# SOIL INOCULATION WITH AZOTOBACTER

BY PAUL EMERSON

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A DISSERTATION SUBMITTED TO THE GRADUATE FACULTY OF THE IOWA STATE COLLEGE OF AGRICULTURE AND ME-CHANIC ARTS IN PARTIAL FULFILLMENT OF THE REQUIREMENTS FOR THE DEGREE OF DOCTOR OF PHILOSOPHY

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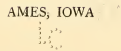
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# Soil Inoculation with Azotobacter

By PAUL EMERSON

AGRICULTURAL EXPERIMENT STATION IOWA STATE COLLEGE OF AGRICULTURE AND MECHANIC ARTS

> AGRONOMY SECTION Soil Bacteriology



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# Soil Inoculation With Azotobacter\*

# BY PAUL EMERSON.

Following the discovery of the nitrogen fixing powers of the symbiotic bacteria in the soil, early investigators found that the power of utilizing the free atmospheric nitrogen was not confined to the symbiotic bacteria alone. They noted increases in soils which had borne no legumes and they found that fallow soils in particular increased appreciably in nitrogen content. These facts stimulated researches which led to the discovery of many forms of bacteria which are able, when growing alone, to fix nitrogen from the air. The chief of these is now known as the azotobacter group.

It seems likely that the azotobacter will prove more effective in fixing nitrogen than the symbiotic bacteria, although the general requirements of the two classes of organisms are very similar. The azotobacter are active in practically all soils regardless of the kind of crop grown when conditions for their growth are satisfactory. These conditions are probably much the same as for the symbiotic bacteria except that these latter organisms require the presence of a specific legume for fixing the greatest amount of nitrogen. Azotobacter require a certain amount of carbonaceous material in the soils and are usually stimulated by. a small amount of nitrogen, but the exact optimum conditions for their growth are as yet unknown. These organisms are active in causing nitrogen increases in many soils, but the feasibility of introducing them into the soil or of attempting to increase their nitrogen-fixing powers by artificial means, and the effect of the presence of growing plants on their efficiency are questions as yet unanswered, although Lipman has indicated that under proper conditions successful inoculation may be accomplished in soils and Bottomley has successfully grown pure cultures of these organisms in the presence of growing plants with favorable results.

# HISTORICAL

Beijerinek (2) isolated and described the first azotobaeter (in 1901). He found two species, one of which he named Azotobacter chroöcoccum and the other Azotobacter agilis. The former was isolated from the soil and the latter from a sample of water taken from one of the canals in the city of Delft. Two years later Lipman (36) added a third species, A. vinelandii, to the hist and the following year isolated and described two more, giving them the names of A. beijerinckii and A. woodstownii. Of the five organisms of this species, A. chroöcoccum, A. beijerinckii

\*Thesis submitted in partial fulfillment of the requirements for the Degree of Doctor of Philosophy at the Iowa State College.

and A. vinelandii are considered the most important in soil inoculation studies.

The frequency with which investigators in all parts of Europe and America have isolated azotobacter from various soils, indicates that they are widely distributed. Christensen (10) found that they were present throughout northwestern Europe, the activity of the organism apparently depending on the basicity of the soil. This view was later supported by the works of Fisher (14), Löhnis and Pillai (45) and others. Ashby (1) studied the soils of Mombasa, East Africa, Cairo, Egypt and Rothamsted, England and found azotobacter forms present in most cases. Lipman and Burgess (42) working with forty-six samples of soil from various parts of the world, found that over one-third of them contained azotobacter, the predominant form being A. chroöcoccum. Many of the soils examined were muscum specimens and had been kept in tightly stoppered bottles for long periods of time.

