 days; 48 bugs bit tail of mouse 4g. Repays; 45 bugs bit tail of mouse 4g. 26 days; 17 bugs bit tail of mouse 4g. 26 days; 6 bugs bit tail of mouse 4g. 52 days; 6 bugs bit tail of mouse 4g. 62 days; 6 bugs bit tail of mouse 4g. 62 days; 6 bugs bit tail of mouse 4g. 72 days; 5 bugs bit tail of mouse 4g. 62 days; 5 bugs bit tail of mouse 4g.	18 hours; 62 bugs bit tail of mouse 67.	18 hours; 67 bugs bit tail of mouse 68.	
Transmission due to eating infected bugs. Length of time after date of infection of bugs when dying or dead bugs were eaten by a white mouse. "Positive" means death from tularemia.	1 day; fed 3 dead bugs to mouse 37. Positive. 1 day; fed 3 dead bugs to mouse 38. Positive. 31 days; fod 5 dying bugs to mouse 39. Nogative. 35 days; fed 1 dead bug to mouse 40. Nogative. 53 days; fed 1 dead bug to mouse 98. Nogative.	1 day; fed 3 dying bugs to mouse 42. Positive, 1 day; fed 3 dying bugs to mouse 43. Positive, 27 days; led 14 dead bugs to mouse 44. Positive, 30 days; fed 5 dying bugs to mouse 45. Positive, 40 days; fed 4 dead bugs to mouse 45. Positive, 64 days; fed 1 dead bugs to mouse 46. Positive, 64 days; fed 1 dead bug to mouse 90. Positive.			
Infectivity of bug fees: Number of days after date of infec- tion of bugs when their fresh fees were (1) injected into a guinea pig or (2) fed to a white mouse "Positive" means death from tulare- mia.	(1) Feccs injected: 20 days. Positive. 35 days. Positive. (2) Feccs fed: 30 days. Negative. 40 days. Negative.	(1) Feces injected: 14 days. Fostive. 24 days. Postive. 25 days. Postive. (2) Feces fed: 15 days. Negative. 27 days. Negative.	Feces injected: 35 days. Positive.	35 days. Positive.	,
Various lots of bugs; each lot was infected by biting a separate mouse.	Lot No. 254, 44 bugs in fect od Sept. 25, 1921.	Lot Nos. 255-256. 126 bugs infocted Sept. 26, 1921.	Lot No. 263. 62 bugsinfected Oct. 4, 1921.	Lot No. 264. 62 bugsinfected Oct. 4, 1921.	

Lot No. 262. 80 bugsinfected Oct. 2, 1921.	35 days. Positive.		18 hours; 80 bugs bit tail of mouse 69. Negative.	
Lot No. 272. 66 bugsinfected Oct. 11, 1921.	30 days. Positive.	-	48 hours; 66 bugs bit tail of mouse 70. Negative.	
Lot No. 275. 60 bugsinfected Oct. 15, 1921.	25 days. Positive.		48 hours; 60 bugs bit tall of mouse 71. Negative.	
Lot No. 276. 39 bugsinfected Oct. 15, 1921.	25 days. Positive.		48 hours; 39 bugs bit tail of mouse 72. Negative.	
Lot No. 283. 79 bugsinfected Oct. 21, 1921.	2 days. Positive.		48 hours; 79 bugs bit tail of mouse 73. Negative.	
Lot No. 284. 27 bugsinfected Oct. 21, 1921.	2 days. Positive.	:	48 hours; 27 bugs bit tail of mouse 74. Negative.	

V. TRANSMISSION OF TULARÆMIA BY THE MOUSE LOUSE POLYPLAX SERRATUS (BURM.).

By Edward Francis, Surgeon, and G. C. Lake, Passed Assistant Surgeon, United States Public Health Service.

In two experiments healthy white mice were placed in contact in glass aquarium jars with white mice which had been inoculated subcutaneously with diluted heart's blood of mice dead from tular-emia. Not only did the inoculated mice die from tular-emia but the healthy mice contracted the disease and died with typical lesions.

TRANSMISSION BY CONTACT.

One of the experiments just cited lasted 15 days, from June 24 to July 9.

On the first day of this experiment, 24 healthy mice and 2 mice inoculated two days previously were introduced into a glass aquarium jar 18 inches in diameter. As soon as an inoculated mouse died it was replaced by another inoculated mouse in the third day of the disease; thus the jar was kept constantly supplied throughout the experiment with two inoculated mice in the later stages of the By the fifteenth day of the experiment all of the 24 healthy mice had died, 16 having died with the typical condition of the spleen due to Bacterium tularense. Nine of the 16 spleens were rubbed on the shaved abraded skin of 9 guinea pigs, all of which died with the typical lesions of tularæmia. Of the 16 mice referred to, 1 died on the fifth day of the experiment, 4 died on the seventh day, 3 on the eighth day, 1 on the ninth day, 1 on the tenth day, 1 on the eleventh day, 1 on the twelfth day, 2 on the thirteenth day, 1 on the fourteenth day, and 1 on the fifteenth day. Eight of the 24 mice died from causes other than tularæmia, 4 having died during the first three days of the experiment, and 4 having died later in the experiment.

In a similar experiment, 4 inoculated white mice and 26 healthy white mice were introduced into a glass aquarium jar on August 20, after which date no additional mice were added. The four inoculated mice died within the first five days, with typical lesions of tularæmia. On September 15 the last remaining healthy mouse died with typical lesions of the spleen, which was rubbed on the abraded skin of a guinea pig and caused its death on the sixth day from tularæmia. During the interval of 25 days between August 20 and September 15, the 26 healthy mice all died, the majority of them showing typical lesions of tularæmia.

After the death of the last mouse in the aquarium jar and its removal therefrom, the jar was left undisturbed for eight days. At the expiration of this time eight healthy mice were introduced into

the jar. They remained well, thus showing the absence of any residual infection.

Throughout these two contact experiments we searched the mice and the bedding for parasites and tested the infectivity of the urine of infected mice, and also noted whether dead infected mice had been mutilated by contacts, our object being to determine the mode of transmission in these experiments. The several possible factors concerned in the transmission will now be considered.

INFECTIVITY OF MOUSE URINE.

Throughout our experiments we kept going in white mice strain "S" of *Bacterium tularense* by subcutaneous inoculations of the heart's blood of a recently dead infected mouse into a healthy mouse. Urine voided naturally by living infected mice was collected between 72 and 96 hours after inoculation, since death quite regularly occurred by the end of 96 hours.

The urine collected from four mice was injected subcutaneously into 4 guinea pigs, respectively. The guinea pigs all died acutely with typical lesions of tularæmia. The amounts of urine injected were 10 gtts., 12 gtts., ½ c. c., and ½ c. c. The infectivity of similar samples of mouse urine was also tested by feeding the urine to white mice. The first mouse ate 12 gtts. of such urine mixed with corn meal. The second mouse ate ½ c. c. urine mixed with corn meal on six consecutive occasions spaced about a week apart, thus consuming about 3 c. c. Both mice remained well. We concluded that urine, though infected, was not an agent of transmission in nature.

CANNIBALISM AND INFECTION.

White mice mutilate and sometimes completely eat their dead comrades. While we made no observation on whether a healthy mouse will contract the infection by eating a mouse dead from tularæmia, we did feed six white mice with a small amount of the liver of a rabbit which had just died with typical lesions of tularæmia. All the mice died within five days: One died on the third day, two on the fourth, and three on the fifth day. In each instance the mouse's spleen was rubbed on the shaved, abraded skin of a guinea pig, causing acute death with typical lesions of tularæmia. In our contact experiments, infected mice, when found dead, occasionally showed evidence of mutilation by their comrades.

INFESTATION OF WHITE MICE WITH LICE AND MITES.

Examination of the mice revealed the presence of the blood-sucking louse *Polyplax serratus* (Burm), the blood-sucking mite *Liponyssus isabellinus*, and two species of non-blood-sucking mites. No other

parasites were found either on the mice or in the hay used for bedding.

The lice were present in variable numbers; some mice had none; two had about 60 each; 30 lice was considered a large number for a single mouse; about 15 were commonly found on a mouse. The mites were found in much smaller numbers; most mice had none; 20 was the largest number found on a mouse; 9 was the next largest number found; the number on a mouse hardly ever exceeded 4 or 5.

EXPERIMENTAL TRANSMISSION BY THE MOUSE LOUSE, POLYPLAX SERRATUS.

Experiments were planned to determine definitely the rôle of the mouse louse as an agent in the transfer of the infection from infected mice to healthy mice. Mice were inoculated subcutaneously with diluted blood taken from the heart of a mouse dead from tularæmia, and upon the death of the inoculated mouse his hair was pulled out, transferred to a sheet of white paper and examined for lice. The hair was well teased apart with needles, and any moving object was readily seen in the white hair on the white background. A glance at any moving object with a hand lens was sufficient to exclude the possibility of the parasite being other than a louse. The few hairs to which a louse was clinging were transferred to a clean petri dish, and thus the lice were collected into a pile of very few hairs. The small pile of hairs was then picked up with a forceps and transferred to beneath the hair of the back of a healthy white mouse which was being held. The lice almost instantly left the pile of hairs and disappeared among the hairs of the living mouse. Mice to which lice were thus transferred were placed separately in clean jars for observation.

The time which elapsed between the removal of lice from a dead mouse and their transfer to a healthy mouse probably never exceeded an hour. No effort was made to learn how long lice would remain infected. We did, however, remove from three mice three lots of lice, numbering about 25 to the lot, and kept them on hair in petri dishes at room temperature in August; they were all dead by the end of 48 hours. The time which elapsed between the death of a mouse and the removal of its lice was likewise not definitely determined, although it never exceeded 18 hours. An effort was made to remove the lice as soon as possible. Lice were collected in the morning from mice which died during the previous night, and lice were collected during the day from mice dying during the day.

These transmission experiments, as conducted, excluded the entrance of several possible factors which might have operated to make the contact experiments successful, namely, the agency of the blood-sucking mites, and the eating of infected mice, or possibly infected secretions or excretions, by healthy mice.

The strain of Bacterium tularense (S) used was one isolated from a human case in Utah in the summer of 1920.

First series.—In this series, transmission of tularæmia to healthy mice was effected by the transfer to them of lice removed from mice dead after subcutaneous inoculation with diluted heart's blood of infected mice. Eleven healthy mice were thus infested, the number of lice used for each mouse varying from 10 to 43. Nine of the 11 died with typical lesions of the spleen due to tularæmia, and, moreover, the spleen in each instance, when rubbed on the shaved, abraded skin of a guinea pig, caused the death of this animal with the typical lesions of the disease. Two mice of this series remained negative; 1 had been infested with 10 lice and the other with 33 lice.

Second series.—Transmission of tularæmia was effected in the second series by the transfer of lice from louse-infected mice of the first series to healthy mice. Six healthy mice were thus infested, the number of lice transferred to each mouse varying from 5 to 25. Three of the six mice died with typical lesions of the spleen due to tularæmia, and the spleen in each instance, when rubbed on the shaved, abraded skin of the abdomen of a guinea pig caused the death of this animal with the typical lesions of the disease. Three mice of this series remained well; they had been infested with 5, 9, and 20 lice, respectively.

INFECTIVITY OF THE MITE LIPONYSSUS ISABELLINUS.

Ten mites of the species Liponyssus isabellinus were collected while on the ends of the hairs about to leave the body of a white mouse dead from tularæmia after subcutaneous inoculation with infected blood. The mites were rubbed in a mortar with saline solution and the suspension was injected subcutaneously into a white mouse, causing its death in 4½ days, with typical lesions of tularæmia. A portion of the spleen of the latter mouse was used for subcutaneous injection of one guinea pig, while another portion was rubbed on the shaved abraded skin of another guinea pig; both guinea pigs died acutely with typical lesions of tularæmia. Had these blood-sucking mites been present in sufficient numbers, transmission experiments would have been conducted along the lines followed in the louse transmission.

FAILURE OF ATTEMPTS TO RENDER MICE LOUSE-FREE AND MITE-FREE.

We endeavored to render white mice free from parasites. Nicotine sulphate in water (1:1000), 95 per cent alcohol, undiluted kerosene, and undiluted gasoline were used. A lousy mouse dipped into either of these agents suffered considerable toxic effects; he was rendered apparently free from parasites for a few days, but if examined 7 to 10

days later, he was again lousy. These agents did not kill the eggs. We failed to find a delousing agent into which a mouse could be dipped 4 or 5 times at intervals of a week without causing death or injury to the mouse. Had we been able to render our mice lice-free and mite-free we would have conducted with them a contact experiment similar to our two contact experiments with the expectation that transmission to contacts would not occur.

THE ABSENCE OF LICE FROM HOUSE MICE.

An observation was made on the occurrence of lice on the ordinary house mouse found in various parts of the laboratory. The mice were caught during the night in snap traps and were collected each morning and examined. The hair of 56 mice were pulled out and searched for lice and mites. Only 1 louse and 16 mites were found. This louse was the rat louse *Polyplax spinulosa* (Burm). No specimen was found of the mouse louse *Polyplax serratus* (Burm) so commonly found on our white mice.

The apparent absence of lice from house mice has a bearing on the question of whether tularæmia would become epizoötic in house mice if by chance infected ones got at large.

The susceptibility of "gray" mice to the infection has already been shown by McCoy.² In an experiment of ours, house mouse No. 1, inoculated subcutaneously with the heart's blood of an infected white mouse, died with typical lesions of tularæmia. House mouse No. 2, inoculated with the heart's blood of No. 1, died with typical lesions of tularæmia. The infection was similarly carried over to No. 3. The spleen of No. 3, when rubbed on the shaved, abraded skin of a guinea pig caused its death with typical lesions of tularæmia.

SUMMARY.

The transmission of tularæmia was effected in 12 out of 17 attempts through the agency of the mouse louse (Polyplax serratus) by the transfer of lice from white mice dead of tularæmia to healthy white mice, the intervals elapsing between infestation of the healthy mice and their deaths varying from 5 to 12 days, the average being 74 days. The number of lice transferred in the 12 successful attempts varied from 12 to 43, the average being 25. The intervals which elapsed between the deaths of infected mice and the transfer of their lice to healthy mice varied from a few minutes to 18 hours. Transmission of tularæmia by lice was thus effected to two series of mice, the first series being infected by lice removed from the louse-infected mice of the first series.

² A plaguelike disease of rodents. By George W. McCoy, Passed Assistant Surgeon, United States Public Health Service. Public Health Bulletin No. 43, April, 1911.

When inoculated mice were dropped into a jar in contact with lousy healthy mice, the infection killed off all the healthy mice in 25 days. Transmission in this case was probably due to lice.

Blood-sucking mites of the species *Liponyssus isabellinus* removed from an infected white mouse were crushed and injected subcutaneously into another white mouse causing its death from tularæmia.

The urine of infected white mice was infective for guinea pigs when injected subcutaneously into the latter. Similar urine failed to infect white mice when fed to them on corn meal.

The mouse louse *Polyplax serratus* commonly found on our white mice was absent from 56 house mice caught in snap traps in the laboratory.

Acknowledgment.—The determinations of specimens of lice and mites were made by Dr. H. E. Ewing, of the Bureau of Entomology, Department of Agriculture.

Transmission of tularxmia in white mice by the mouse louse Polyplax serratus.

	Infect	ed mice from wh were removed.			Length of time	guinea pig with
Series.	No. of mouse.	Date of death.	Number of lice transferred from infected mouse to healthy mouse.	Healthy mouse to which lice were transferred.	be- tween infesta- tion with lice and death of mouse.	typical fesions of tulvræmie, the griinea pig hav- ing been rubbed on the shaved abraded skin of the abdomen with the spleen of the louse-in- fected mouse.
First series: Transmission	SM156	July 2, 1921	15	Mouse No. 1	Days.	Positive.
of tularæmia by lice,	SM158	do	14		9	Do.
transferred from inocu- lated mice to healthy	SM155 SM160	July 3, 1921	3 5	Mouse No. 2	9	Ъ0.
mice.	SM161	do	15	do		
	SM162	do		do Mouse No. 3		Do.
	SM163 SM164	July 4, 1921	15 12	Mouse No. 3	9	ъ.
·	SM163	do	15	Mouse No. 4.	6	Do.
	SM166	do	8	do		
	DC6 DC2	July 5, 1921	6 30	do Mouse No. 5.	7	Do.
ļ	DC2	July 5, 1921	30	Mouse No. 6.	9	Do.
	SM172	July 6, 1921	30	Mouse No. 7.	5	Do.
	SM173	do		do		
• 1	SM176	July 7, 1921	15	Mouse No. 8.	6	Do. Do.
	SM178 SM181	do July 9, 1921	12 2 0	Mouse No. 9. Mouse No. 10	12	Negative.
i	SM 181 SM 185	July 9, 1921	13	mouse No. 10		regative.
	DC11	July 10, 1921	iŏ	Mouse No. 11		·Do.
Second series: Transmis-	1	July 7, 1921	14	Mouse No. 20	5	Positive.
sion of tularæmia was	7	July 11, 1921	25	Mouse No. 21	7	Do.
effected by lice trans-	5	July 12, 1921				Negative.
ferred from louse-in-	2	do	.5	Mouse No. 23		Do.
fected mice of first series	8	July 13, 1921	12	Mouse No. 24 Mouse No. 25	7	Positive. Negative.
to healthy mice.	9	July 19, 1921	9	MIOUSE NO. 20		regative.

VI. CULTIVATION OF BACTERIUM TULARENSE ON MEDIUMS NEW TO THIS, ORGANISM.

By EDWARD FRANCIS, Surgeon, United States Public Health Service.

The only culture mediums reported heretofore for the cultivation of *Bacterium tularense* are congulated hen's egg yolk, originally used by McCoy and Chapin,³ and hen's ovomucoid with a trace of yolk, recommended by Wherry and Lamb.⁴ All attempts at cultivation on other laboratory mediums had failed.

The writer now reports the cultivation of this organism on (1) serum glucose agar, (2) glucose blood agar; (3) blood agar, (4) each of the foregoing mediums plus a piece of fresh, sterile rabbit spleen.

It is a common practice in laboratories to adapt an organism to growth on ordinary mediums after original isolation from man or animals has been accomplished by cultivation on some special medium. The mediums which are here reported for *Bacterium tularense* were used for original isolations of this organism from animals, and therefore the question of acquired adaptability to a new medium brought about by previous cultivation on a special medium is not involved.

The strains used for this cultural work were three human strains from Utah and one ground-squirrel strain from California, all obtained in 1920. These strains were originally obtained by the inoculation of human or squirrel tissue into guinea pigs, and they were subsequently passed for many generations through guinea pigs or rabbits, always by subinoculation of infected tissues. Original isolation from animals of pure cultures of these strains was accomplished by use of the proposed mediums; the resultant cultures have never been on egg medium either before or since their first isolation.

COMPOSITION OF MEDIUMS.

(1) Serum glucose agar.—Beef infusion containing 1 per cent peptone and 1½ per cent agar adjusted to a reaction having a p_H of 7.6 is kept on hand in stock. When needed, the stock agar is melted and brought to 45° C. in a water bath, at which temperature there is added 1 per cent glucose from a sterile 50 per cent solution of glucose and 5 per cent sterile horse serum. This is immediately tubed, slanted, and incubated 48 hours to insure sterility.

³ Bacterium tularense, the Cause of a Plague-like Disease of Rodents. By George W. McCoy and Charles W. Chapin, Passed Assistant Surgeons, United States Public Health Service. Public Health Bulletin No. 53, January, 1912.

Further Observations on a Plague-like Disease of Rodents with a Preliminary Note on the Causative Agent, Bacterium tularense. By George W. McCoy and Charles W. Chapin, Passed Assistant Surgeons, United States Public Health Service. The Journal of Infectious Diseases, Vol. X, No. 1, January, 1912, pp. 61-72.

⁴ Infection of Man with Bacterium tularense. By William B. Wherry and B. H. Lamb. Journal of Infectious Diseases, 1914, vol. 15, p. 331.

- (2) Glucose blood agar.—This is the same as (1) except that 5 per cent defibrinated rabbit blood is substituted for the horse serum.
- (3) Blood agar.—This is the same as (2) except that no glucose is added.
- (4) Mediums (1), (2), and (3) plus spleen tissue.—A spleen is removed from a healthy rabbit and under sterile precautions is cut into pieces of about 3 mm. diameter. One piece is rubbed on the slanted surface of each tube of a portion of mediums (1), (2), and (3) and the piece of spleen is left remaining on the surface of each slant just above the water of condensation. After 48 hours' incubation the tubes, if sterile, are ready for inoculation.

COAGULATED HEN'S EGG YOLK.

At this place it will be well to give also the composition of the coagulated hen's egg yolk described by McCoy and Chapin. Fresh eggs are scrubbed with a brush in soap and water, if fecal matter is present on the shells, and then placed in a wire basket. The basket containing the eggs is dipped into 95 per cent alcohol for a few seconds, after which time it is withdrawn and the small amount of alcohol which still remains on the basket and eggs is ignited in order to remove the alcohol and help sterilize the shells.

While one person with clean hands holds an egg, grasping it at each end, an assistant strikes the shell in its middle with a sterile knife with sufficient force to crack the shell. The whites are separated from the yolks by decanting from one half of the shell to the other, thus allowing the whites to drain away while the yolks are saved and collected in a sterile beaker.

The volume of yolks is measured in a sterile graduate and to this is added sterile normal saline solution in the proportion of 40 per cent saline solution to 60 per cent egg yolk. Mix thoroughly. Tube in sterile test tubes, using a sterile funnel.

Place the tubes in metal racks constructed so as to allow one-half-inch space between the tubes for circulation. Heat the racked tube, in a slanting position for the first half hour at 70° C., and for the second half hour at 72° C. A uniform temperature is best maintained for this purpose in a water-jacketed chamber. The chamber should contain about a half inch of water above which the racks of tubes are exposed in the moist, heated air. After coagulation, paraffined sterile cork stoppers are substituted for the cotton plugs and the tubes are incubated upright for three or four days to insure against a slow-growing contamination.

Instead of the jacketed chamber one may, with patience, use an Arnold steam sterilizer, a board having been placed at the bottom to protect the tubes from the direct steam.

The finished medium should be soft; that is, the surface of the slant should yield slightly when pressed with a platinum loop, and to that end the medium should not be overheated. A glazed surface results from overheating. The tubes should be stored in the cold room, unexposed to the light. The water of condensation in a batch of medium which grew the organism very well showed a reaction having a p_{π} of 6.8. No titration or adjustment of reaction has been done on batches of this medium used for routine cultivation of the organism in the laboratory.

CULTIVATION ON SERUM GLUCOSE AGAR.

Serum glucose agar was successfully used (1) for original isolation of strains from the spleens of infected rabbits, (2) for second and third isolations of strains from the spleens of infected guinea pigs, (3) for subcultures (without the addition of a piece of fresh, sterile rabbit spleen), and (4) for subcultures (with the addition of a piece of fresh, sterile rabbit spleen).

(1) Original isolation of strains.—Human strains "J" and "G" and ground-squirrel strain "S F" were isolated on serum glucose agar by planting a piece of infected rabbit spleen on serum glucose agar in each instance. (See Table I, and Table III, animals 1, 3, and 4.) These human strains had been carried over from July 3 and September 9, 1920 (the dates on which they left the humans). to April 11, 1921 (the date on which they were cultured), in laboratory animals, i. e., guinea pigs and rabbits. These human strains having been carried over exclusively by animal passages for 9 and 7 months. respectively, original isolations were made on serum glucose agar. The ground-squirrel strain had been carried over from May. 1920 (the date on which it left a California ground squirrel), to April 10. 1921 (the date on which it was cultured), in laboratory animals, guinea pigs, and rabbits. This California ground-squirrel strain having been carried over exclusively by animal passages for 11 months, original isolation was made on serum glucose agar.

On April 10 and 11, 1921, each of the three strains was inoculated on a serum glucose agar slant by transferring to the surface of the medium a piece about 3 mm. in diameter, taken under sterile precautions, from the spleen of a rabbit dead from tularæmia; the piece of spleen was rubbed over the surface of the medium as forcibly as the consistency of the latter would permit and then left to remain on the solid medium just above the water of condensation. The inoculated tubes were placed at 37° C. for eight days, at the end of which time they were observed for the first time and found to have a growth which was not examined microscopically, but the tubes were replaced in the incubator under the impression that the growths

were contaminations. The tubes remained at 37° for a month longer without observation, at the end of which time they were given another examination preliminary to discarding. The presence of the growth saved the tubes from the discard and they remained another month at room temperature, when, on June 23 and August 23, the growths were examined microscopically and found to simulate *Bacterium tularense*.

Confirmation of these original cultures was obtained when subcultures in the fifth generation on glucose blood agar plus a piece of fresh, sterile rabbit spleen caused acute death with typical lesions of tularæmia in a set of guinea pigs which had been rubbed on the shaved abraded skin of the abdomen with those subcultures.

Further confirmation was obtained when cultures derived from the spleens of the above set of guinea pigs caused acute death with typical lesions of tularæmia in a second set of guinea pigs, the second set having been rubbed on the shaved abraded skin of the abdomen with subcultures in the fifth generation on glucose blood agar plus a piece of fresh sterile rabbit spleen. (For details see Table I.)

(2) Second and third isolations.—The two sets of guinea pigs just referred to not only gave confirmation to the identity of the original cultures which were isolated on serum glucose agar but afforded opportunities for the second and third isolations of those strains from animals on serum glucose agar as follows: On the death of a guinea pig of either set a piece of its spleen was planted on a tube of serum glucose agar and incubated at 37° C. Ten tubes were thus planted from 10 guinea pigs. (See Table III, animals 5 to 15.) Five tubes showed growth and five showed no growth.

The five growths therefore constituted second and third isolations of our three human strains and one ground squirrel strain on serum glucose agar.

- (3) Subcultivation on serum glucose agar.—Plain serum glucose agar was used for subcultures as follows: Starting with a piece of infected spleen of a rabbit or guinea pig, this was planted on a tube of serum glucose agar and the resultant growth was transferred for its second generation to a tube of serum glucose agar. Thirteen serum glucose agar tubes were thus inoculated with six cultures having an antecedent history as indicated. Four of the tubes developed growth after an average of six days and nine failed to grow. (See Table III, footnotes 4, 5, and 6.) The four growths constituted subcultivation on serum glucose agar.
- (4) Subcultivation on serum glucose agar (plus a piece of fresh sterile rabbit spleen).—Subcultivation from serum glucose agar to serum glucose agar plus a piece of fresh sterile rabbit spleen was successful in 14 out of 27 attempts; the average time which elapsed before the appearance of growth on the 14 tubes was 2½ days.

Three cultures were thus carried for the third to sixth generations on serum glucose agar plus a piece of fresh sterile rabbit spleen, the first generation in each case having been obtained on a tube of serum glucose agar inoculated with a piece of spleen of an infected guinea pig. (See Table III, animals 5, 6, and 12.) The fifth generation of culture 1 was injected subcutaneously into a guinea pig, causing its death in three days with typical lesions of tularæmia. The sixth generation of culture 1 was rubbed on the shaved abraded skin of a guinea pig and caused its death on the twelfth day with typical lesions of tularæmia. (See Table I, footnote 2.) The sixth generations of cultures 2 and 3 when injected subcutaneously each failed to kill a guinea pig.

Summary.—Serum glucose agar per se is a poor medium for the cultivation of Bacterium tularense because subcultures from serum glucose agar to serum glucose agar grew in only 30 per cent of the instances. Serum glucose agar is a fair medium for the cultivation of Bacterium tularense if the surface of the medium is provided with a piece of fresh spleen tissue. Such tissue may be supplied in either of two ways.

If the inoculating material is in substance a piece of fresh tissue, as is the case when a piece of the spleen of an infected rabbit or guinea pig is planted on a serum glucose agar tube, growth may be expected in 48 per cent of the tubes. Growth did occur in 11 out of 23 such attempts, the average time before the appearance of growth being $8\frac{1}{2}$ days.

If the inoculating material is a culture, as is the case in subcultivation, and a piece of fresh sterile rabbit spleen has been supplied to the surface of the serum glucose agar tubes, growth may be expected in 51 per cent of the tubes. Growth did occur in 14 out of 27 such attempts, the average time which elapsed before the appearance of growth on the 14 tubes being $2\frac{1}{2}$ days.

It was noted, however, that the growth of *Bacterium tularense* on serum glucose agar under all conditions tends to become scanty and its virulence tends to become either diminished or lost.

CULTIVATION ON GLUCOSE BLOOD AGAR.

Glucose blood agar was successfully used (1) for original isolation of strains from the spleens of infected rabbits, (2) for second and third isolations of strains from the spleens, liver, and heart's blood of infected guinea pigs, (3) for subcultures (without the addition of a piece of fresh sterile rabbit spleen), and (4) for subcultures (with the addition of a piece of fresh sterile rabbit spleen).

(1) Original isolation of strains.—Human strains "J" and "S" were isolated on glucose blood agar by planting a piece of infected

rabbit spleen on a glucose blood agar tube in each case. (See Table III, animals 1 and 2.)

These human strains had been maintained in the laboratory in guinea pigs and rabbits from September 9 and July 3, 1920 (the dates on which they left the humans), to April 11 and 13, 1921, the dates on which they were first cultured; they had not been on any culture medium during that period. On April 11 and 13 these strains were inoculated on glucose blood agar slants by transferring to the surface of the medium a piece about 3 mm. in diameter taken under sterile precautions from the spleen of a rabbit dead from tularæmia. The tubes were incubated at 37° and not observed for 8 days, at the end of which time growth was present in case of the "J" strain, but no notation was then made of growth of "S" strain, although it was noted at a later date. As in the case of original cultures on serum glucose agar these cultures remained unobserved in the incubator for one month longer, when, on May 24, strain "J" was subcultured and it was subcultured next again on June 22; whereas strain "S" went from April 13 to July 4 before the first subculture was made.

Confirmation of the original cultures was obtained when subcultures in the fifth generation on glucose blood agar, plus a piece of fresh, sterile rabbit spleen caused acute death with typical lesions of tularæmia in a set of guinea pigs which had been rubbed on the shaven, abraded skin of the abdomen with those subcultures.

Further confirmation was obtained when cultures derived from the spleens of the above set of guinea pigs caused acute death with typical lesions of tularæmia in a second set of guinea pigs, the second set having been rubbed on the shaved, abraided skin of the abdomen with subculture in the fifth generation on glucose blood agar, plus a piece of fresh, sterile rabbit spleen. (For details see Table II.)

(2) Second and third isolations.—The two sets of guinea pigs just referred to not only gave confirmation to the identity of the original cultures which were isolated on glucose blood agar, but afforded opportunity for the second and third isolations of these strains from animals on glucose blood agar as follows: On the death of a guinea pig of either set, a piece of its spleen was planted on a tube of glucose blood agar and incubated at 37° C. Thirteen tubes were thus planted from 13 guinea pigs, including liver and heart's blood in two instances. (See Table III, animals 5 to 15.) All of the 13 tubes showed growth after an average of a little less than four days.

These 13 growths therefore constituted second and third isolations of these two human strains on glucose blood agar.

(3) Subcultivation on glucose blood agar.—Plain glucose blood agar was used for subculture as follows: Starting with a piece of infected spleen of a rabbit or guinea pig, this was planted on a tube of glucose

blood agar and the resultant growth was transferred for its second generation to a tube of glucose blood agar. Five glucose blood agar tubes were inoculated with five cultures having an antecedent history as indicated. None of the tubes developed any growth. (See Table III, animals 21 to 25.)

Four cultures grown for two or three generations on glucose blood agar, plus a piece of fresh sterile rabbit spleen, were subsequently transferred for two or three generations to glucose blood agar; the transfer was accompanied by a falling off in the abundance of growth. Moreover, a falling off in virulence also took place because the last generation of each culture was rubbed on the shaven, abraded skin of one or two guinea pigs, with the result that one guinea pig died acutely on the seventh day, two died tardily on the twelfth day, one died subacutely on the twenty-third day, and two guinea pigs, vaccinated with the fourth culture, remained well. (See Table I, footnote 1, and Table II, footnotes 1, 2, and 3.)

(4) Subcultivation on glucose blood agar (plus a piece of fresh, sterile rabbit spleen).—Fifteen cultures which had been isolated by planting a piece of the spleen of an infected rabbit or guinea pig on either glucose blood agar or serum glucose agar, with or without the addition of a piece of fresh, sterile rabbit spleen were subcultured from the second to the fifth generations on glucose blood agar plus a piece of fresh sterile rabbit spleen; the fifth generation in each instance was rubbed on the shaven abraded skin of a guinea pig causing its death acutely with typical lesions of tularæmia. (See Tables I and II, and Table III, footnote 5.)

This medium was successful in 88 out of 103 attempts at subcultivation of the organism; the average time which elapsed before the appearance of growth on the 88 tubes was two days; growth appeared before the end of 24 hours in 61 of the 88 tubes.

Isolation of cultures from animal tissues to glucose blood agar plus a piece of fresh sterile rabbit spleen was successful in 21 out of 27 attempts. The animal tissues consisted of a piece of the spleen of 19 infected rabbits and guinea pigs, a piece of the liver of one infected guinea pig, the heart's blood of two infected guinea pigs, and the heart's blood of five infected white mice. The average of time before the appearance of growth on the 21 tubes was five days. (See Table III.)

Summary.—Glucose blood agar per se is not a good medium for the cultivation of Bacterium tularense. The plain glucose blood agar is a good medium only when the infected material with which it is inoculated is in substance a piece of fresh tissue as represented by a piece of the infected spleen of a rabbit or guinea pig. This was the case when 25 rabbits or guinea pigs dead from tularæmia each furnished a piece of spleen which was inoculated on a tube of glucoso

blood agar. Growth appeared after an average of $4\frac{1}{3}$ days on 21 tubes; no growth appeared on 4.

Subcultures were made on 56 tubes of glucose blood agar; growth appeared after an average of three days on 26 tubes and failed to appear on 30. The growth, when subcultured on plain glucose blood agar became scanty and of lowered virulence.

The falling off in growth and virulence which accompanied the transfer of cultures to plain glucose blood agar is accounted for by the absence of a piece of fresh sterile rabbit spleen from the medium.

On the other hand, glucose blood agar plus a piece of fresh sterile rabbit spleen is a good medium, both for the isolation of Bacterium tularense from animals and for subcultivation. This medium was successful in 21 out of 26 attempts at isolation of the organism from the tissues of 26 animals. This medium was successful in 88 out of 103 attempts at subcultivation of the organism. Fifteen subcultures each in the fifth generation on this medium were rubbed on the shaven abraded skin of 15 guinea pigs, causing acute death in each instance with typical lesions of tularæmia, thus indicating no loss of virulence after subcultivation on this medium.

CULTIVATION ON BLOOD AGAR.

Blood agar was used (1) for second and third isolation of strains from rabbits and guinea pigs and (2) for subcultures.

(1) Second and third isolations.—No attempt was made at original isolation of our strains from animals on blood agar, but second and third isolations of these strains from animals were accomplished on blood agar. (See Table III, animals 5, 8, 10, 11, and 13.)

The organism was isolated on a blood agar slant which had been inoculated with a piece of the spleen of animal No. 5 dead after vaccination with the fifth generation of a culture which was originally isolated on serum glucose agar and subsequently grown for four generations on glucose blood agar plus a piece of fresh sterile rabbit spleen. Practically the same statement can be made concerning blood agar cultures obtained from animals 8, 10, 11, and 13.

Isolation on blood agar slants was also accomplished from the spleens of animals Nos. 21, 22, 23, 24, and 25 (see Table III), all of which had been inoculated with the heart's blood of Mouse 206. For one year previous to Mouse 206 the strain had had a continuous passage through mice, tame rabbits, and guinea pigs, back to human case "S" in Utah except that once during that year (from May 27 to June 6, 1921) the strain was carried on coagulated egg yolk for ten days.

(2) Subcultivation.—Subcultures on blood agar failed to grow in 13 out of 15 attempts; growth appeared on two tubes after three and six days, respectively:

Summary.—For the isolation of Bacterium tularense from infected animals, this medium was successful in 9 out of 15 attempts; growth appeared on the nine tubes after an average of seven days. The success of this medium for isolation from infected animals was undoubtedly due to the transfer to the medium of a piece of fresh tissue as represented by the piece of infected spleen of rabbit or guinea pig with which the tubes were inoculated. For subcultivation of this organism, blood agar per se is a poor medium, growth having been obtained only two times out of 15 attempts at subcultivation on this medium.

CONCLUSION.

The view heretofore held that *Bacterium tularense* will grow only on a culture medium containing egg yolk is no longer tenable.

The present paper contains reports of the growth of this organism in subcultures on serum glucose agar, glucose blood agar and blood agar. Growth on the above mediums per se is, however, scanty and of lowered virulence.

But these mediums take on an exalted value for the cultivation of this organism when supplied with a piece of fresh tissue; this tissue may be supplied either by the piece of spleen of the infected rabbit or guinea pig with which the medium is inoculated or a piece of fresh sterile spleen of a rabbit may be transferred to the medium, thereby preparing it to grow a subculture with which it may subsequently be inoculated.

The success of the cultural experiments here reported can not be ascribed to adaptation from a special medium to an ordinary medium because our mediums were employed for original isolations of the strains. The work here reported is with strains of *Bacterium tularense* which have never been on egg medium either before or since isolation; the only exception to this statement is contained in the very limited work done on animals 16 to 29 of Table III, in which the strain had had a few days' cultivation on coagulated egg yolk as exemplified by the statement at the bottom of page 109.

From the data presented in Table III there appears to be very little difference between the efficiency of glucose blood agar plus a piece of fresh rabbit spleen, and coagulated egg yolk. I am however of the opinion that coagulated egg yolk, carefully prepared, is still the best medium for routine isolation and cultivation of *Bacterium tularense*.

Sub-TABLE I.—Serum glucose agar used for original isolation of human strains "J" and "G" and squirrel strain "SF" of Bacterium tularense. cultures made on glucose blood agar and serum glucose agar, each being supplemented by a piece of fresh sterile rabbit spleen.