#### DESCRIPTION OF AZOTOBACTER.

Beijerinck characterizes the azotobacter as stout bacteria, 4-6 microns or less in length, sometimes longer, occurring as large diplococci or short rods in young cultures, the hyaline cells often containing a vacuole and the entire organism enclosed in a mucilagenous wall of varying thickness. They have a single polar flagellum or bundles of 4-10 polar flagella of about the same length as the organism itself. Beijerinck found no spores. Vagler (65) writes that the older colonies produce involution forms similar to those of yeasts while Heinz (22) and Fisher (15) showed that the organisms can resist drying for six to nine months. Later investigations by Mulvania (50) and Löhnis and Smith (47) demonstrated that the organism produces spores and completes a very complicated life cycle. Descriptions of azotobacter and detailed cultural characteristics of the organism were given by Lipman (35), Beijerinck (2), Prazmowski (54), Warmbold (70), Bonazzi (6), Löhnis and Westerman (48), Löhnis and Hanzawa (44), Jones (27) and others.

#### ACTIVITIES OF AZOTOBACTER.

Beijerinck first claimed that the isolated pure cultures of azotobacter were able to fix the atmospheric nitrogen in appreciable amounts; later, however, when working with Van Delden (4), he retracted this statement, claiming that pure cultures did not have this ability and that only in the presence of very small celled organisms called radiobacter could the free nitrogen of the air be fixed in the soil. Gerlach and Vogel (18), Heinz (23), Lipman (37) and Freudenreich (17) proved conclusively that the earlier conclusions of Beijerinck were correct and that the organism may fix considerable amounts of nitrogen in pure cultures. Lipman accounts for the fact that Beijerinek did not get a fixation of nitrogen in pure cultures by showing that the organism will not fix nitrogen unless the reaction of the medium is made neutral or slightly alkaline. When Beijerinek later accepted this suggestion he found that his pure cultures were able to fix atmospheric nitrogen.

#### STUDIES OF AZOTOBACTER.

Very few investigators have attempted to inoculate soils with azotobacter or other non-symbiotic nitrogen fixing bacteria under conditions approximating those in the field. The influence of various kinds of sugars, cellulose, inorganic salts, and various organic compounds on the nitrogen-fixing power of the organisms have been studied extensively. Gerlach and Vogel (19), Pringsheim (55), Krainsky (33), Koch (30), Hoffman and Hammer (25) and Stranak (61) have found that various sugars and cellulose materially increase their nitrogen fixing powers while Fisher (16), Christensen (10), Löhnis and Pillai (46), Wilfarth and Wimmer (59) Kaserer (28), Rosing (59), Vogel (66), Greaves and Anderson (20) and Pringsheim (56) have shown that small amounts of lime, very small amounts of nitrogen, various inorganic salts and even a very small amount of arsenic will stimulate the nitrogen fixing power of the organisms in the presence of certain carbon compounds. Stoklasa (60) studied the products of the activities of the azotobacter organisms, confining his researches largely to the amounts and kinds of gases produced under different circumstances, under the influence of various substances supposed to be energy sources, and under varying temperature conditions. His results have been more or less confirmed by the works of Thiele (64), Hoffman (24), Kellerman and Smith (29) and Ehrenberg (13).

The activity of the azotobacter in soils in general, and particularly under laboratory conditions, was fully shown by the works of Lipman (39), Voorhees and Lipman (68), Löhnis (43), Kuhn (34), Freudenreich (17), Dvorak (12), Remy (57), Remy and Rosing (58), Jacobitz (26), Stranak (62), Headden (21), Peterson and Mohr (52), Koch and Seydel (31), Omeliansky and Ssewerowa (51), Warmbold (71) and others who demonstrated that under various conditions and in almost every type of known soil these organisms are able to fix appreciable amounts of the free atmospheric nitrogen. Only a few of these investigators, however, have made any attempt to secure an active flora of these organisms in the soil. Vogel (67) inoculated pure cultures of azotobacter into soils that had been treated with grape sugar, in some cases adding comparatively large amounts of nitrate of soda. In pot experiments with oats and mustard, increases were noted for the inoculated series, altho the pots receiving nitrate of soda gave the greatest yields. When the experiment was repeated in the field the inoculated plots gave smaller yields than the uninoculated, and the inoculation appeared to have an injurious effect upon the erop.

A short time later Lipman and Brown (41) tried inoculation experiments with A. vinelandii and A. beijerinckii. They sunk four foot cylinders open at both ends into the soil, filled the cylinders with soil and inoculated the soil with the organisms. The first summer the soils were left bare and then a rotation of crops was followed and oats, corn and rye grown in succession. While considerable variation was found in the nitrogen content of the crops and in the dry weight, the general conclusion reached was that the activities of the organisms did not increase the nitrogen content of the soil. The results do not preclude the possibility that inoculation with the organisms in question may be made of practical value, provided proper conditions for the best growth of the organisms are secured. Bottomley (7) and Bottomley and Hall (9) experimented with oats, barley and some root crops, and arrived at the same conclusions as did Lipman and Brown. Stranak (63) also inoculated soils with azotobacter and found a pronounced increase in the growth of potatoes, grain and beets.

Altho the experiments dealing with the inoculation of soils with azotobacter have been inconclusive, it is believed that under proper conditions such inoculation may be extremely profitable.

# EXPERIMENTAL

The wide distribution of non-symbiotic nitrogen fixing bacteria in many types of soils is practically parallel with the distribution of the symbiotic organisms, and since it is practical and profitable to inoculate soil with the latter, even tho the particular organism may be present, the following questions quite naturally arise:

1. If the azotobacter are not present in the soil, can inoculation be profitably accomplished?

2. What soil conditions are necessary for the greatest fixation of nitrogen by these organisms?

These questions have an important bearing on the problem of the maintenance of permanent fertility in soils from the nitrogen standpoint and may govern the choice of the proper method of farming. Some commercial concerns have placed cultures on the market, claiming that they contain sufficient numbers of the nonsymbiotic nitrogen fixing bacteria to enable the farmer to solve his nitrogen problem without growing legumes. However, results of experiments showing that such cultures are capable of inoculating the soil were not found in the present investigation.

# INCREASING THE NITROGEN FIXING POWER OF PURE CULTURES.

Very little work has been done along the line of breeding pure cultures of bacteria to an increased efficiency in their specific actions, in fact, practically all the experiments have been carried out with the idea of finding a method whereby the organism could be kept alive for long periods without periodic transfers. The earliest investigation along this line was that of Czaplewski (11) who limited the amount of air in the tube by saturating the plug with paraffine. Later Lunt (49) found that certain cultures of water bacteria may be kept alive much longer in sterile water than in ordinary culture media. In some cases he kept certain organisms alive for two years by this method. Bolley (5) secured good growths of B. amylovorus and Bact, dianthi in agar and in bouillon by making transfers from cultures that had been hermetically sealed for nine years. It is not stated whether or not the organisms were tested for their pathogenicity and hence their virulence is left in doubt. This work supports that of Czaplewski in showing that cultures can be kept alive for long periods of time if the transpiration is reduced to a minimum. Some commercial concerns claim that they are able to increase the efficiency of their particular cultures of legume bacteria by alternate inoculations first on agar, then into sterile greenhouse soil, growing the specific legume to which the organism in question is adapted, and re-isolating the organism from the nodules produced on the roots of the legume. If this is possible for the symbiotic bacteria then it seems probable in the case of the nonsymbiotic organisms. The following questions naturally suggest themselves:

1. Can the nitrogen fixing power of azotobacter be increased by periodic transfers on nitrogen free media?

2. Can the nitrogen fixing powers of azotobacter be increased by growing the organism in the presence of growing plants?

In outlining work to answer the above questions it was realized that a large number of bacteria should be used. A number of large celled nitrogen fixing organisms that had all the staining reactions of the azotobacter type and closely resembled it in size and shape, were isolated in pure cultures from soil secured from the humus plots at the Iowa station and were designated with laboratory numbers. At the same time pure cultures were secured and their activities determined along with these of the unnamed cultures. The pure cultures were kindly furnished by Dr. J. G. Lipman of the New Jersey Agricultural Experiment Station and also by the American Museum of Natural History of New York,