California ground squirrel strain "SF." (See Table III, animal No. 4.)	Piece of infected rabbit spleen planted on serum glucose agar. Subcultured on glucose abood agar plus a piece of fresh sterile rabbit spleen. Subcultured on above medium. Aug. 23. Subcultured on above medium. Aug. 25. Subcultured on above medium. Aug. 26. Subcultured on above medium. Aug. 27. Subcultured on above medium. Aug. 28. Subcultured on above medium. Aug. 29. Subcultured on above medium. Aug. 29. Subcultured on above medium. Aug. 29. Subcultured on above medium. Subcultured fifth generation. Aug. 29. Subcultured, fifth generation. Subcultured on above medium. Subcultured, fifth generation. Subcultured on above medium. Subcultured o
	rum Apr. 1921 piece Aug. 5 dug. Aug. Aug. Aug. Aug. Aug. Aug. Aug. A
Human strain "G." (See Table III, animal No. 3.)	
Hum	1921. Apr. 11. June 22. July 14. July 14. July 13. July 25. Aug. 21. Aug. 22. Aug. 24. Sept. 1.
Human strain "J." (See Table III, animal No. 1.)	Plece of infected rabbit spleen planted on serum Apr. 11. Subcultured on plucose blood agar plus a plece of fresh sterile rabbit spleen. Subcultured, fifth generation. Subcultured, fifth generation. Subcultured, fifth generation. Guinea pig dying, chlordormed, planted pieces of its spleen on 5 mediums. (See Table III, animal No. 7.) Subcultured from glucose blood agar to glucose blood agar plus a piece of fresh sterile rabbit. Subcultured from glucose blood agar to glucose blood agar to glucose blood agar to glucose blood agar to glucose blood agar plus a piece of fresh sterile rabbit. Subcultured on above medium. Salden spleen above medium. Sept. 1. Subcultured on above medium. Sept. 1. Subcultured on above medium. Sept. 1. Subcultured on above medium. Sept. 1.
Hum	1921. June 23. July 4. July 14. July 18. July 18. July 18. July 18. July 18. July 18. July 25. Aug. 21. Sept. 2. Sept. 3. Sept. 5.

1 This culture of June 22 being in the second generation was also subcultured for the third, fourth, and fifth generations on glucose blood agar without the addition of a piece offresh sterile rabbit spleen. The fifth generation was then vaccinated on two guinea pigs on July 23. Both guinea pigs were dead Aug. 4 with typical lesions of tularramia.

1 Sopt. 14. Guinea pig injected subcutaneously with 3-day culture of the fifth generation.

2 Sopt. 15. Subcultured on above medium.

2 Sopt. 17. Guinea pig dead with typical lesions of tularramia.

3 Sopt. 18. Vaccinated a guinea pig with a loop of 4-day culture of the sixth generation.

3 Sopt. 19. Subcultured, fifth generation.

3 Sopt. 11. Subcultured, sixth generation.

h

. Subcultures on glucose blood aga;	
TABLE II. Glucose blood agar used for original isolation of human strains "J" and "S" of Bacterium tularense. Subcultures on glucose blood	plus a piece of fresh sterile rabbit spleen.

Human strain "S." (See Table III, animal No. 2.)	Apr. 11. Piece of infected rabbit spleen planted on glucose Apr. 12. Place of infected rabbit spleen planted on glucose blood agar. Apr. 11. Place of infected rabbit spleen planted on glucose blood agar. Apr. 11. Place of infected rabbit spleen application of the spleen planted on glucose blood agar. Juno 22. Subcultured on above medium. Juno 23. Subcultured on above medium. July 19. Subcultured on above medium. Aug. 19. Subcultured on above medium. Aug. 21. Subcultured on above medium. Aug. 22. Subcultured on above medium. Aug. 23. Subcultured from glucose blood agar plus a piece of the start plus and the subcompanies and th
Human strain "J." (See Table III, animal No. 1.)	agar plus a piece d agar plus a piece lune 22. Subcultured on glucose blood agar plus a piece lune 24. Subcultured on above medium. lune 24. Subcultured on above medium. luly 3. Subcultured on above medium. luly 9. Subcultured on above medium. luly 14. Subcultured on above medium. luly 18. Vaccinated a guinea pig with a loop of a 4-day culture of the sixth generation. luly 18. Vaccinated a guinea pig with a loop of a 4-day culture of the sixth generation. look agar plus a piece of the splean on 7 mediums. (See Table III, animal No. 6.) animal No. 6. Subcultured from glucose blood agar plus a piece of test splean on 7 mediums. Aug. 24. Subcultured from glucose blood agar plus a piece of test schlean on above medium. Aug. 25. Subcultured from glucose blood agar plus a piece of test schlean on above medium. Aug. 28. Subcultured from glucose blood agar plus a piece of test schlean on above medium. Aug. 28. Subcultured from glucose blood agar plus a piece of test schlean on above medium. Aug. 28. Subcultured from glucose blood agar plus a piece of test schlean on a samo medium. Aug. 28. Subcultured from glucose blood agar plus a piece of test schlean on a samo medium. Aug. 28. Subcultured from glucose blood agar plus a piece of test schlean on a samo medium. Aug. 28. Subcultured from glucose blood agar plus a piece of test schlean on a samo medium. Aug. 29. Subcultured from glucose blood agar plus a fiece of test schlean on a samo medium. Aug. 29. Subcultured from glucose blood agar plus a fiece of test schlean on a samo medium. Aug. 29. Subcultured from glucose blood agar plus a fiece of test schlean on a samo medium. Aug. 29. Subcultured from glucose blood agar plus a fiece of test schlean on a samo medium. Aug. 20. Subcultured from glucose blood agar plus a piece
Human strain "J." (Sec Table III, animal No. 1.)	Apr. 11. Five of infected rabbit spleen planted on glucose blood agar. May 24. Subcultured on glucose blood agar plus a piece of fresh steriforabbit spleen. June 22. Subcultured on above medium. July 10. Subcultured on above medium. July 11. Subcultured on above medium. July 12. Subcultured fifth generation. July 27. Guinea pig with a loop of a 4-day culture of the fifth generation. July 27. Guinea pig dying, chlorformed, planted pieces of its spleen on 7 mediums. (See Table III, animal No. 5.) Aug. 8. Subcultured from glucose blood agar plus a piece of fresh sterile rabbit spleen to same medium. Aug. 14. Subcultured to same medium. Aug. 23. Subcultured to same medium. Aug. 24. Subcultured to same medium. Aug. 25. Subcultured to same medium. Aug. 26. Vaccinated a guinea pig with a loop of a 48-hour culture of the sixth generation. Sept. 9. Guinea pig dying, chlorformed, showed typical lesions of tularæmia.

This subcluture plane. This sixth generation was on July 22 vaccinated on two guines page of tiefs sterile rabbit spiech. This sixth generation was also subcultured for the fourth, fifth, and sixth generations on glucose blood agar. This sixth generation was on July 23 vaccinated on two guines need fresh sterile rabbit spiech. This sixth generation was also subcultured for the form blood agar to blood agar. This sixth generation was also subcultured on the above medium.

July 17. Subcultured on above medium.

July 32. Vaccinated a guines pig with a loop of 6-day culture of the fifth generation.

July 33. Quines pig dying, chloroformed, planted pieces of its spiece on 8 mediums.

July 34. Guines pig dying, chloroformed, planted pieces of its spiece on 8 mediums.

July 35. Vaccinated a guine pig with a loop of 6-day culture of the fifth generation.

July 36. Guines pig dying, chloroformed, planted pieces of its spiece on 7 mediums. This subculture planted July 10 being in the fourth generation was also subcultured for the fifth and sixth generations on glucose blood agar without the addition of a piece of fresh sterile rabbit spiecn. This sixth generation was on July 23 vaccinated on a guinea pig which on Aug. 15 was dying and was chloroformed; it showed the typicallesions of

Table III.—Comparative value of several mediums for the isolation of Bacterium tularense from the spleens and heart's blood of 29 infected animals.

				Medium	on which piece	Mediums on which pieces of infected tissue were planted	e were planted.		
No. of mail-	Either a piece of the spleen or the heart's blood of infected animals was planted.	Date planted.	Coagulated egg yolk.	Glucose blood agar slant plus a piece of fresh sterile rabbit spicen.	Glucose blood agar slant.	Blood agar slant.	Serum glucose agar slant.	Plain agar slant.	Glucore fermenta- tion tube.
	(A) Original Isolations of Bacterium Tularense.	1921.			:			;	
-	Human strain "J" from spleen of rabbit J45R	Apr. 11	Good growth		Growth after 8 days.1		ie.	No growth. No growth.	Nogrowth.
8	Human strain "S" from spleen of rabbit S51R	Apr. 13	Good growth		Growth 1		- I	qo	Do.
က	Human strain "G" from spleen of rabbit GCR3	Apr. 11	Growth		No growth		Growth after	do	Do.
∢ .	California ground squirrel strain "SF" from spleen of rabbit SF25R.	Apr. 10	Growth, second day.		do		Growth after 9 days.1	do	Do.
	(B) SECOND AND THIRD ISOLATIONS OF ABOVE STRAINS.		; ,		-		1	-	
r.	5 Spleen of guinea pig dead after vaccination with July 27 subculture from animal No. 1.	July 27	Growth, third day.	About 100 colonies, fifth	About 50 colo- onies, fourth	5 colonies, fifth Growth, fifth Nogrowth. Nogrowth.	Growth, fifth day.	No growth.	Nogrowth.
9.	do	July 31	Growth at end of 24 hours.	About 100 col- onies, fourth	do.3		3 colonies, fourth day.	do	Do.
7	July 25	July 25	Contaminated.	ರ	Pure culture,	Contaminated. Contaminated.	Contaminated.		
<i>y</i> 5	Spleen of guines pig dead after vaccination 2 with	do	Growth	Good growth,	Good growth,	Growth, fifth	Nogrowth	No growth.	Do.
<i>3</i> 3	Subcutture from animal No. 2. 9 Spleen of guinea pig dead after vaccination 2 with		Sept. 1 No growth	Good growth,	Good growth,	No growth	do	do	Do.
10		July 30	do	mira asy.	do.	Good growth,dodo	do	do	Do.
	Subentured on choose blood sear plus a piece of fresh sterile rabbit spleen from second to fifth generations; the fifth generation was rubbed on the shaved abraded skin of a	h storile rat	bit spleen from s	second to fifth ge	nerations; the fit	th generation wa	as rubbed on the	shaved abrac	ed skin of a

Subcultured on glucose blood agar plus a piece of fresh sterile rabbit spleen from second to fifth generations; the fifth generation was rubbed on the shaved abraded skin of guine pilety enshing its death acutely with typical lesions of tularramia.

Wacchintion means that the shaved abraded skin was rubbed with a culture or with a piece of infected tissue.

**Subcultures on this same medium for the second, third, and fourth generations grew well.

**A subculture on this same medium for the second and spenerations showed growth.

**Subculture on this same medium for the second and third generations and on serum glucose agar plus a piece of fresh sterile rabbit spleen for the fourth to seventh generations;

**Subculture on this same medium for the second and third generations and on serum glucose agar plus a piece of fresh sterile rabbit spleen for the fourth to seventh generations.

**A subculture on this same medium for the second and third generation falled to grow.

**A subculture on this same medium for the second and third generation falled to grow.

TABLE III.—Comparative value of several mediums for the isolation of Bacterium tularense from the spleens and heart's blood of 29 infected animals—Continued

No growth. fermenta-tion tube. Glucose å Š å å ådo Contami-....do.... No growth No growth No growth Plain agar slant. nies, ninth No growth.... Mediums on which pieces of infected tissue were planted. 30 colonies sf-Growth at end Serum glucose agar slant. ter 21 days.7 of 4 days. 1 dozen colo-nies, ninth day. Growth at end Blood agar slant. of 5 days. No growthdo do.4. Good growth, second day.9 Growth at end growth, colonies, rowth, seventh day. Good growth, Glucose blood agar slant. third day. of 48 hours. fifth day.3 Growth. 000 C day.4 Good growth, Good growth, fifth day.1 Growth at No growth.... do.1 Growth, sevsecond day. Growth eighth day.8 25 colonies, third day. 900 Growth Growth, sixth agar slant plus a piece of fresh Good growth, third day. ninth enth day.3 Glucose blood sterile rabbit around dozen day. ours. day. nies,do....do.... Good growth, second day. No growth.... Growth at end No growth.... Growth....do.... Good growth, third day. 9 Good growth. of 24 hours. Coagulated egg yolk. ond day. No growth Growth. ...qo... : ...qo... Aug. 14 2 2 8 æ ង Date planted. Sept. Sept. ...do ...do July ...de July July July Spleen ofguinea pig dead after subcutaneous injection with cultures grown on glucose blood agar plus a Liver of guinea pig No. 15. cutaneously with heart's blood of mouse. Heart's blood of white mouse SM214 inoculated subcutaneously with culture on egg. Heart's blood of white mouse SM215 inoculated sub-Heart's blood of guinea pig No. 14...... cutaneously with heart's blood of mouse. Heart's blood of white mouse SM207 inoculated sub-Heart's blood of guinea pig No. 11..... Spleen of guinea pig dead after vaccination with sub-Spleen of guinea pig dead after subcutaneous injec-Heart's blood of white mouse SM206 inoculated sub-Spleen of guinea pig dead after vaccination 2 with Spleen of guina pig dead after vaccination with spleen of guinea pig dead after subcutaneous injection ABOVE Either a piece of the spleen or the heart's blood of unfected animals was planted. tion with cultures grown on glucose blood agar. (C) ISOLATIONS FROM VARIOUS SOURCES. ö AND THIRD ISOLATIONS with subculture from animal No. 4. ece of fresh sterile rabbit spleen. STRAINS—Continued cutaneously with culture on egg. subculture from animal No. 10. culture from animal No. 3. SECOND â 11 8 2 9 2 z 2 = è, ani-mal.

Electric blood of white mouse BM217 incentated sub- cutaneously with heart's blood of mouse BM20. Spleen of rabbit RL24 incontated with spleen of mouse BM20. Spleen of guinea pig vaccinated with spleen of mouse BM20. Spleen of guinea pig vacci						-					
bod of white mouse SM217 incentated sub- subti RL22 incentated sub- sub- sub- sub- sub- sub- sub- sub-		•	Nogrowth.				6	 6	គំ	ģ	
bod of white mouse SMZ17 inoculated sub- lay with hear're blood of mouses free blood of mouse SMZ06. Third day.	фо	qo	qo			qo	qo	ф	ф	фо	
and of white mouse SM217 incentated sub- state about series and of mouse SM217 incentated sub- state block of mouse SM207 incentated sub- strate block of mouse SM206 incentations		Good growth,	No growth	Good growth, seventh day.	No growth	qo.	т.	ф	Growth		112184
bod of white mouse SM217 incomisted sub- rabbit RL29 incoulsted subcutaneously rar's blood of mouse SM206. do Growth do		No growth	Good growth, ninth day.	Growth, eighth day.	Growth, third	Growth, seventeenth	No growth	•		No growth	తాబచ్చి
bod of white mouse SM217 incoulated sub- lay with heart's hood of mouse Tablit RL29 incoulated subcutaneously Tree blood of mouse SM206. Tablit RL29 incoulated subcutaneously Tree blood of mouse SM206. Tablit RL28 incoulated subcutaneously Tree blood of mouse SM206. Tablit RL28 incoulated subcutaneously Tree blood of mouse SM206. Tree blood growth. T		Growth,	Good growth,	Growth, third day.	do.4	Growth, four- teenth day.	No growth	do	Growth,	Growth, seventh day.	84.¥.
bod of white mouse SM217 innoulated sub- lay with heart's blood of mouse SM207 innoulated sub- tr's blood of mouse SM206. Tabbit RL28 incoulated subcutaneously rr's blood of mouse SM206. Tabbit RL28 incoulated subcutaneously rr's blood of mouse SM206. Tabbit RL28 incoulated subcutaneously rr's blood of mouse SM206. Tabbit RL28 incoulated subcutaneously rr's blood of mouse SM206. Tabbit GRE incoulated subcutaneously rr's blood of mouse SM206. Tabbit GRE incoulated with spleen of mouse pig vaccinated with spleen of mouse and ay. This spond growth, tuines pig vaccinated with spleen of mouse and ay. This spond growth, thind day. This spond growth, thind day. Thind	No growth	Good growth,		No growth		do.	Good growth,	Good growth,	Good growth, fourth day.	No growth	22.7.7.7.7.7.7.7.7.7.7.7.7.7.7.7.7.7.7.
bod of white mouse SM217 inoculated sub- lay with heart's hood of mouse. The bolt RL20 inoculated subcutaneously Trie blood of mouse SM206. The Blood of M206. The Blood of M206. The Blood of M206. The Blood of M206. The Blood	qo	Good growth,	Good growth,	Growth, second day.	до	No growth	Good growth,	Good growth,	Good growth, third day.	Good growth, fifth day.	23 74.2
ood of white mouse SM217 inoculated sub- laly with heart's flood of mouse. The blood of mouse SM20. The blood of mouse of the plean of the p	July 17	July 26	July 29	July 30	 8	July 29	do	Aug. 5	Aug. 14	Aug. 30	
21 Spien of Spien of Spien of Spien of Spien of With hea Spien of With hea Spien of With hea Spien of	20 Heart's blood of white mouse SM217 incoulated sub-	21 Spleen of rabbit RL20 incoulated subcutaneously with heart's blood of mouse SM206.	22 Spieen of rabbit RL28 incomisted subcutaneously with heart's blood of mouse SM206.	23 Spleen of rabbit RL3 inoculated subcutaneously with heart's blood of mouse SM206.	with heart's blood of mouse SM206.	25 Spleen of rabbit GCR inoculated subcutaneously with heart's blood of mouse SM206.	Spleen of guinea pig vaccinated	Spleen of guines pig vaccinated	Spleen of guinea pig vaccinated with spleen of mouse C4.	Spieen ofguines pig vaccinated with spieen of mouse 31.	Total number of "growths" on each medium Total number of "no growths" on each medium Percentage of "growths" on each medium

1 Subcultured on glucose blood agar plus a piece of fresh sterile rabbit spleen from second to fifth generations; the fifth generation was rubbed on the shaved abraded akin was rubbed on the shaved abraded akin was rubbed with a culture or with a piece of infected tissue.
 2 Subcultures on this same medium for the second, third, and fourth generations grew well.
 3 Subcultures on this same medium for the second generation showed growth.
 3 A subculture on this same medium for the second generation and on serum glucose agar plus a piece of fresh sterile rabbit spiece for the third to sixth generations; the sixth generation and on serum glucose agar for the second generation and on serum glucose agar for the second generation and on serum glucose agar for the second generation and on serum glucose agar for the second generation and on serum glucose agar for the second generation and on serum glucose agar for the third to sixth generations; the sixth generation and on serum control of the guines of tularemia.

A subculture on coagulated egg yolk grew well.

THE BASAL METABOLISM OF INFANTS FED ON DRY MILK POWDER.

. By FRITZ B: TALBOT, M. D., Boston, Mass., assisted by Miss MARGARET E. MORIARTY.

This report is part of an investigation on the use of dry milk powder in infant feeding made by the United States Public Health Service. A preliminary report made by Surg. W. H. Price appeared in Public Health reports, April 2, 1920.