#### MEDIA USED.

The nitrogen free medium used throut the experimental work was a modification of that proposed and used by Lipman (35), and its composition was as follows:

Distilled water	1,000 cc
Di-potassium phosphate	0.2  grams
Magnesium sulphate	0.2 grams
Calcium chloride	0.02 grams
Dextrose	10.0 grams
10% Ferric chloride solution	2 drops

The solution was brought to boiling and made neutral to phenolphthalein by the addition of N/10 NaOH. If a solid medium was desired 1% powdered agar was added. Sterilization was accomplished by placing in the autoclave at ten pounds for 20 minutes.

Inoculation was seeured by scraping off a two days' growth from the agar slants with a sterile needle and transferring it to flasks containing 50 ee. of the above solution. In order to determine whether the nitrogen fixing power of the organisms was stimulated by the addition of nitrogen, the above solution with the addition of 1 mg. of nitrogen as sodium nitrate was used.

#### PRELIMINARY STUDIES.

All of the organisms of the azotobaeter type including both the pure cultures and the unnamed cultures, were inoculated into 50 c. c. of both of the above solutions and tested for their nitrogen fixing powers. The inoculated solutions were incubated for three weeks at room temperature  $(22-25^{\circ} \text{ C})$  and then Kjeldahlized. The amount of nitrogen fixed by each organism in the different solutions is shown in table I. The same cultures were transferred 12 times at three to four day intervals on nitrogen free media and their nitrogen fixing power tested in solutions with and without nitrogen. The results appear in table II.

The laboratory organisms used in table I had been freshly isolated and purified from the soil, the named cultures had been kept on agar slants for varying periods of time. During the time that the inoculated culture solutions were incubating the transfers were being made in preparation for the inoculations for table II.

Comparing the two tables we find that a majority of the organisms decreased in their ability to fix atmospheric nitrogen, altho a few showed a slight increase or at least retained their efficiency. From these the following eight were selected for further study: No. 4, No. 22, No. 26, No 27, A. vinelandii, A. chroococcum, A. beijerinckii and A. chrooccoccum (HCM). These eight organisms were studied under both laboratory and greenhouse conditions.

# TABLE I-NITROGEN-FIXATION BY PURE CULTURES IN SOLUTION WITH AND WITHOUT NITROGEN.

#### TABLE II-NITROGEN-FIXATION BY PURE CULTURES IN SOLUTION WITH AND WITHOUT NITROGEN.

After each organism has been transferred twelve times on nitrogen-free media at three to four day intervals.

	N. Fixed	l in Mgs.
Organism	Solution wiithout Nitrogen	Solution with Nitrogen
	2 Print	Lo Put
	lifi	Solu with Nitr
	N RY	0 \$4
		1
Lab. No. 1	2.24	0.14
Lab. No. 2	2.38	1.54
Lab. No. 3	0.28	
Lab. No. 4	0.98	1.96
Lab. No. 6	1.96	1.82
Lab. No. 7	3.22	1.68
Lab. No. 9	0.42	2.10
Lab. No. 10	0.42	0.84
Lab. No. 11	0.42	0.70
Lab. No. 12	lost	1.40
Lab. No. 14	1.12	lost
Lab. No. 15	2.80	1.12
Lab. No. 16	7.14	lost
Lab. No. 18	0.28	0.42
Lab. No. 19	1.12	1.54
Lab. No. 20	0.56	1.96
Lab. No. 21	0.56	2.10
Lab. No. 22	0.42	2.52
Lab. No. 23	0.70	2.52
Lab. No. 24	0.70	1.82
Lab. No. 25	0.28	1.82
Lab. No. 26	0.56	2.10
Lab. No. 27	1.12	5.60
A. vinelandii	lost	2.66
A. chroococcum (HCM)	4.20	3.08
A. chroococcum	0.84	1.54
A. chroococcum (Colo)_	0.70	2.52
A. beijerinckii	0:84	1.68
A. beijerinckii No. 5	1.96	2.38
And a second strength of the second strength		