The present study on the basal metabolism of 13 normal infants fed on dry milk powder was conducted at the Massachusetts General Hospital, and it was through the helpful cooperation of Miss Helen Falvey, temporary supervising nurse, United States Public Health Service, and the Baby Hygiene Association of Boston, that this part of the investigation was made possible.

The clinical status of the babies studied was obtained from their records at the Baby Hygiene Association. The babies were also given a superficial physical examination on the day that the metabolism was determined. In general, it may be said that they were satisfied with their food, were gaining weight, and had well-digested stools. As in the cases reported by Surg. Price, group 2 received 164 grams (equivalent to 1½ cupfuls) of whole dry milk powder to 1 quart of boiled water, giving approximately the following: Fat 4 per cent, sugar 5.7 per cent, and protein 3.71 per cent. Group 3 received reconstructed, or emulsified, milk made from dry skim milk powder, unsalted butter fat, and water emulsified in a centrifugal apparatus. The constituents of the reconstructed milk were, approximately, fat 4 per cent, sugar 5.1 per cent, and protein 3.1 per cent.

The technique of investigation was the same as that used by Benedict and Talbot in previous investigations on infant metabolism. (See publications of the Carnegie Institution of Washington, 201, 233, and 302.) The infants were brought to the Massachusetts General Hospital in the morning. They were fed immediately before they entered the respiration chamber to find out the effect, if any, of the dry milk powder on the basal metabolism and compared with normal infants fed at the breast or bottle fed with ordinary milk dilutions.

The following is a summary of the histories, showing the clinical status of the infants.

SUMMARY OF CASE HISTORIES.

CASE 1: A. P., a male infant 9 months old. Full term, normal labor, birth weight unknown. Always bottle fed and had done well. When 3 months old weighed 4.025 kilos, and was put on a mixture of whole dry milk powder, water, and malt sugar. Four months later weighed 7.2 kilos. At time of experiment, when 9 months old.

¹ Dried Milk Powder in Infant Feeding. Reprint No. 588.

weighed 7.944 kilos, had 7 teeth, stools were normal. In addition to the dry milk mixture, he was receiving orange juice. Rectal temperature on day of metabolism experiment, 99% and 100° F.

Summary.—A normal infant, close to expected weight of 8.39 kilos for age.

Case 2: M. F., a female infant, 5½ months old. Full term, normal labor, birth weight unknown. Breast fed for two weeks and then put on whole milk and maltose. On this mixture she had much gas and slight eczema of face. When 3½ months old weighed 6.35 kilos, and was put on a whole dry milk powder, and is recorded as having improved steadily. When seen at 5½ months of age weighed 7.895 kilos, was bright, happy, and well proportioned, with firm flesh. She had been receiving orange juice since birth. At the time of metabolism experiment she was receiving whole dry milk powder 6½ ounces, boiled water 48 ounces; 6 ounces every three hours.

Summary.—A well, healthy infant, weighing 845 grams more than the expected weight of 7.05 kilos.

Case 3: R. S., a male infant, 93 months old. Full term, normal labor, breast fed during the first three months. When weaned, was put on a whole milk and maltose mixture, but did not do well. At 4 months of age was given a whole dry milk mixture and improved steadily; bowels regular, stools normal. He was receiving orange juice daily. No teeth. When metabolism was determined, he was receiving whole dry milk powder 6 ounces, water 32 ounces.

Summary.—An average, normal infant, weighing 8.675 kilos. Expected weight 8.66 kilos.

CASE 4: M. H., female infant, 8½ months old. Premature birth; said to have weighed less than 0.9 kilo. Never breast fed. Until 2½ months of age was on a whole milk formula and gained very slowly. Since then on a dry milk powder mixture gaining consistently. At time metabolism was determined was receiving 6 ounces of whole dry milk powder to 1 quart of water—7 ounces, boiled water 1 ounce. Getting orange juice irregularly. She was well rounded, firm flesh, and had two teeth.

Summary.—Normal, underweight infant, weighing 6.776 kilos. Expected weight, had she been at term, would be 8.23 kilos.

Case 5: J. N., a male infant, 7½ months old. Birth weight unknown. Always bottle fed. For first three months received a mixture of whole milk, water, and maltose; the next two months fed on a condensed milk mixture, and at 5 months on a mixture of whole dry milk powder. At this time was in a fair condition, perspiring freely, and had several boils. At time metabolism was determined he was receiving 5 ounces of dry milk powder to 32 ounces of water—5 ounces, water 2 ounces, condensed milk 2 teaspoonfuls—and also orange juice. Improved steadily.

Summary.—Of average normal development, weighing 7.854 kilos. Expected weight 7.98.

Case 6: E. S., a male infant, 9 months old. Full term, instrumental delivery, always bottle fed. For the first three months fed on a grade A milk formula, but did not thrive. Then given a whole dry milk powder mixture and did well, but remained under weight for age. Two teeth. At time metabolism was determined was getting 6 ounces of whole dry milk powder to 1 quart of water, 2 tablespoonfuls of maltose—8 ounces every 3 hours, 6 feedings.

Summary.—A moderately underweight infant, flesh firm, but evidence of mild rickets, weighing 7.059 kilos. Expected weight 8.39.

Case 7: H. C., a female infant, 81 months old. Full term, normal labor, breast fed for one week; then given Nestle's food, was not satisfied, regurgitated, troubled with constipation, and had eczema. At 11 months of age given a whole dry milk powder mixture and improved steadily. At time metabolism was determined she was receiving 6 ounces of whole dry milk powder, 47 ounces of water, and 4 tablespoonfuls of maltose—8 ounces every 3 hours, 6 feedings.

Summary.—Normal infant, weighing 9.072 kilos. Expected weight 8.175.

CASE 8: H. H. T., a male infant, 113 months. Full term, normal at birth, breast fed for one month. Then given grade A whole milk formula, but did not gain well. At 4 months of age was given a reconstructed dry milk mixture and improved steadily. At time metabolism was determined he was receiving an emulsified milk mixture and cereals. He had never had orange juice. Ten teeth.

Summary.—At 113 months weighed 8.618 kilos. Expected weight 9.43.

CASE 9: F. C., a female infant, 6 months old. Full term, Cæsarean birth. Partly breast and partly bottle fed up to date of metabolism determination, the minor part of the food said to be breast milk. Has always been well. At 2 months of age was given an emulsified milk mixture as follows: Reconstructed dry milk, 20 ounces; boiled water, 10 ounces; maltose, 2 level tablespoonfuls—6 ounces every 3 hours, 6 feedings. Receiving orange juice.

Summary.—At 6 months weighed 6.974 kilos. Expected weight, 7.27 kilos.

Case 10: P. N., female infant, 6½ months old. Full term, normal delivery, breast fed for two months, and was then given a mixture of grade A milk, but did not do well. At 5 months was given 30 ounces of reconstructed milk, 12 ounces of boiled water, and 4 level tablespoonfuls of maltose—6 ounces every 3 hours, 7 feedings. Improved steadily. Was getting orange juice. Vaccinated 6 days before metalbolism was done.

Summary.—At 6½ months weighed 7.087 kilos. Expected weight 7.499 kilos.

CASE 11: I. V., a female infant, age 3 months. Fed on a reconstructed milk mixture. Weight at 3 months, 4.337 kilos. Expected weight, 5.56 kilos.

CASE 12: C. C., a female infant, age 11½ months. Full term, normal delivery. Breast fed for 3 months. Then given a mixture of grade A milk. Was not sick, but did not gain well. At 7 months was put on a reconstructed milk mixture; improved, but did not gain very rapidly. At time metabolism was determined was receiving undiluted reconstructed milk, cereals, beef juice, and orange juice.

Summary.—At 11½ months weighed 7.272 kilos. Expected weight 9.373.

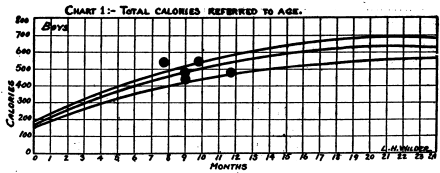
The following table gives the metabolism findings of these infants.

Case No.	Sex.	Age.	Height.	Weight.	Average weight for age.	Number of basal metabo- lism de- termina- tions.	Total calories.	Calories per kilo, body weight.	Calories per sq. m., body surface.	Pulse.
1 2 3 4 5 6 7 8 9 10 11	MEMEMERE	Mo. 99 55 55 55 55 55 55 55 55 55 55 55 55	Cm. 76 72 72 67 62.5 68 71 69.5 66	Kilos. 7. 944 7. 895 8. 675 6. 776 7. 854 7. 069 9. 072 8. 618 6. 974 7. 088 4. 337 7. 272	Kilos. 8.390 7.054 8.660 8.290 7.984 8.390 8.150 9.430 7.270 7.499 5.560 9.373	12222121123323	458 513 541.5 421.5 542 478 570 478 478 478 503 288 559	57. 7 65 64. 5 61. 5 67. 8 63. 8 55. 3 68. 5 71 66. 4 73. 3	1,078 1,220 1,215.5 1,124 1,295 1,229 1,236 1,074 1,074 1,074 1,405	113 118 119 103 114.5 118 106 111 117 128 113.5

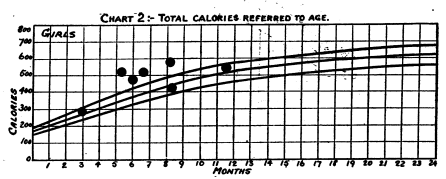
TABLE I .- Metabolism findings of babies fed on dry milk powder.

Since none of these infants was more than 20 per cent below the expected weight for its age, they did not fall in the class of severe malnutrition or marasmus (see Talbot, Severe Infantile Malnutrition, American Journal Diseases of Children, 1921, 22, 358). The clinical impression was that this group of children were doing well, eight of the infants being within 10 per cent of the average weight for the age, and may be considered average normal infants.

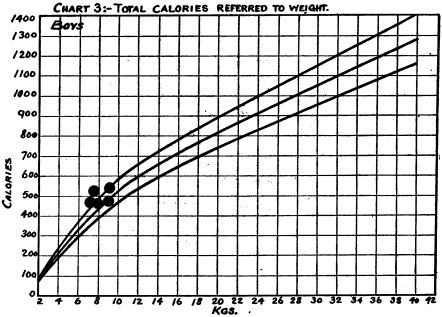
The accompanying charts show the distribution of the cases as compared to the normal standards. The heavy black line represents the average metabolism; the lines on either side represent the 10 per cent variation which may be considered normal.



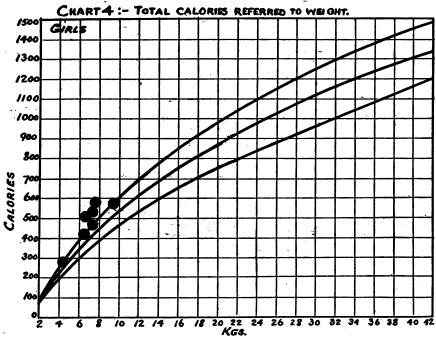
Basal metabolism of boys during first 24 months of age, showing the total calories per 24 hours.



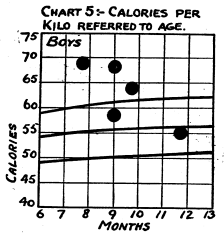
Basal metabolism of girls during first 24 months of age, showing the total calories per 24 hours.



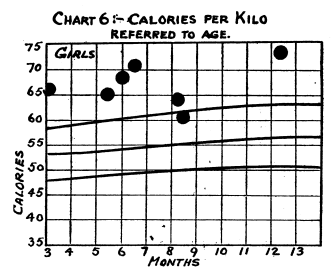
Basal metabolism of boys, showing total calories in 24 hours at different weights. The curve is projected from 32 kg. upward.



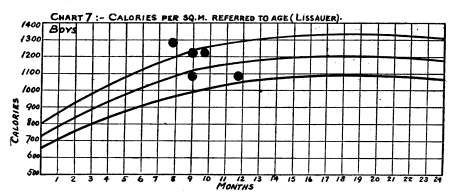
Basal metabolism of girls, showing total calories in 24 hours at different weights. The curve is projected from 32 kg. upward.



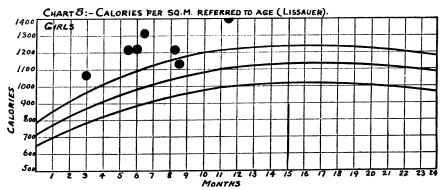
Basal metabolism of boys, showing calories per kilogram of body weight per 24 hours.



Basal metabolism of girls, showing calories per kilogram of body weight per 24 hours.



Basal metabolism of boys, showing calories per square meter of body surface per 24 hours during first 24 months of age.



Basal metabolism of girls, showing calories per square meter of body surface per 24 hours during first 24 months of age.

SUMMARY.

The metabolism studies in this series of cases show a tendency for the boys to fall within, or very close to, the standard variations, the greatest deviation being in the calories per kilogram of body weight. The metabolism of the female infants, on the other hand, ran higher, and, with few exceptions, fell more than 10 per cent above the average, but are not outside of the extreme normal variation. This difference in the metabolism of the different sexes is in accord with what Benedict and Talbot² found in their series of observations; that is, that it was much more difficult to predict the metabolism of female than of male infants, and that the deviation from the average was much greater in the former than in the latter.

The results of the findings of this study on the basal metabolism of infants fed on dry milk powder show either a normal or a slight elevated "basal" metabolism. Coincident with most of the cases of an elevated metabolism, there was an elevation in temperature to 99° to 100° F.; but, as far as the writer has been able to determine, such slight elevations of temperature are not sufficient to explain the slight increase in the metabolism.

Case 11 had been vaccinated against smallpox six days previous to the metabolism determination. There is no published data on the effect of vaccination on the metabolism, but it seems reasonable to suppose that the metabolism would be more likely to increase on the tenth or twelfth day during the "take" rather than earlier.

All precautions were taken to insure quiet periods; and careful pulse and kymograph and visual records were kept of the infants during the metabolism periods to warrant recording the results as purely "basal" findings.

The basal metabolism of this series of infants fed on dry milk powder mixtures tends to be slightly higher than that of average normal infants, but is within normal limits. This may have been due to the relatively high protein content of the food, but the deviations from the average are not great enough to permit any striking conclusions to be drawn.

INFLUENZA IN ENGLAND AND GERMANY.

Official information relative to the occurrence of influenza in England, received by cable January 17, 1922, states that the disease is prevalent in several parts of England and is spreading. The epidemic period is short in each area affected, being from four to six weeks. The type of disease is stated to be of a much milder form than was that of the epidemic of 1918. Broncho-pneumonia is reported to be the chief complication.

Benedict and Talbot: Carnegie Ins., Washington, Pub. No. 302.

Outbreaks similar to that in England are reported from continental Europe, and the British authorities believe that further general spread is probable.

According to information dated January 4, 1922, received from an officer of the Public Health Service and based on German newspaper reports, epidemic influenza is prevailing in Berlin, Baden, Mannheim, and Wurttemberg, and it is stated that the hospital facilities in Berlin are being overtaxed with the care of influenza patients.

DEATHS DURING WEEK ENDED JAN. 7, 1922.

Summary of information received by telegraph from industrial insurance companies for week ended Jan. 7, 1922, and corresponding week, 1921. (From the Weekly Health Index, Jan. 10, 1922, issued by the Bureau of the Census, Department of Commerce.)

	Week ended Jan. 7, 1922.	Corresponding week, 1921.
Policies in force	48, 628, 495	45, 142, 899
Number of death claims	6, 740	7, 362
Death claims per 1.000 policies in force, annual rate	7.2	8. 5

Deaths from all causes in certain large cities of the United States during the week ended Jan. 7, 1922, infant mortality, annual death rate, and comparison with corresponding week of 1921. (From the Weekly Health Index, Jan. 10, 1922, issued by the Bureau of the Census, Department of Commerce.)

	Estimated		ended , 1922.	Annual death rate per		hs under year.	Infant mor- tality
City.	population July 1, 1921.	Total deaths.	Death rate.1	1,000, corre- sponding week, 1921.	Week ended Jan. 7, 1922.	Corresponding week, 1921.	rate, week ended Jan. 7, 1922. ²
Total	26, 586, 734	6,907	13.5	13. 6	915	957	
Akron, Ohio	3 208, 435	31	7.8	8.4	9	.4	95
Albany, N. Y	115,071	35	15. 9	12.2	4	3	90
Atlanta, Ga	207, 473	66	16.6	13.8	8	6	· · · · · · · · · · · · · · · · · · ·
Baltimore, Md	750, 864	228	15. 8	14.2	31	30	87
Birmingham, Ala	186, 133	45	12.6	17.1	7	-10	
Bridgeport, Conn.	* 143, 555	32	11.6	13.6	4	6	50
Buffalo, N. Y.		128	12.8	12.7	18	22	71
Cambridge, Mass	110,444	34	16. 1	14.6	7	7	128
Camden, N. J.	119,672	36 633	15. 7 11. 9	10.9 12.6	6 96	117	92
Chicago, Ill	2,780,655 403,418	123	15.9	16.9	9	10	60
Cleveland, Ohio.	831, 138	199	12.5	11.3	21	17	54
Columbus, Ohio	245, 358	58	12.3	14.5	5	4	53
Dallas, Tex.	165, 282	58	18. 3	12.0	ğ	4	- 55
Dayton, Ohio	³ 152,559	38	13.0	7.9	4	5	68
Denver, Colo.	263, 152	97	19. 2	19.6	7	6	
Detroit, Mich		236	11.5	12.8	46	58	89
Fall River, Mass.	120,668	28	12.1	18.6	4	4	56
Fort Worth, Tex	111, 423	15	7.0	100	i	*	- 50
Grand Rapids, Mich	141, 197	29	10.7	12.6	$\hat{\mathbf{z}}$	4	33
Houston, Tex	144,310	48	17. 3	11.9	9	3	
Indianapolis, Ind	325, 632	95	15. 2	14.1	7	18	53
Jersey City, N. J.	302, 788	77	13. 3	14.1	15	18	96
Kansas City, Kans	103, 884	20	10.0	13.6	2	4	46
Kansas City, Kans. Kansas City, Mo.	336, 157	110	17. 1	11.5	15	16	10
Los Angeles, Calif.	614, 160	180	15. 3	16.6	14	17	58
Lonisville, Ky		85	18.8	15. 9	5	io	54
Annual rate per 1,000 population.	===,000 1	•• ,	23.0	2017	•		

² Deaths under 1 year per 1,000 births—based on deaths under 1 year for the week and estimated births for 1921. Cities left blank are not in the registration area for births.

* Enumerated population Jan. 1, 1920.

Deaths from all causes in certain large cities of the United States during the week ended Jan. 7, 1922, infant mortality, annual death rate, and comparison with corresponding week of 1921. (From the Weekly Heatth Index, Jan. 10, 1922, issued by the Bureau of the Census, Department of Commerce)—Continued.

	Estimated		ended ', 1922.	Annual death rate per	Deaths under 1 year.		Infant mor- tality	
City.	population July 1, 1921.	Total deaths.	Death rate.	1,000, corre- sponding week, 1921.	Week ended Jan. 7, 1922.		rate, week ended Jan. 7, 1922.	
Lowell, Mass. Memphis, Tenn Milwaukee, Wis. Minneapolis, Minn Nashvilie, Tenn New Bedford, Mass. New Haven, Conn New Orleans, La. New York, N. Y Newark, N. J Norfolk, Va. Oakland, Calif. Omnha, Nebr. Paterson, N. J Philadelphia, Pa Pittsburgh, Pa. Portland, Oreg. Providence, R. I Richmond, Va. Rochester, N. Y St. Louis, Mo. St. Paul, Minn Salt Lake City, Utah San Francisco, Calif. Seattle, Wash.	113, 757 165, 656 468, 386 392, 815 122, 036 125, 042 167, 007 394, 657 5, 751, 867 424, 885 121, 260 226, 472 197, 066 137, 463 1, 866, 212 264, 859 239, 645 175, 686 305, 229 237, 781 121, 595 522, 546 231, 531 121, 595 123, 595 124, 859 125, 126 127, 126	36 67 96 90 35 30 30 1,486 125 49 49 43 35 498 157 66 70 54 65 70 54 65 70 54 69 70 70 70 70 70 70 70 70 70 70 70 70 70	16. 5 21. 1 10. 7 11. 9 15. 0 10. 8 9. 4 18. 5 15. 3 17. 2 11. 3 11. 4 13. 3 13. 9 15. 2 16. 0 11. 1 12. 1 16. 9 8. 1	20. 2 14. 8 9. 9 14. 6 18. 4 19. S 12. 8 14. 7 13. 8 14. 5 12. 9 14. 0 16. 8 9. 5 17. 0 19. 6 9. 1 13. 3 14. 7 15. 8 10. 0 11. 5	7 9 9 16 9 2 4 2 2 19 9 210 4 8 8 6 6 6 32 5 1 4 4 9 9 13 8 6 6 8 8 3 3 3 3 3	4 14 14 77 8 9 16 184 222 3 4 4 7 61 28 6 8 12 23 6 9 9 11 7	118 118 118 119 1102 1102 1111 111 1111	
Springfield, Mass. Syracuse, N. Y. Toledo, Ohio. Trenton, N. J. Washington, D. C. Wilmington, Del. Wonkers, Mass. Yonkers, N. Y.	135, 877 177, 265 253, 696 122, 760 3 437, 571 113, 408 184, 972 103, 324	37 42 56 51 115 28 50 31	14. 2 12. 4 11. 5 21. 7 13. 7 12. 9 14. 1 15. 6	13. 8 16. 8 14. 8 14. 0 15. 6 18. 9 14. 7 10. 6	1 9 8 19 5 5	7 9 3 23 7 7	15 108 88 122 109 175 54 104	

³ Enumerated population Jan. 1, 1920.

PREVALENCE OF DISEASE.

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

UNITED STATES.

CURRENT STATE SUMMARIES.

Telegraphic Reports for Week Ended Jan. 14, 1922.

These reports are preliminary and the figures are subject to change when later returns are received by the State health officers.

the State Realth omeets.		• ;	
ALABAMA.	Cases.		ases.
Cerebrospinal meningitis	1	Measles	2
Chicken pox	59	Mumps	8
Diphtheria	13	Pneumonia	5
Hookworm disease	35	Scarlet fever	. 45
Ophthalmia neonatorum		Smallpox	
Pneumonia	14	Tuberculosis	
Poliomyelitis		Typhoid fever	. 3
Scarlet fever			-
Smallpox	18	CONNECTICUT.	
Typhoid fever		Cerebrospinal meningitis.	. 2
		Chicken pox	
ARKANSAS.		Conjunctivitis (infectious)	. 3
Chicken pox	18	Diphtheria:	
Diphtheria	8	Bridgeport	. 18
Influenza	40	Hartford.	
Malaria	17	New Haven	
Measles	2	Scattering	
Pellagra	2	German measles	
Scarlet fever	6	Influenza	
Tuberculosis		Measles:	
Typhoid fever		Glastonbury	. 28
		New Haven	
CALIFORNIA.		Stamford	
Diphtheria	205		
Lethargic encephalitis—Butte County.	1	Scattering	
Measles	14	MumpsOphthalmia neonatorum	
Scarlet fever	86		
Smallpox:		Paratyphoid fever	
Bakersfield	19	Pneumonia (lobar)	
Kern County	36	Poliomyelitis	. 1
Monterey County	11	Scarlet fever:	_
San Jose.	19	Bridgeport	
Santa Clara County	27	Hartford	
Scattering		New Haven	
Typhoid fever	5	Stamford	
•		Scattering.	
COLORADO.		Smallpox	
(Exclusive of Denver.)		Trichinosis.	
•	1	Tuberculosis (all forms)	
Chicken pox		Typhoid fever	
Diphtheria	33	Whooping cough	54
	/19	6)	

DELAWARE. Cases.		PATRITA NY A	70.000
Chicken pox	g l		`ases.
	4 0	Cerebrospinal meningitis:	
	1	Allen County	
	2	Randolph County	
• • • • • • • • • • • • • • • • • • • •	1 1	Steuben County Diphtheria	118
Scarlet fever:	I	Rabies in animals—Orange County	
Lewes	3 I s	Scarlet fever	
Wilmington 38		Smallpox	. 33
Scattering 22 Tuberculosis:	² 1	Ty phoid fever	6
Wilmington 10		****	
	. 1	IOWA. Diphtheria	39
Typhoid fever	~ ! *	Scarlet fever.	
FLORIDA.		Smallpox	
	٠ ا		
Diphtheria 19 Influenza 6	9	KANSAS.	
Malaria1		Cerebrospina! meningitis	
	4 0	Chicken pox	
Smallnox	5 I	Diphtheria	
Typhoid fever	Di	German measles	
GEORGIA.		nfluenza	
the state of the s	١.	dumps	
Cerebrospinal meningitis	* E	Pneumonia	
Chicken pox	^у I т	Poliomyclitis	
Diphtheria 34 Dysentery (amebic) 1		cariet fever	. 248
Dysentery (amebic)	18	Smallpox	. 65
Hookworm disease 29	9 T	Puberculosis	
Influenza	a 1	Typhoid fever	
Malaria2	2 Y	Whooping cough	. 22
Mumps 2	2	LOUISIANA.	
Pneumonia 22	1 1	Diphtheria	. 23
Poliomyelitis	1 1	nfluenza	
Scarlet fever	1 1	Paratyphoid fever	. 1
Smallpox	. 0	carlet fever	
Tetanus 2 Tuberculosis (all forms) 18	۰۱۰	mallpox	
Typhoid fever	1 1	Typhoid fever	. 14
Whooping cough4	4	MAINE.	
ILLINOIS.		hicken pox	
		Diphtheria	
Cerebrospinal meningitis—Chicago 1		German measlesnfluenza	
Diphtheria:	1 -	fumps	
Aurora	· I _	neumonia	
East St. Louis	~	carlet fever	
Rockford11	i S	ma`lpox	. 2
Streator 9		'ubercu!osis	
Scattering		Pyphoid fever	
Influenza49		Vhooping cough	. 10
Lethargic encephalitis—Chicago		MARYLAND.1	
Pneumonia	¹ c	erebrospinal meningitis	. 2
Poliomyelitis: Charleston		hicken pox	
Chicago		Diphtheria	
Scarlet fever:		German measles	
Chicago 121		nfluenza	
Rockford	7 I .	feasles	
Scattering		fumps	
Smallpox:		Pneumonia (all forms)	
Chicago		cabies	
Peoria	· .	carlet fever	
Typhoid fever		eptic sore throat	
Whooping cough		ubereulosis	
1 777 - 1 2 - 1 77 - 1	-		

1 Week ended Friday.

MARYLAND—continued.	ses.	NEBRASKA.	ases
Typhoid fever			
Whooping cough.		Chicken pox.	
		Diphtheria:	
Massachus etts .		Omaha	
Cerebrospinal meningitis		Scattering	. 1
Chicken pox			
Conjunctivitis (suppurative)		LincolnOmaha	
Diphtheria German measles		Scattering	
Influenza.		Mumps.	
Lethargic encephalitis		Scarlet fever:	
Measles		Seward County	. 1
Mumps		Scattering	. 7
Ophthalmia neonatorum		Smallpox:	
Pneumonia (lobar)		Gering	
Poliomyelitis		Scattering	
Scarlet fever		Typhoid fever	• '
Septic sore throat		NEW JERSET.	
Typhoid fever		Cerebrospinal meningitis	. :
Whooping cough		Chicken pox.	
	••	Diphtheria	
MINNESOTA.		Influenza.	
Cerebrospinal meningitis	1	Measles	. 18
Chicken pox	18	Pneumonia	
Diphtheria	91	Scarlet fever	
Influenza		Smållpóx	
Measles		Trachoma	
Pneumonia	6	Typhoid fever	
Scarlet fever		Whooping cough	. 100
Smallpox		NEW MEXICO.	
Tuberculosis	45 4	Chicken pox	. 26
Typhola lever	*	Diphtheria	
MISSISSIPPI.		Measles	. 1
Diphtheria	28	Mumps	
Scarlet fever	15	Pneumonia	
Smallpox		Scarlet fever	
Typhoid fever	12	Smallpox	
MISSOURI.		Tuberculosis	
Cerebrospinal meningitis	2	Typhoid fever	_
Chicken pox	54	w nooping coagn	•
Diphtheria		NEW YORK.	
Epidemic sore throat	22	(Exclusive of New York City.)	
Glanders	1	•	
Influenza	16	Corebrospinal meningitis	
Measles	10	Diphtheria	
Mumps	8	Influenza	
Ophthalmia neonatorum	3	Measles	
Pneumonia	34	Pneumonia	
PoliomyelitisScarlet fever.	1	Typhoid fever	
Scarlet lever	65	Whooping cough	133
Tetanus	3		200
Tuberculosis	59	NORTH CAROLINA.	
Typhoid fever	4	Chicken pox	129
Whooping cough	6	Diphtheria	54
		German measles	4
MONTANA.		Measles	8
Cerebrospinal meningitis—Lewistown	1	Scarlet fever	65
Diphtheria	24	Septic sore throat	10
Poliomyelitis—Billings	1	Smallpox	23
Scarlet fever	36	Trachoma	4
Smallpox	33	Typhoid fever	99

OREGON.	ases.	WASHINGTON. C	ases.
Chicken pox	. 24	Cerebrospinal meningitis—Adams County	. 1
Diphtheria:		Chicken pox	. 48
Portland			
Scattering		•	
Measles	_	1	
Mumps			
Pneumonia		Smallpox. Tuberculosis.	
Scarlet feverSmallpox:	. 10	Typhoid fever	
Portland	. 17	Whooping cough.	
Scattering		, and the second	
Tuberculosis	-	WEST VIRGINIA.	
Typhoid fever		Diphtheria:	
Whooping cough		Fairmont	. 9
SOUTH DAKOTA.		Scattering	
Chicken pox	. 18	Scarlet fever	
Diphtheria		Smallpox	
Measles		Typhoid fever	. 4
Pneumonia		WISCONSIN.	
Scarlet fever		Milwaukee:	
Smallpox		Chicken pox	138
Tuberculosis		Diphtheria	
Whooping cough	. 1	Measles	. 1
TEXAS.		Pneumonia	. 9
Diphtheria		Scarlet fever	. 30
Measles		Smallpox	
Pneumonia		Tuberculosis	
Poliomyelitis	-	Typhoid fever	
SmallpoxScarlet fever		Whooping cough	. 19
· ·	. 10	Scattering:	
VERMONT.	. 40	Cerebrospinal meningitis	
Chicken pox		Chicken pox Diphtheria	
DiphtheriaInfluenza		German measles.	
Measles.		Influenza.	
Mumps		Lethargic encephalitis	
Pneumonia		Measles	
Scarlet fever		Pneumonia	
Smallpox	5	Poliomyelitis	2
Typhoid fever	3	Scarlet fever	127
Whooping cough	30	Smallpox	50
VIRGINIA.		Tuberculosis	
Smallpox:		Typhoid fever	
Botetourt County	3	Whooping cough	35
Delayed Reports fo	r W	eek Ended Jan. 7, 1922.	
•		KENTUCKY—continued.	
DISTRICT OF COLUMBIA.			ses.
	ases.	Measles: Ca Jefferson County	
Chicken pox	26	Scattering	6
Diphtheria		Mumps	
Influenza Lethargic encephalitis	î	Pellagra	1
Measles	1	Pneumonia	48
Scarlet fever.		Scarlet fever	22
Smallpox	7	Septic sore throat	2
Tuberculosis	21	Smallpox:	
Typhoid fever	1	Jefferson County	10
Whooping cough	3	Scattering	12
KENTUCKY.		Trachoma	4
Chicken pox	6	Tuberculosis:	_
Diphtheria:		Jefferson County	8
Jefferson County	18	Scattering	3
Scattering	39	Typhoid fever	7 4
Influenza	17	Whooping cough	*
¹ Deaths.			

SUMMARY OF CASES REPORTED MONTHLY BY STATES.

The following summary of monthly State reports is published weekly and covers only those States from which reports are received during the current week:

State.	Cerebrospinal meningitus.	Diphtheria.	Influenza.	Malaria.	Measles.	Pellagra.	Poltomyelitis.	Scarlet fever.	Smallpox.	Typhoid fever.
DECEMBER, 1921. District of Columbia. Florida. Louisiana Massachusetts. Michigan Nebraska. Vermont. Wisconsin	1 1 1 14 3	191 137 102 1, 088 1, 448 209 44 618	6 41 21 46 24 4	79 36 2	12 13 4 835 320 115 31 62	16 6	2 1 10 5 2 1	56 20 55 740 1, 164 363 191 679	18 20 33 118 161 5 224	26 56 88 52 101 11 6 31

PLAGUE (RODENT).

Galveston, Tex.

One rat was reported positive for plague at Galveston, Tex., January 17, 1922. The rat was trapped December 28, 1921.

CITY REPORTS FOR WEEK ENDED DEC. 31, 1921.

ANTHRAX.

	City.	 Cases.	Deaths.
New York: New York		 1	
	BERIBERI.		
California: San Francisco		 1	

CEREBROSPINAL MENINGITIS.

The column headed "Median for previous years" gives the median number of cases reported during the corresponding weeks of the years 1915 to 1920, inclusive. In instances in which data for the full six years are incomplete, the median is that for the number of years for which information is available.

City.	Median for pre-		City.	Median for pre-	Week ended Dec. 31, 1921.		
	vious years.	Cases.	Deaths.		years.	Cases.	Deaths.
California: San Francisco. Connecticut: Bridgeport. Illinois: Chicago Indiana: Muncie. Iowa: Marshalltown Massachusetts: Holyoke Michigan. Highland Park. Port Huron	0 0 1 0	2 1 2 1 1	1 1 2	Missouri: Kansas City. St. Louis. New Jersey: Hackensack. New York: Buffalo. New York. Rhode Island: Providence. Tennessee: Chattanooga Washington: Tacoma West Virginia: Huntington.	0 0 0 3 0 0	1 2 1 2 8	1 3 1 1

DIPHTHERIA.

See p. 136; also Telegraphic weekly reports from States, p. 126, and Monthly summaries by States, p. 130.

INFLUENZA.

City.	Cases.	Deaths.	City.	Cases.	Deaths
Alabama: Birmingham		3	Minnesota: Minneapolis. St. Cloud.	1	
Los Angeles	3 2	1	New Jersey: Newark	11	
Florida: TampaGeorgia:	4	1	Trenton		
Atlanta Savannah	4 1	2 1	New York Niagara Falls.	53	
Illinois: Chicago Danville	13 1	4	Ohio: Akron	1	ļ
Kentucky: Covington			Cincinnati	<u>2</u>	
Louisiana: New Orleans Maine:		•••••	Toledo	1	
Auburn	1		Pailadelphia		
Baltimore	6	1	Charleston		
Brookline Pittsfield	1 1 1	i	El Paso Virginia: Roanoke		
SaugusMichigan: Detroit	3	1	West Virginia: Charleston		
:	, [,]	LEPR	osy.		1
California: Los Angeles	1		Ohio: Dayton	3	
	LETH	LARGIC E	NCEPHALITIS.		
California: San Francisco	2		Oregon: Portland		
		MALA	ARIA.		
Florida: Tampa	3		Tennessee: Memphis		1

MEASLES.

See p. 136; also Telegraphic weekly reports from States, p. 126, and Monthly summaries by States, p. 130.

PELLAGRA.

City.	Cases.	Deaths.	City.	Cases.	Deaths.
Alabama: Montgomery Georgia: Atlanta Missouri: Springfield		1	Pennsylvania: Philadelphia Texas: Waco.		1

New Orleans.

PNEUMONIA (ALL FORMS).

City.	Cases.	Deaths.	City.	Cases.	Death.
Alabama:			Kansas:		
Birmingham	.	.] 6		1	
Mobile		. 1		1	
Montgomery	2		Lawrence	1	·····
Arizona:	j		Topeka		. 2
Tucson		. 2	Kentucky: Covington	l	1 .
Arkansas: Fort Smith	ı	2	Louisville.	14	1 3
Little Rock	2	1 -	Louisiana:		
California:	1 -		Baton Rouge		. 1
Alameda	 	. 2	New Orleans		. 14
AlamedaBakersfield		2	Maine:	1	1
Relkeiea		. 1	Auburn		. 1
Los Angeles		. 2	Bangor Biddeford Portland	1	
Los Angeles	36	25	Biddeford		. 1
Oakland	8	5	Portland		. 2
Riverside		1	Sanford	1	
Sacramento		3	Maryland: Baltimore	36	24
San Bernardino		1 4	Cumberland	30	3
San Diego	6		Massachusetts:	-	
San Francisco	14	3	Attleboro		1
Santa Cruz	2	l i	Attleboro Beverly		i
Stockton		1 4	Boston		20
Colorado:		1 -	1 Deceletor	7	1 0
Denver		- 11	Cambridge	4	2 2 2
Connecticut:		·	Chelsea		2
Bridgeport	3	l	Chicopee		1
Bristol	6		Clinton		1
Manchester	2		Easthampton Everett	2	
Meriden	2	<u>.</u>	Everett		1
New Haven		5	Fall River.	• • • • • • • • •	5
Waterbury	3	2	Framingham Greenfield		1 1
Delaware:		3	Greenield	4	1 1
Wilmington District of Columbia:	•••••		Haverhill Holyoke	3	4
Washington		. 19	Lawrence		i
Florida:	••••••	. 10	Loomington		2
Tampa	2	1	Lowell	4.0	1 3
Jeorgia:			i Lvnn	7	3 3
Atlanta		13	Malden Medford	1	
Brunswick		1	Medford		1 2 3 1 2
Savannah		3	Melrose	• • • • • • • • •	2
Valdosta	1	• • • • • • • • • • • • • • • • • • • •	Melrose. New Bedford Newburyport Newton Pittsfield Plymouth Quincy Somerville Springfield Wakefield Wakefield	• • • • • • • • •	3
llinois: Alton	4	1	Newton	• • • • • • • • • •	1
Aurora		î	Pitteffuld	• • • • • • • • •	í
Bloomington		•	Plymouth		î
Blue Island		1	Quincy		$ar{f 2}$
Champaign	3 1		Somerville	2	
Chicago	199	52	Springfield		2
Chicago Chicago Heights Danville.		1	Wakefield	1	
Danville	3				
Decatur		1	Wostfield	••••	1
East St. Louis Elgin Evanston	• • • • • • • • • • • • • • • • • •	3 4	Worcester	••••••	8
Evanston		*	Michigan: Ann Arbor	1	
Fromert		····i	Detroit	41	21
Freeport		il	Flint	31	4
Jacksonville		2	Grand Rapids	5	í
Kewanee	i		Labramina	ĭ	-
La Salle	2		Jackson.	8	
Oak Park		2	Kalamazoo	8	1
Quincy	3		Pontiae	2	
Rockford	1		Port Huron		1
Rock Island	3		Minnesota:	i	_
Springfield	8		• Duluth		4
ndiana:	ا م	.	Faribault	••••••	1
EIRNART	2	1	Hibbing	•••••	2 5
Indianapolis	• • • • • • • • • • • • • • • • • • • •	4	Minneapous	•••••	5
Indianapous		11	Hibbing Minneapolis Rochester St. Paul		1 2
La Favatta		1 1 1	Missouri:	••••••	Z
Mishawaka		11	Independence	1	1
Muncie		3	Independence	18	10
South Bend		1	St. Joseph		3
		$\hat{2}$	Montana:		•
Terre Haute		21			
ndiana: Elkhart Gary Indianapolis Kokomo La Fayette Mishawaka Muncie South Bend Terre Haute ova: Council Bluffs		3	Billings	1 2	

PNEUMONIA (ALL FORMS)—Continued.

City.	Cases.	Deaths.	City.	Cases.	Deaths.
lebraska:			Ohio-Continued.		
Lincoln		. 4	East Cleveland	. 1	1
Omaha.] 9	Findlay	1	
New Hampshire:	• • • • • • • • • •	' "	Hamilton.	1	1 '
Concord		2	Kenmore		
			Tomoston		
lew Jersey:	_		Lancaster		·i :
Atlantic City	1		Lima		·1
Bloomfield	4		Mansfield		.
East Orange	2	1	Niles	4	1
Garfield	1	1	Norwood	.	
Hackensack	2		Piqua		.1
Harrison	3		Salem	1]
	, ,	3	Conductor		
Hoboken	· • • • • • • • • • • • • • • • • • • •		Sandusky		
Jersey City	2		Springheid		
Kearny	1		Toledo. Youngstown		·i
Montelair.		. 2	Youngstown	1	.t
Morristown	2	1 1	Zanesville		Ì
New Brunswick		4			1
			Obleheme City	1.0	ı
Newark	72	6	Oklahoma: Oklahoma City		1
Orange	6	1	n Olegon.		1
Passaic	3	2	Portland	l	!
Paterson	2		Pennsylvania:	t	
Perth Amboy	_	2	Philadelphia	74	5
Plainfield	4	3	Rhode Island:		1
r mainield	. 9	ľ			
Summit			Cranston		
Trenton	8	3	Providence		1
West New York		1	South Carolina:		1
West Orange	4	1	Charleston	1	
ew York:	-		Charleston	1	1
Auburn		1	South Dakota:	1	
Auburn	••••••		South Dakota.	l	! .
Buffalo	29	5	Sioux Falls		
Elmira	2	1.	Tennessee:	i	i
Glens Falls	1		Memphis	1	
Ithaca		1	Nashville		
Jamestown	5		Texas:		l
Mount Vernon	š		Dallas		
	•		Danas		
Newburgh	••••••	1	El PasoFort Worth		i
New York. North Tonawanda	381	165	Fort Worth		
North Tonawanda	1		Houston		
		2	Waco		!
Peekskill	2		Utah:		
Dent Charten	4		Salt Lake City		
Port Chester	. 4		Bait Dake City	•••••	
Pougn reepsie		1	Vermont:		
Rochester	13	14	Burlington		
Rome	3	1	Rutland	l	
Schenectady	6	5	Virginia:		
Syracuse	J	4	Norfolk	ا ا	,
Mass		4	Petersburg. Portsmouth Richmond.	••••••	'
Troy			Petersburg	• • • • • • • • • •	
White Plains		2	Portsmouth		
Yon ers	8	1	Richmond		
orth Carolina:		-	Roanoke		
Charlotte Durham Wilmington Winston-Salem		1	777 A 372		
Daraham		i	Divesold	1	
Durnam	• • • • • • • •		Bluefield		
Wilmington		2	Charleston		:
Winston-Salem		3	w neening		
nio:		,	Wisconsin:		
Akron	4				
	-	i	BeloitJanesville		
Alliance	•••••••	1	Kenosha		
Ashtabula	1		Kenosna		
	1		Milwaukec	9	
Barberton		4	Racine		
Barberton					
Barberton		12	Wyoming:		
Barberton		13	Wyoming:	g	
Barberton			Casper	. 5	:
Barberton		13		. 5 1	

POLIOMYELITIS (INFANTILE PARALYSIS).

The column headed "Median for previous years" gives the median number of cases reported during the corresponding weeks of the years 1915 to 1920, inclusive. In instances in which data for the full six years are incomplete, the median is that for the number of years for which information is available.

City.	Median for pre-			City.	Median for pre-	Week ended Dec. 31, 1921.		
	vious years.	Cases.	Deaths.		vious years.	Cases.	Deaths.	
California: San Francisco Illinois:	0	1		New York: New York Utah:	0	1		
ChicagoIowa: Cedar Rapids	0	1		Salt Lake City Washington: Bellingham	0	1	1	
New Jersey: Clifton		1				•		

SMALLPOX.

The column headed "Median for previous years" gives the median number of cases reported during the corresponding weeks of the years 1915 to 1920, inclusive. In instances in which data for the full six years are incomplete, the median is that for the number of years for which information is available.

City.	Median for pre- vious		k ended 31, 1921.	City.	Median for pre- vious		c ended 31, 1921.
	years.	Cases.	Deaths.		years.	Cases.	Deaths
Alabama:				Missouri:			
Mobile	0	4	1	Kansas City	5	18	11
California:	"		1 -	St. Louis	i	ĩ	1
Bakersfield	0	20	1	Nebraska:			1
Berkeley		ĩ	1	Omaha	7	2	1
Los Angeles		5		North Carolina:		_	1
Oakland	Ō	4		Winston-Salem	2	1	
Sacramento	l o	1		Ohio:			1
San Diego	0	1		Dayton	0	3	
Stockton	0	1		Fremont	0	3	
Colorado:	1			Sandusky	0	1	
Denver	9		2	Springfield	0	2	
Trinidad	l	1		Oklahoma:	1 1		1
Connecticut:				Oklahoma City	2	8	l
Bridgeport	0	5		Tulsa	2	3	
District of Columbia:	i i		i	Oregon:	i i		1
Washington	0	2		Portland	5	22	
Georgia:	1			Pennsylvania:	l 1		1
Atlanta	3	2		Philadelphia	0	1	
Savannah	0	2		Pittsburgh	0	2	
Illinois:	l i			South Dakota:			l
Centralia	0	2		Sioux Falls	0	1	
Chicago	1			Tennessee:			1
Freeport	0	1		Nashville	0	3	j
Peoria	7	4		Utah:	. !	_	
Indiana:	1			Salt Lake City	4	8	ļ
Bloomington	0	3		Virginia:	_		
Gary	0			Danville	0	1	
Indianapolis	6			Washington:	1		1
Logansport	0			Aberdeen		6	
Marion	0			Bellingham	4	2	
South Bend	0	1		Everett	0	2	
Iowa:			1	Seattle	3	2	•••••
Burlington	0	6		Spokane			
Cedar Rapids	2		• • • • • • • • • • • • • • • • • • • •	Tacoma	0	13	
Des Moines	2	2	• • • • • • • • •	Walla Walla	1	3	
Mason City	2			Yakima	5	4	· · · · · · •
Muscatine Kansas:	0	4		West Virginia: Bluefield	1	2	
Hutchinson	1	16		Parkersburg	ő	2	• • • • • • •
Kansas City	ō			Wiscensin:	V I	2	· · · · · · · · · · · ·
Leavenworth	ŏ			Janesville	0	1	
Michigan:	o į	*	• • • • • • • •	Manitowoc	ŏ	8	••••••
Detroit	8	2		Milwaukee	3	4	••••••
Highland Park	ől			Racine.	ő	i	••••••
Jackson	ŏ	il		Superior	2	4	· · · · · · •
Minnesota:	٠	*		Wyoming:	-	*	• • • • • • •
Hibbing	0	2		Casper	1	2	
Minneapolis	13	6		ousper		-	• • • • • • •
St. Paul	6	1		ł			
~ * am	•	-		i i			

TETANUS.

City.	Cases.	Deaths.	City.	Cases.	Deaths.
Georgia: Savannah Illinois: Chicago Louisiana: Baton Rouge Massachusotts: Boston	3 2 1	3 1 2 2	New Jersey: Trenton New York: New York Oklahoma: Oklahoma City		1 1 2

TUBERCULOSIS.

See p. 136; also Telegraphic weekly reports from States, p. 126.

TYPHOID FEVER.

The column headed "Median for previous years" gives the median number of cases reported during the corresponding weeks of the years 1915 to 1920, inclusive. In instances in which data for the full six years are incomplete, the median is that for the number of years for which information is available.

City.	Median for pre- vious		k ended 31, 1921.	City.	Median for pre-		ended 1, 1921.
-	years.	Cases.	Deaths.		vious years.	Cases.	Deaths
Alabama:				Missouri:			
Mobile	. 0	4		Joplin	0	. 1	
Arkansas:			1	Kansas City	1	1	1
Little Rock	0	1		St. Louis	5	1	
California:	0		l	New Jersey:		_	
Berkeley		1		Newark	0	2	
Long Beach		3		New York:	0	1	
Colorado:	ı "I			Buffalo	2	2	1
Trinidad	ll	1		New York	15	8	ļ
Connecticut:		-		Rochester	i	ž	İ
Norwich		2		Trov	ī	ī	
Georgia:	i i			Watertown	0	Ī	
Albany		1		Ohio:			
Savannah	0	1		Cincinnati	0	2	
Illinois:	ا ما			Cleveland	1	1	
Champaign	0	1		Columbus	0	1	
Decatur	0	1	·····i	Dayton Newark	0	1	
Quincy Indiana:	"	• • • • • • • •	-	Sandusky	ö	1	
Richmond	ا ه	1		Spring eld	ŏ	i	1
Kansas:	١٠١	•		Steubenville	ŏ	i	• • • • • • • • • • • • • • • • • • • •
Coffevville	lol	1		Toledo	ŏ		
Fort Scott	o l		1	Zanesville	ŏ	1	
Kentucky:				Pennsylvania:	1	_	
Covington	0	1		Philadelphia	3	1	
Louisiana:	_ [Pittsburgh	1	2	
New Orleans	2	8	1	South Carolina:			
Maine:	0	2	I	Charleston	0		2
Bangor	ŏ	1		Texas:	0	1	
Sanford	١	1	••••••	Dallas	0 !	2	
Baltimore	4	4	1	Fort Worth	ŏl	í	• • • • • • •
Massachusetts:	- 1	- 1	-	Utah:	•	•	••••••
Boston	1	2		Salt Lake City	0	1	
Brockton	ōl	· 1		Virginia:	- 1	_	
Brookline	o l	1		Roanoke	0 .		1
Gardner	0	1		Washington:		_	
Newburyport	0	1		Walla Walla	0	1	· · · • · · · ·
Waltham	0	1	•••••	Yakima	0	1	
Michigan:	اہ		- 1	West Virginia:	0	3	
Detroit	2	1	•••••••	Huntington Wisconsin:	٧	3	• • • • • • •
Minnesota: Minneapolis	1	1		Wisconsin: Green Bay	0	2	
St. Paul	il	4	•••••	Manitowoc	ő	í	• • • • • • •
Der Taut	* [*	• • • • • • • • •	maniwww	٠	- 1	• • • • • • •

· RABIES IN ANIMALS.

City.	Cases.
Missouri: Kansas City	1

SCARLET FEVER.

See p. 136; also Telegraphic weekly reports from States, p. 126, and Monthly summaries by States, p. 130.

DIPHTHERIA, MEASLES, SCARLET FEVER, AND TUBERCULOSIS.

	Popula- tion Janu-	Total deaths	Diph	theria.	Мо	asles.		arlet ver.		ber- osis.
City.	ary 1, 1920, subject to correction.	from all causes.	Cases.	Doaths.	Casos.	Docths.	Casos.	Deeths.	Cesos.	Deaths.
		¦				-	 	ļ	l	
Alabama: Birmingham	178, 270	58	2		1	j	2	į	3	4
Mobile	60, 151	19	lí				3		3	li
Montgomery	43, 464	14	ī	1						2
Arizona:			İ	-	İ	ĺ	İ	1	1	
Tucson	20, 292	18			·				ļ	10
Fort Smith	28,811	4	1			1	l	l	ł	1
Hot Springs	11,695	3	li			1				
Little Rock	64, 997						5			
California:			١ ـ	!		ĺ	1	ļ	l	Į
AlamedaBakers eld	28, 806 18, 638	11 9	1				i			·····i
Borkeley.	55, 886	9	3				2			
Eureka	12,923	9	ĭ				ļ <u>.</u>			
Glendale	13, 536	13								2
Long Beach	55, 593	27	68		4		24	2		;
Los AngelesOakland	576, 673 216, 361	191 74	27	1 4	3		5		62	14
Pasadena	45, 354	17	2.						ĭ	2
Richmond	16, 843	3	5	1					ļ <u>.</u>	
Riverside	19, 341	11	1				-		1	1
Sacramento	65, 857	25	6	1					2	3 1
San Bernardino San Diego	18, 721 74, 683	8 37		• • • • • •	····i	•••••	6		3	5
San Francisco.	508, 410	187	60	4			8		30	18
Santa Ana	15, 485	2	1						1	
Santa Barbara	19, 441	6				••••				
Santa Cruz	10, 917	6 12	·····	····i	····i		1 4	• • • • • •		• • • • • •
Colorado:	40, 296	12	9	•	•	•••••	· •	•••••	•••••	
Denver	256, 369	71		3						14
Greeley	10, 883	4								1
Connecticut:			8	3						_
Bridgeport Bristol	143, 538 20, 620	36 6	8	3	1	• • • • • •	14 1	•••••	4	. 8
Manchester (town)	18, 370	3						•••••	-	•••••
Meriden (city)	29,842								i	-
Milford (town)	10, 193	2					2			1
New Haven	162, 519	47	9	1	10	• • • • • •	9			2
New London Norwich (town)	25, 688 29, 685	6 7	1 2	•••••	4	•••••	1		····i	• • • • • •
Stonington (town)	10, 236	3					1			•••••
Waterbury	91,410	20	2	1	3		10		3	.
Delaware:				_	1					_
Wilmington	110, 168	28	1	1		•••••	14		• • • • • •	1
D strict of Columbia: Washington	437, 571	153	30	2	4		18	2	27	16
Florida:	101,011	100	۰ 🐱	- 1	- 1			- "	~ 1	10
Tampa	51, 252	31	4						1	3
Georgia:	1	1	_ 1		- 1	- 1		1		
Albany	11,555		1 6				8	•••••	:	•••••
Atlanta Augusta	200, 616 52, 548	73	1			•••••	i		3	5
Brunswick	14, 413	3								
Macon	52, 995	- 1	2							

CITY REPORTS FOR WEEK ENDED DEC. 31, 1921—Continued. DIPHTHERIA, MEASLES, SCARLET FEVER, AND TUBERCULOSIS—Continued.

	Popula- tion Janu-	Total deaths	1 -	htheria.	Me	asles.		arlet ever.		iber- osis.
City.	ary 1, 1920, subject to correction.	from all causes	Cases.	Deaths.	Cases.	Deaths.	Cases.	Deaths.	Cases.	Deaths.
Georgia—Continued.			1							
Rome	13, 252 83, 252					.		.		2
. Savannah	83, 252 10, 783	35	2	}			. 2		3	3
Idaho:	10, 705	0		1				-		-
Boise	21, 393	5	1		ļ		4	ļ	l	
PocatelloIllinois:	15, 001	3						.	ļ	
Alton	24, 682	7	1		l		7		l	l
Aurora. Bloomington.	36, 397	11	5				ļ		i	
Blue Island	28, 725 11, 424	14	3						 .	
Centralia.	12, 491	4	3	1						
Champaign	15, 873						2			
Chicago	2, 701, 705	575	170	13	43		108	4	131	52
Chicago Heights Danville	19,653 33,750	8	2				4		i	•••••
Decatur	33, 750 43, 818	8	7				i			····i
East St. Louis Elgin	66, 740	15 10	7	1			2			•••••
Evanston	37, 215	8	i		•••••		····i		····i	•••••
Forest Park	66, 740 27, 454 37, 215 10, 768			.	1		.			
Freeport	19,669 23,834	7	6 1	1						
GalesburgJacksonville. Kewanee.	15, 713	ģ			•••••		3 2		1	•••••
Kewanee	16,026	7		1			. .			
Le Salle	13, 050 13, 552	0	1		• • • • • •		• • • • • •			• • • • •
Oak Park	39, 830	12	5		3	•••••	4		····2	•••••
Pekin.	12,086		2				1			•••••
PeoriaQuincy	76, 121 35, 978	20 17	2 2				4 3		;-	•••••
Rockford	65, 651	14		1			3		1	1
Rock Island	35, 177	8]		i	• • • • • •
Springfield	59, 183	18	. 7				2		· · · · · ·	• • • • • •
Bloomington	11, 595	2								
Elkhart	24, 277	4	6				i			
FrankfortGary	11, 585 55, 378	3 14	<u>2</u>			• • • • • •	···· ₂ ·			• • • • •
Huntington	14,000	4					4			
Indianapolis	314, 194 30, 067	92	31 1	2	2		8		3	7
LaFayette	22, 486	7 10	i					•••••	···i	1
Logansport	21, 626	8							î i	i
Marion. Mishawaka	23, 747 15, 195	9 3	6			• • • • •	4			
Muncie	36, 624	12	•••••				····i			····i
Newcastle	14, 458	0					1			
Richmond	26, 765 70, 983	12 10	2 1		•••••	•••••	1	-	···i	••••
Terre Haute	66, 083	14	9	1	i		î			
Iowa: Burlington	24, 057	3			1		1	1		
Cedar Rapids	45, 566	0	···i							••••
Council Bluffs	36, 162	10	5							•••••
Des Moines Dubuque	126, 468 . 39, 141		5 3	2 .			12			••••
Iowa City	11, 267		ĭ							
Marshalltown	15, 731						3 .			
Ma on City Muscatine	20, 065 16, 068	4			••••• •	•••••	2 .	-		••••
Ottumwa	23,003 .		i				5			•••••
Sioux City	71, 227		9	-			4 .			•••••
Kansas:	36, 230 .				1 -	•••• •				
Atchison	12,630		2 2				3 .			
Coffeyville Fort Scott	13, 452	2	2	-	-	-				••••
Hutchinson	10, 693 23, 298	3	4		•••••		···· ₂ ·		···i	• • • • •
Kansas City	23, 298 101, 177		4				1 .			• • • • •
Lawrence. Leavenworth	12, 456	2	3		1 .		3 .		1 .	•••••
Parsons.	12, 456 16, 912 16, 028	7	2				1 .			····i
		- 4	- ,.				- 1.		,	-

CITY REPORTS FOR WEEK ENDED DEC. 31, 1921—Continued. DIPHTHERIA, MEASLES, SCARLET, FEVER, AND TUBERCULOSIS—Continued.

_	Popula- tion Janu-	Total deaths	Diph	theria.	Меа	sles.		arlet ver.		iber- losis.
City.	ary 1, 1920, subject to correction.	from all causes.	Cases.	Deaths.	Cases.	Deaths.	Cases.	Deaths.	Cases.	Deaths.
Kansas-Continued.										
SalinaTopeka.	15, 085 50, 022	7 26	7 9		i		5 1		5	i
Kentucky: Covington	57, 121	11	1	ļ			1		ļ	
LouisvillePaducah	234, 891 24, 735	59	20	2	102		2		9	i
Louisiana: Baton Rouge	21, 782	. g.	3	<u> </u>			2	 	1	1
New Orleans	387, 219	133	12	1	·····		8	ļ	12	12
AuburnBangor	16, 985 25, 978	5	2				5			
BathBiddeford	25, 978 14, 731	7								
Lewiston	18,008 31,791 69,272	4 3	3	i	i		2			i
PortlandSanford	69, 272 10, 691	23 1	2	<u> </u>			24			
Maryland: Baltimore.	733, 826	201	51	3	63		51	İ	15	21
Cumberland	29, 837	21	3	ļ			î			î
Massachusetts: Amesbury	10, 036	1		 		ļ	1	ļ		
Arlington Attleboro	18, 665 19, 731	3 8	11	1			1			1
Belmont	10,749	1	.2				ī		ļ <u>.</u>	
BeverlyBoston	22, 561 748, 060	9 214	67	4	1 51	····· <u>2</u>	52		1 48	9
BraintreeBrocton.	10, 580 66, 138	5 18	3 13	•••••			4		_i -	3 2
Brookline	37, 748	13	1				2		5	1
Cambridge Chelsea	109, 694 43, 184	32 16	2 2		4	•••••	3 4		3	2
Chicopee	36, 214	9 2	ī	1			1		1	2
Clinton Danvers	12, 979 11, 108	. Z			1		2		• • • • • •	
Dedham	10, 792 11, 261	4	•••••	•••••		•••••			····i	
Everett	40, 120	4	3		4		6		4	i
Fall RiverFraningham	120, 485 17, 033	32 6	4	1	• • • • • •		6 1	• • • • • •	4	1
Gardner	16, 971	4	1		9	•••••	3		1	i
Greenfield	15, 462 53, 884	5 16	····i		9	1	···i		····2	•
Holyoke. Lawrence.	60, 203 94, 270	17 19	3 5	1 2	*****		1 2	• • • • • •	1 1	1
Leominster	19, 744	4	3				2		1	i
LowellLynn	112, 479 99, 148	26 27	5 8	1 1	2		····i		2	1 1 2
Malden Me ford	49, 103 39, 038	11 11	9	•••••	14		12 1		1	2
Melrose	18, 204	7			1					•••••
Methuen New Bodford	15, 189 121, 217	5 25	17	····i	4		7		4	····· <u>·</u>
Newburyport	15, 618	25 7					₂			····
Newton	46, 054 22, 282	12 8	8				1		1	•••••
Northampton	21, 951 19, 552	11 3		•••••		•••••			•••••	i
PeabodyPittsfield	41, 751	14	4				3	i	···i	2
PlymouthQuincy	13, 045 47, 876	3 7	3		17		13		3	• • • • • • • • • • • • • • • • • • •
SaugusSomerville	10, 874 93, 091 14, 245 129, 563 13, 025	2 19	3 1 7	1	2		5		• • • • • •	••••••
Southbridge	14, 245	0			î i		! .		4	•••••
Springfield	129, 563	36	3 4	•••••			7	1	7	5
Waltham	90,919 [8					4			•••••
Watertown Webster	21, 457 13, 258	5 2	2	:::::		:::::i.	3	:::::	3	· · · · · •
Webster	13, 443	3 11			2	-				•••••
Westfield	18, 604 15, 057	i i								· · · · · · · ·

CITY REPORTS FOR WEEK ENDED DEC. 31, 1921—Continued. DIPHTHERIA, MEASLES, SCARLET FEVER, AND TUBERCULOSIS—Continued.

	Popula- tion Janu-	Total deaths	Diph	theria.	Mea	ısles.		arlet ver.		ber- osis.
City.	ary 1, 1920, subject to correction.	from all causes.	Cases.	Deaths.	Cases.	Deaths.	Cases.	Deaths.	Cases.	Deaths.
Massachusetts—Continued.		1	1	1	1			1		
WinthropWoburn	15, 455 16, 574	4	ļ	· ·····						• • • • • •
Worcester	179, 754		4	i			7	i	3	6
Michigan: Ann Arbor.	19, 516	10	5		1			ł		1
Ann ArborBattle CreekBenton Harbor	36, 164		5		25		2			
Benton Harbor Detroit	12, 233 993, 739	188	99	5	67	····i	62	····i	41	20
Flint	91, 599	17	7	1		ļ <u>.</u>	9	ļī.		
Grand Rapids	137, 634 48, 615	30	10	4	·····		7	·····	1	· · · · · •
Hamtramck Highland Park	46, 499	10	î				i			
Ishpeming	10, 500 48, 374	18	2	.		• • • • • • • • • • • • • • • • • • • •	18			
Kalamazoo.	48, 858	13	15		i		21			.
Marquette	12,718	10	4	i						-
Pontiac Port Huron	34, 273 25, 944	9	2	ļ <u>.</u>			6		i	
Sault Ste. Marie	12, 096	4			•••••		2		1	1
Minnesota: Duluth	98, 917	21	2				3		7	2
Karihault	11,089	7	l	1						
Hibbing	15, 089 12, 469	3	1		1	• • • • •				
Minneapolis	380, 582	85	21	i	ii		37	1	9	7
Rochester	13, 722 15, 873	12			• • • • • •	•••••	1 4			1
St. Cloud	234, 595	43	ï	2	2		15		7	2
Winona	19, 143					• • • • • •	5		1	
Missouri: Independence	11,686	2	4							
Joplin	29, 855		3				4			•••••
Kansas City	324, 410 77, 9.9	136 30	27 1	3	1	•••••	15 3	1	17	11 3
St. JosephSt. LouisSpringfield	772,897	206	70			1	25		18	8
Springfield	39, 631	13	• • • • • •							. 3
Anaconda	11,668	5								
Billings	15, 100	3 5	<u>2</u>				3			•••••
Great Falls	24, 121 12, 668	7					····i		1	····i
Nebraska:	· 1	10					ا ،	1	- 1	
LincolnOmaha	54, 934 191, 601	12 46	2 6	····i	20		3 1			i
Nevada:		_		-			-			
Reno New Hampshire:	12, 016	2	•••••	•••••	•••••	•••••	•••••	•••••		· · · · · •
Berlin	16, 104	4								-
Concord	22, 167 13, 029	9	i				2		1	
Keene	11, 210	4								
New Jersey: Asbury Park	12,400	2		.	.	l	1	1	1	
Atlantic City	50, 682	9	2				2		i	
Bayonne	76, 754		6			•••••	3		3	-
BellevilleBloomfield.	15,660 22,019	2	····2		···i		4			
Clifton	26, 470	3	2				2			.
East Orange	50, 710 11, 627	6	2		···i	•••••	7		2	
EnglewoodGarfield	19, .81	3							î	•••••
Hackensack	17,667 15,721	4	•••••		•••••		3 5			· · · · · ·
Harrison	68, 166	14	3			· · · · · · · ·	1			i
Jersey City.	297,864 26,724		15		23		9		10	····i
Kerny	26, 724 28, 810	6 8	3				4		1	i
Morristown	19 549	8 7 12	1							· · · · • •
New Brunswick Newark	32, 779 414, 216 33, 268 63, 824	108	28	••••	49	• • • • • • •	09	· · · i	33	· · · · • •
Orange	33, 268	6			1 .		9			· · · · •
Passaic	63, 824	16	5	1	-		5		1).	••••

CITY REPORTS FOR WEEK ENDED DEC. 31, 1921—Continued. DIPHTHERIA, MEASLES, SCARLET FEVER, AND TUBERCULOSIS—Continued.

	Popula- tion Janu-	Total deaths	1 -	htheris	. Me	asles.		arlet ver.	T	uber- losis.
City.	tion Janu- ary 1, 1920, subject to correction.	from all causes	,,	Deaths.	Cases.	Deaths.	Cases.	Deaths.	Cases.	Deaths.
New Jersey—Continued. Paterson Perth Amboy	135, 866 41, 707	12			. 2		. 2		. 3	
Phillipsburg Plainfield Rahway Summit	16, 923 27, 700 11, 042 10, 174	3 8 1 4	2		i		8 2		i	
Trenton Union West Hoboken	119, 289 20, 651 40, 068	32	. 4		1		14 1 3		5	
West New York	29, 926 15, 573	3 3		-	: '''i'		1 4			
Albuquerque	15, 157 36, 192 66, 800	10 15 11	3 2	1			1 11			1
Buffalo Elmira Geneva.	45, 305 14, 648	113 20 5	49 1	1	17		36 2	2 1	28	13 1 1
Glens Falls	16, 638 15, 025 17, 004 38, 917	10 5 11	2 1 2 6	1		1	4		1 2	i
Little FallsLockport	13, 029 21, 308 42, 726 30, 366	2 7 10	1		i		3 5		2 2	
Newburgh New York Niagara Falls North Tonawanda	30, 366 5, 621, 151 50, 760 15, 482	13 1,364 7 5	276 6 2	25	269 2	5	283 14 1	3	1 137 1	1 89
Olean	20, 506 15, 868 16, 573	10 5 6	í		3	•••••			1	
Poughkeepsie Rochester Rome	35, 000 295, 750 26, 341	5 69 11 3	1 23 3				5 7		11	3 1
Saratoga Springs Schenectady. Syracuse Troy Watertown	13, 181 88, 723 171, 717 72, 013	56 26	2 32 2	2			7 16	1	1 3 11 2	1 3 1
White Plains Yonkers	31, 285 21, 031 100, 226	10 4 26	2 1 5	i	1		6 3 13	1	1	•••••
North Carolina: Charlotte Durham Greensboro.	46, 338 21, 719 19, 861	15 8 7	4				1 2		1	1 2
Rocky Mount	12, 742 13, 884 33, 372	1 1 15								·····ż
Winston-SalemOhio: AkronAlliance	48, 395 208, 435 21, 603	18 34 6	2 4 1	1	2		3 22 1		2	
Ashtabula Barberton Bucyrus	22, 082 18, 811 10, 425	5 5 1	2 1 1				i		2	· · · · · · · ·
Canton Chillicothe Cincinnati Cleveland	87, 091 15, 831 401, 247 796, 836	26 2 121	17 1 22 35	<u>1</u>	12 38		3 1 3 47		9	12
Columbus Dayton East Cleveland	237, 031 152, 559 27, 292 20, 474	68 37 3	11 4	1	1		6 4 2		5 1	4
ElyriaFindlayFreemont	12, 468	3 8 1 7	į				ī			•••••
Hamilton Kenmore Lancaster Lima	39, 675 12, 683 14, 706 41, 306	7 7 7	5 6 3 3	1	1		1 1		1	1

¹ Pulmonary tuberculosis only.

CITY REPORTS FOR WEEK ENDED DEC. 31, 1921—Continued. DIPHTHERIA, MEASLES, SCARLET FEVER, AND TUBERCULOSIS—Continued.

	tion Janu- deaths		1 -	theria.	Ме	Measles.		Scarlet fever.		iber- losis.
City.	ary 1, 1920, from subject to all	from all causes.		Deaths.	Cases.	Deaths.	Cases.	Deaths.	Cases.	Deaths.
Ohio-Continued.					<u> </u>	l				
Lorain	37, 295 27, 824	ii		-	. 3		. 7		. 1	
Marion	27, 891	11	4	-			3			1
Middletown	23,594	1	ī							
New Philadelphia	26, 718	13	. 5		2		8 2		• ••••	. 1
Niles	10, 718 13, 080	3					2		i	
Norwood	24, 966	5	ļī.	.			2			
Piqua	15,044	3		.		·		.	. 4	1
Salem	10, 505 22, 897	10		·			2			
Sandusky Springfield Steubenville	60, 840	17	ii				l î			
Steubenville	28, 508	9					- 1		1	
Tiffin	14, 375	5			i		····			
Toledo Youngstown	243, 109 132, 358	59	34	2			7 9	2	. 1	1 1
Zanesville.	29, 569	12	4				2			1
Oklahoma:				1			1			
Oklahoma City	91, 258	. 29	5				6			3
Tulsa	72,075		6				4			
Oregon: Portland	258, 288	75	24	1	l		12	1	4	4
Pennsylvania:	•			1					_	
Allentown	73, 502		1 7				1 4			
AltoonaBethlehem	60, 331 50, 358		3				9			• • • • • • • • • • • • • • • • • • • •
Braddock	20, 879		2							
Bradford	15, 525		1							
Bristol	10, 273 10, 504		1 1		·····2					-
Carrick	19,011		2				5			
Easton	33, 813		ļ .				ĭ		i	
Erie	93, 572		7		1		3		5	
Farrell. Harrisburg.	15, 586 75, 917	• • • • • • • •	4	• • • • • •	1		5			
Harrisburg. Hazleton	32, 277		7				i			
Johnstown	32, 277 67, 327		9		5		2			
Lancaster	53, 150		3				4			· · · · · ·
Lebanon	24, 643 45, 975		4		15	• • • • • •				-
McKees Rocks	16, 713		12		1		3		1	
Mount Carmel	17, 469		2							
Nanticoke	22,614	• • • • • • • •	6 7		• • • • • •		•••••			
New Castle Norristown	44, 958 32, 319	• • • • • • • •	2		• • • • • •		4	• • • • • • •	• • • • • •	
Philadelphia	1, 823, 158	503	76	7	6		158	4	61	30
Pittsburgh	588, 193		41		11		49		7	
PottstownReading.	17, 431	•••••	3 6	• • • • • •		• • • • • •	16		• • • • • •	-
Seranton.	107, 784 157, 783		12				4			· · · · · •
Shamokin			2				î			· · · · · · ·
Sharon	21, 747 21, 747 24, 726		1		35					· · · · · ·
Shenandoah	10, 908	•••••	1				•••••		•••••	• • • • •
Tamaqua	12, 363		3							• • • • • •
Uniontown	15, 692				1		1			
Washington	21, 480		3		1		3			· · · · · •
Wilkes-Barre	73, 833 12, 495		6		19		4			•••••
York.	47, 512		3				5			
Rhode Island:			- 1				-			· · · · · · ·
Cranston	29, 407	3			•••••		2			1
Pawtucket Providence.	64, 248 237, 595	14 78	3 12		5		1 2			1 3
South Carolina:	-				١		-			0
Charleston	67, 957	30	2							1
Columbia	37, 524 23, 127	·····;·	2				4		1	· · · · · ·
South Dakota:	20, 121	3					•••••			•••••
Sioux Falls	25, 176	12	2							
Chattanage		i	- 1			- 1	ا ،		- 1	
ChattanoogaKnoxville.	57, 895 77, 818		····i		•••••		2	-	···· ₂ ·	·····2
	11,010		1].	•••••	•••••	!	4	1	21	Z

CITY REPORTS FOR WEEK ENDED DEC. 31, 1921—Continued. DIPHTHERIA, MEASLES, SCARLET FEVER, AND TUBERCULOSIS—Continued.

***	Popula- tion Janu-	Total deaths	1	theria.	Ме	Measles.		arlet ver.		Tuber- culosis.	
City.	ary 1, 1920, subject to correction.	from all causes.		Deaths.	Cases.	Deaths.	Cases.	Deaths.	Cases.	Deaths.	
Tennessee—Continued.											
Memphis	162, 351	51	21		.		. 6		. 5	3	
Nashville	118, 342	53	4	2	1		. 2		. 1	4	
Texas: Beaumont	40, 422	6	1 1	1		1 .	1	1	1	1	
Corpus Christi	10 599	4	l î								
Dallac	158, 976 77, 543 106, 482 44, 255 138, 076	43			22		2		. 2	4	
El Paso	77, 543	36	. 4	····i			1		i	9	
Galvestan	44, 255	9	0	1			1		1		
Houston	138, 076	i 29	3	1			3	1		6	
Waco	38, 500	18			ļ				ļ	1	
Utah: Salt Lake City	118, 110	. 23	3		1		16	1.	1	l	
Vermont:	110, 110	~	1 "		٠ .		10				
Barre	10,008					ļ	1				
Burlington	22,779	5	1				3				
RutlandVirginia:	14, 954	7						ļ			
Alexandria	18,060	. 3		1		l	l	l	l	1	
Danville	21, 539	4	2							ļ <u>.</u>	
Lynchburg Norfolk	29, 956 115, 777	. 6	3						2	···· <u>·</u>	
Norioik	31 002	19	6	1		••,•••	3		2	6 3	
Petersburg Portsmouth	31,002 54,587 171,667	17	1				2				
Richmond	171,667	48	6		47		5		14	2	
Roanoke	50, 842	17	8	1			1			1	
Washington:	07 644		3					i	ĺ	1	
Everett	27, 644 315, 652		7		i		7			•••••	
Spokane	104, 437		14				2				
Tanoma	96, 965	•••••	6				3		2		
Walla Walla	15, 503	••••••	2 2		1	••,•,•••	2	• • • • • •		·····	
Yakima	18, 539	•••••	, z		• • • • • • •	• • • • • •	Z			•••••	
West Virginia: Bluefield Charleston	15, 282	4	2		1						
Charleston	39, 608 17, 851	14	3 7				3	1			
Fairmont. Huntington. Morgantown.	17, 851	.1	7	1			1				
Huntington	50, 177 12, 127	11	1			•••••	1			2	
Moundsville	10, 669	3	<u>.</u>						•••••	•••••	
Parkersburg	20, 050 1	6	2								
Wheeling	54, 322	16				•••••	2		• • • • • •	1	
Wisconsin: Appleton	19, 561		4				1				
Ashland	11, 334		i								
Roloit	21, 284	4	3				3			•••••	
Eau Claire	20,880	•••••			1		1			•••••	
Eau Claire. Fond du Lac. Green Bay. Janesville	23, 427 31, 017	6 5	6		····i				····i	•••••	
Janesville.	18, 293	6	5				i		î		
Kenosna	40, 472	10	21	2			3		2	•••••	
Madison	38, 378	6	10				4			•••••	
Marinette Milwaukee	13, 610 457, 147 33, 162	••••••	1 37		····2		1 16		27	•••••	
Oshkosh	33, 162	5									
Racine	58, 593	23	7				18		2	3	
Sheboygan Superior	30, 955	•••••	;-			•••••	1			•••••	
Superior	39, 624 18, 661	13	1 3				10		····i	•••••	
West Allis.	13, 765		î								
Wyoming:	· 1		-								
Casper	11,447	8		1					3	1	
Cheyenne	13, 829	4	1	1			3				

FOREIGN AND INSULAR.

AUSTRALIA.

Plague—Queensland.1

Plague has been reported in Queensland, Australia, as follows:

Brisbane.—Week ended November 5, 1921, five cases with two deaths; 8 rodents and one guinea pig found infected; week ended November 19, 1921, six cases with two deaths; 10 rodents and one cat found infected.

Cairns.—Week ended November 5, 1921, one case. Three rodents found infected; week ended November 19, 1921, two cases with one death.

Cooktown.—Week ended November 5, 1921, one case (pestis minor).

Port Douglas.—Week ended November 19, 1921, one case with one death.

CUBA.

Communicable Diseases - Habana.

Communicable diseases have been reported at Habana as follows:

	Dec. 21-	Dec. 21-31, 1921.		
Disease.	New cases. Deaths.		ing under treat- ment Dec. 31, 1921	
Chicken pox. Diphtheria. Infantile tetanus. Leprosy. Malaria. Poliomyelitis (infantile paralysis). Scarlet fever.	3 1 1 1 23 1 3	1 2 1	3 2 11 124	
Typhoid fever	. 8	3	2 16	

¹ From the interior 10.

INDIA.

Animal Anthrax-Burma.

During the month of October, 1921, outbreaks of animal anthrax were reported in two districts of Burma by the Veterinary Department.

² From the interior 14.

¹ Public Health Reports, Jan. 13, 1922, p. 75.

MEXICO.

Plague-infected Rodents - Tampico.

The finding of plague-infected rodents at Tampico, Mexico, has been reported as follows: Week ended December 31, 1921, three infected rodents; total plague-infected rodents found from January 1 to December 31, 1921, 322. Week ended January 7, 1922, plague-infected rodents found, one.

POLAND.

Cholera - Smallpox - Typhus Fever - Aug. 14-Sept. 10, 1921.

During the period August 14 to September 10, 1921, cholera, small-pox, and typhus fever were reported in Poland as follows:

Cholera.—Four cases, with one death.

Smallpox.—Cases, 102, with 24 deaths; highest mortality reported in Kracow.

Typhus fever.—Cases, 738, with 52 deaths; highest mortality reported in Stanislawow.

The statistics do not include reports from Brest-Litovsk, Minsk, and Wilno districts.

UNION OF SOUTH AFRICA.

Anthrax-Transvaal.

A fatal case of anthrax occurring in a European resident of Maraisburg, Transvaal, has been reported as occurring August 19, 1921. It is stated that the deceased developed an anthrax sore (malignant pustule) on the chin a few days after purchasing a shaving brush of Japanese origin, which on bacteriological examination was found to be anthrax infected, facts indicating that the fatal infection was conveyed by the brush.

During the week ended November 4, 1921, an outbreak of anthrax was reported on the Vlakplaats farm, Heidelberg, Transvaal, where several fatal cases of animal anthrax had been previously noted. In a group of 21 persons belonging to two families who appear to have eaten meat from an infected animal, 12 cases of anthrax with four deaths have been reported.

Infection Found in Shaving Brushes.

Early in the year 1920 consignments of shaving brushes of Japanese origin at Cape Town, East London, and Bleemfontein were found to contain anthrax infection. These consignments were suitably dealt with, and further importation was prohibited; but many firms in the Union had already large stocks of these brushes.

CHOLERA, PLAGUE, SMALLPOX, TYPHUS FEVER, AND YELLOW FEVER. Reports Received During Week Ended Jan. 20, 1922.

CHOLERA.

Place.	Date.	Cases	B. Death	Remarks.
India				Oct. 9-15, 1921: Deaths, 4,174.
Calcutta	Nov. 20-26	. 1	6 1	3
Rangoon Poland	do		1	1 Aug. 14-Sept. 10, 1921: Cases, 4; deaths, 1.
	 PLA	GUE.		1
Australia:	T -	Ī.	T	
Queensland— Brisbane	Oct. 30-Nov. 5			2 8 rodents and 1 guinea pig found
Do	Nov. 13-19			infected. 2 10 rodents and 1 cat infected.
Cairns	. Oct. 30-Nov. 5	1		3 rodents.
Do Cooktown.	Nov. 13-19 Oct. 30-Nov. 5	2		Pestis minor.
Port Douglas	Nov. 13-19	i		resus minor.
Azores:	1		1	
Livramonto	. Dec. 4-10do	1		Vicinity of Ponta Delgada.
Ribeira Grande	. do	9	1 3	9 miles from port (Ponta Del- gada).
Brazil: Bahia	Nov. 6-12	1	1 1	
Ceylon:	1101.0-12	.	.	
Colombo Egypt			-	Nov. 20-26, 1921: 1 rodent plague. Jan. 1-Dec. 15, 1921: Cases, 341;
Alexandria	Dec. 12	i		deaths, 143.
Ålexandria	Dec. 9-14	Ī		. [
India	• • • • • • • • • • • • • • • • • • • •		-	Nov. 13-19, 1921: Cases, 1,199; deaths, 869.
Bombay	Nov. 13-19 Nov. 27-Dec. 3	1		.1
Madras Presidency Rangoon	Nov. 21-Dec. 3 Nov. 20-26	326 10		
Indo-China:	100.20-20	10	8	
Saigon			.	. Nov. 20-26, 1921; 1 rodent plague.
Mexico:			1	1
Tampico		••••••••••••••••••••••••••••••••••••••		Dec. 25-31, 1921: Plague-infected rodents, 3; total, Jan. 1-1 ec. 31, 1921: 322. Jan. 1-7, 1922, 1 plague-infected rat.
	l i Smali	- DOT		
	SMAL	LPUX.	· · · · · · · · · · · · · · · · · · ·	
Brazil: Bahia	Nov. 6-12.	1		
Zanada:	1101.0-12	1		
New Brunswick— Counties—				
Charlotte	Dec. 18-24.	3		
Restigouche	do	ĭ		
Ontario—	1	_		
Ottawa Toronto	Jan. 1-7do	3		
hile	do	12	•••••	Nov. 15-21, 1921: Diffused in
`			•••••	southern provinces; not epi- demic.
Concepcion	Nov. 15-21		•	Present. In vicinity, at Hual- qui, 32 cases, 5 deaths.
Coronel	do			Cases numerous.
Curanilahue	Nov. 13–19.	4		
Talcahuano	Nov. 13-19	4		
Temuco	Nov. 15-21	9		
Prestonominican Republic:	Dec. 25-31	2		
Santo Domingo				Dec. 17-27, 1921: Cases 631, in
inland				Surrounding country. Nov. 16–30, 1921: I case.
aiti: Cape Haitien	Dec. 11-24	8		
Port au Princedia:	Dec. 25-31			Present.
Calcutta	Nov. 20-26 Nov. 27-Dec. 3	3 28	2	
	Public Health Service	40 l	7 l 	ula and other sources

¹ From medical officers of the Public Health Service, American consuls, and other sources.

Reports Received During Week Ended Jan. 20, 1922 - Continued.

SMALLPOX-Continued.

Place.	Date.	Cases.	Deaths.	Remarks.
Japan:				
Taiwan Island	. Dec. 1-10	1		
Java:	i		1	j .
West Java—	37. 10.01	١.	i	l i
BandoengBatavia.	Nov. 18-24	1 2	ļ	!
Buiten org	Nov 25 Dec 1	2	2	
Krawang				i
Lebak	Nov. 18-Dec. 1	1 4		Í
Pandezlang	Nov. 25-Dec. 1		l ī	
Tangerang	Nov. 18-Dec. 1	3	ĺ	ĺ
Mexico:	1		_	
Torreon	Dec. 1-31		134	
Poland		!		Aug. 14-Sept. 10, 1921: Cases, 102; deaths, 24. Exclusive of
	1	l	l	102; deaths, 24. Exclusive of
	ļ		ļ	Brest-Litovsk, Minsk, and
Cnains	1	l ··	l	Wilno districts.
Spain: Huelva	Oct. 1-31	İ	1	
Tunis:	1			
Tunis	Dec. 10-16	6	6	
Union of South Africa:	1	1		
Cape Province	Nov. 5-19 Oct. 23-Nov. 12			Outbreaks.
Natal	Oct. 23-Nov. 12			Do.
Orange Free State	Oct. 23-29		l .	Do.
Transvaal	Oct. 23-Nov. 19	• • • • • • • •		Do.
	TYPHUS	FEVE	R.	•
Oh i				
China: Harbin	Nov 28-Dog 4	1		
Mexico: San Luis Potosi	Dec 25_31			Present.
Poland				Aug. 14-Sept.10, 1921: Cases, 738;
				deaths, 52. Exclusive of Brest-
		- 1		Litovsk, Minsk, and Wilno dis-
Union of South Africa:				tricts.
Cape Province	Oct. 23-Nov. 12			Outbreaks.
Natal	Oct. 23-29		•••••	Do.
	YELLOW	FEVE	R	
Movino:				
Mexico: Tierra Blanca	Nov. 6-12	, ,	1	State of Vera Cruz.
Mexico: Tierra Blanca Vera Cruz.	Nov. 6-12 Dec. 20-26	1	1	State of Vera Cruz. Imported.

Reports Received from Dec. 31, 1921, to Jan. 13, 1922.1

CHOLERA.

Place.	Date.	Cases.	Deaths.	. Remarks.
India				Oct. 2-8, 1921: Deaths, 6,374.
Bombay	Oct. 30-Nov. 5		<u>-</u>	
Calcutta	Oct. 23-Nov. 19	14	11	
Karachi	Oct. 23-Nov. 12		1	
Rangoon	Oct. 1-Nov. 19	6	4	
Indo-China:	1			
Saigon	Nov. 6-12	1	1	
Java:				· ·
West Java-	•			
Batavia	Nov. 1-7	2	2	At Lebak.
Philippine Islands:		_	_	
Manila	Nov. 13-Dec. 2	4	1	
Siam:	2.020	-		
Bangkok	Oct. 23-29	1		
Dangava	000.20-20	-	• • • • • • • • • • • • • • • • • • • •	

¹ From medical officers of the Public Health Service, American consuls, and other sources. For reports received from July 2 to Dec. 30, 1921, see Public Health Reports for Dec. 30, 1921. The tables of epidemic diseases are terminated semiannually and new tables begun.

Reports Received from Dec. 31, 1921, to Jan. 13, 1922—Continued.

PLAGUE.

Place.	Date.	Cases.	Deaths.	Remarks.
Asia Minor: Smyrna	Nov. 27-Dec. 3	1	1	
Sydney		ļ		Nov. 6-19, 1921: Plague rats re ported found at distance fron wharves.
Queensland— Brisbane	Nov. 6-26	6	3	Plague-infected rats, 16. Tota cases of plague, Aug. 22-Nov 26, 1921, 29; deaths, 18.
Cairns Ingham Townsville	Nov. 20-26 Nov. 6-12 Nov. 20-26	1 1	1	One plague rat. Nine plague rats.
Azores: St. Michael Island	N 08 D 0			Nov. 27-Dec. 3, 1921; Cases 5 deaths, 5.
Fenaes d'Ajuda Ribeira Grande Brazil: Bahia	Nov. 27-Dec. 3 Nov. 13-Dec. 3 Oct. 30-Nov. 5	10	5 4	Present.
British East Africa: Uganda	Aug. 1-Sept. 30	1	58	Reports of inspectors, deaths, 142; reports of chiefs, deaths
Ceylon: _ Colombo	Oct. 30-Nov. 5	. 1		641. Rodent plague, 2.
Ecuador: Guayaquil	Dec. 1-15	2		Rats examined, 1,458; found infected, 41. Jan. 1-Dec. 8, 1921; Cases, 336
City— Alexandria	Dec. 5-6.	2		deaths, 142.
Suez Province— Keneh	Nov. 22-Dec. 5 Dec. 1	4 1	2	Septicemic.
India Bombay Karachi	Oct. 23-Nov. 12 Nov. 6-19.	3	2 1 392	Oct. 23-Nov. 12, 1921: Cases, 3,244; deaths, 2,398.
Madras Presidency Rangoon Indo-China: Saigon	Nov. 13-26 Oct. 1-Nov. 19	600 53	50	Nov. 6-12,1921; Rodent plague, 1,
Italy: Naples (Province)— Torre Annunziata	Oct. 22.	1		2.00.00 -2,1.02-, 2.000-0 p-u8u0, 2.
Venice	Oct. 27 Oct. 30-Nov. 5	37	31	
BagdadMexico: Tampico	Oct. 1-31	1	1	Dec. 18-24, 1921: Infected rodents
Vera Cruz				found, 2; total, Jan. 1-Dec. 24, 1921; infected rodents, 319. One infected rodent caught Dec. 5, 1921.
Peru				Nov. 17-30, 1921: Cases, 48; deaths 12. Occurring in Cal-
Portuguese West Africa:		•	•	lao, Huacho, Huaras, Lima, Magdalena Vieja, Paita, Sala- verry, and Sechura.
Angola— Loanda	Oct. 9-Nov. 5 Oct. 13	3	2 1	
Siam: BangkokStraits Settlements:	Oct. 23-Nov. 5	1	1	
Singapore	Nov. 6-12 Oct. 9-Nov. 13	9	3	•

Reports Received from Dec. 31, 1921, to Jan. 13, 1922—Continued.

SMALLPOX.

Place.	Date.	Cases.	Deaths.	Remarks.
Bolivia:	1.01.01	42		
La Paz Brazil:	Aug. 1-Oct. 31	1 42	28	
Rio de Janeiro	Nov. 13-26 Oct. 31-Nov. 20	4 2	2	
British East Africa: Uganda	Aug. 1-Sept. 30	7		Reports of inspectors, cases, 4.
Canada: Manitoba— Winnipeg	Nov. 20-Dec. 3	2		
New Brunswick— Charlotte County	1101. 20-20. 0	_		Dec. 17, 1921: 31 cases previously
St. Stephen Restigouche County	Dec. 11-17do.	2 1		reported, occurring at Ander- sonville and Blacks Harbor.
York County Ontario—	do	1		
Niagara Falls Ottawa	do	17		,
TorontoQuebec—	do	.1	···········	
Montreal Saskatchewan—	do	1		,
Saskatoon	Dec. 1-18	6		
Valparaiso	Oct. 23-Nov. 26		34 1	Nov. 92 90 1001: Present
AmoyAntungChungking	Nov. 16-22 Nov. 28-Dec. 4 Nov. 6-19	2		Nov. 23-29, 1921: Present. Present.
Foochow	Nov. 6-26 Nov. 13-Dec. 3			Do. Do.
Harbin	Nov. 14-27	3		Do.
Nanking. Shanghai.	Nov. 20-28 Nov. 20-Dec. 3 Oct. 31-Dec. 4	25	27	Do. Cases, foreign; deaths, Chinese.
Colombia: Cartagena	Nov. 22-28.	-	1	outer, integri, deutino, emittee.
Cuba: Antilla	Dec. 12-18	1	-	At Preston.
Dominican Republic: San Pedro de Macoris	Nov. 20-26	5		An estimate of 500 cases of small- pox in the district of Macoris; of this amount 50 are within
Santo Domingo	Nov. 15-Dec. 5	401		the city limits. In district 146: estimated.
Ecuador: Guayaquil	Dec. 1-15	2		Venecia and San Carlos haciendas.
Egypt: Alexandria	Nov. 26-Dec. 2	1	1	
Haiti: Port au PrinceIndia	Dec. 11-24			Present. Oct. 2-8, 1921; Deaths, 28.
Calcutta	Nov. 13-19	3	3 1	Oct. 2-5, 1821, 17catilis, 25.
BombayCalcutta	Oct. 23-Nov. 12 Oct. 23-Nov. 19	1 6	4	·
MadrasRangoon	Nov. 13-26 Oct. 1-Nov. 19	22	8	
Italy: Genoa Messina—	Nov. 10-20	1		
Messina	Nov. 28-Dec. 4 Nov. 14-Dec. 4	1 2		
Mesopotamia: Bagdad	Oct. 1-31	24	7	
Mexico: Chihuahua	Dec. 5-11	إإ	1	•
Guadalajara	Nov. 1-30 Nov. 20-26 Dec. 18-24	12	2	
Panama:	_ 30. 10 =1			Admitted to hospital by transfer
				from Panama, Nov. 30, 1921, 1 case. Arrived on sailing vessel
Chiriqui Province	Dec. 22.		l	from a village on south coast. Present.

Reports Received from Dec. 31, 1921, to Jan. 13, 1922—Continued.

SMALLPOX—Continued.

Date.	Cases.	1	
I	Casus.	Deaths.	Remarks.
. Dec. 14	1		On December 21, 1921; 1 add tional case from country di trict of Sabanas, admitted i hospital. Total admission Jan. 1-Dec. 21, 1921, 207.
Nov. 13-26	12	5	
1	ļ	4	
Oct. 9-Nov. 5		. 3	
Oct. 1-31	20		
	i	4	
Oct. 23-Nov. 5	1	ļ	
Nov. 1-30		. 36	
Nov. 16-29		l	
	7		Epidemic.
do	2		In vicinity.
Oct. 9-Nov. 13	5	2	
Nov. 26-Dec. 9	10	7	
Nov. 27-Dec. 10	10	2	
TYPHUS	FEVE	R.	
i	83		
l l	a	в	
	-		
Oct. 1-21	4	2	
Oct. 1-31		7	
Nov. 20-26 Dec. 18-24	24	1	
Oct. 1-31	14		
1		9	•
		-	
20000			
Oct. 30-Nov. 5 Nov. 5	1		Outbreak.
YELLOW	FEVE	R.	
Nov. 1-30 Dec. 19	1	1	Present 50 miles northward o
	Nov. 13-26 Oct. 1-Nov. 5 Oct. 9-Nov. 5 Oct. 1-31 do. Oct. 23-Nov. 26 Nov. 1-30 Nov. 16-29 Nov. 6-19 Dec. 10 do. Oct. 9-Nov. 13 Nov. 26-Dec. 9 Nov. 27-Dec. 10 TYPHUS Nov. 1-30 Aug. 1-Oct. 31 Oct. 23-Nov. 23 Nov. 7-27 Nov. 19-25 Oct. 1-21 Oct. 1-31 Nov. 20-26 Dec. 18-24 Oct. 1-31 do Oct. 2-Nov. 23 Nov. 20-Dec. 10 Oct. 30-Nov. 5 Nov. 5 YELLOW	Nov. 13-26. 12 Oct. 1-Nov. 5 5 Oct. 9-Nov. 5 20 Oct. 1-31 20do 87 Oct. 2-Nov. 26 16 Oct. 23-Nov. 5 1 Nov. 1-30 7 Dec. 10 20 Oct. 9-Nov. 13 5 Nov. 26-Dec. 9 10 Nov. 27-Dec. 10 10 TYPHUS FEVE Nov. 1-30 1 4 Aug. 1-Oct. 31 83 Oct. 23-Nov. 25 20 Oct. 1-21 4 Oct. 1-31 14 Oct. 1-31 14 Oct. 1-31 14 Oct. 1-31 14 Oct. 1-31 14 Oct. 1-31 14 Oct. 1-31 14 Oct. 1-31 14 Oct. 1-31 14 Oct. 30-Nov. 25 3 Nov. 20-Dec. 10 10 Oct. 30-Nov. 5 1 Nov. 5 11 Nov. 5 11 YELLOW FEVEL Nov. 1-30 1 1	Nov. 13-26