$ \begin{array}{ c c c c c c c c c c c c c c c c c c c$			in Mgs
Lab. No.       1	Organism	Solution without Nitrogen	Solution with Nitrogen
A. vinelandii         0.00         0.00           A. ehroococcum         1.12         1.82           A. ehroococcum (HICM)         2.52         1.12           A. ehroococcum (Colo)         0.00         1.26           A. beijerinckij         0.42         2.52	Lab. No.       2         Lab. No.       3         Lab. No.       4         Lab. No.       6         Lab. No.       7         Lab. No.       7         Lab. No.       9         Lab. No.       10         Lab. No.       11         Lab. No.       12         Lab. No.       15         Lab. No.       16         Lab. No.       18         Lab. No.       18         Lab. No.       19         Lab. No.       21         Lab. No.       22         Lab. No.       22         Lab. No.       23         Lab. No.       24         Lab. No.       25         Lab. No.       26	$\begin{array}{c} 0.84\\ 0.14\\ 0.14\\ 1.82\\ 0.00\\ 0.28\\ 0.00\\ 0.98\\ 0.00\\ 0.98\\ 0.00\\ 0.42\\ 0.00\\ 0.00\\ 0.00\\ 0.00\\ 0.00\\ 0.00\\ 2.10\\ 0.00\\ 0.00\\ 1.12\end{array}$	$\begin{array}{c} 1.12\\ 0.00\\ 0.98\\ 0.84\\ 0.28\\ 1.26\\ 0.00\\ 1.68\\ 0.28\\ 1.12\\ 1.40\\ 0.14\\ 1.54\\ 1.54\\ 1.54\\ 1.54\\ 1.68\\ 0.28\\ 1.52\\ 1.54\\ 1.68\\ 0.28\\ 1.82\\ \end{array}$
A. beijerinckii No. 5 1.12 0.00	A. vinelandii A. chroococcum A. chroococcum (HCM) A. chroococcum (Colo)- A. beijerinckii	$\begin{array}{c} 0.00 \\ 1.12 \\ 2.52 \\ 0.00 \\ 0.42 \end{array}$	$\begin{array}{c} 0.00 \\ 1.82 \\ 1.12 \\ 1.26 \\ 2.52 \end{array}$

# LABORATORY STUDIES.

The laboratory studies were arranged in a series of three experiments as follows:

1. To determine the effect of transfers made every other day on the nitrogen fixing power of the organisms.

2. To determine the effect of transfers made once each week in sand eultures variously modified.

3. To determine the effect of growing four of the organisms on both agar and sand in large flasks with and without the presence of growing plants.

# Series 1. To Determine the Effect of Transfers Made Every Other Day on the Nitrogen Fixing Power of the Organisms.

Using the eight selected organisms transfers were made every other day on the nitrogen free medium for a period of three weeks. It was feared that such rapid transferring for so long a period on a medium practically free from nitrogen would reduce the vitality of the organisms, accordingly each fifth transfer was made on a modification of the medium consisting in the addition of one milligram of nitrogen as sodium nitrate to each liter of the regular dextrose agar. At the end of the transfer period the organisms were inoculated into the nitrogen free and nitrogen containing solutions incubated for the same periods of time and the amount of nitrogen fixed determined by Kjeldahlizing. The results of the determinations are shown in table III.

TABLE III—THE EFFECT OF TRANSFERS MADE EVERY OTHER DAY FOR FOUR WEEKS ON THE NITROGEN FIXING POWER OF THE ORGANISMS.

	Nitrogen Fixed in Mgs.							
Organism		ion with Nitrogen	out	Solution with Nitrogen				
Lab. No. 4 Lab. No. 22 Lab. No. 26 Lab. No. 26 Lab. No. 27	(a) 0.14 0.00 0.14 0.28	(b) 0.42 0.00 0.14 3.50	(Av.) 0.28 0.00 0.14 1.98	(a) 0.70 0.14 0.28 0.70	(b) 0.42 0.56 0.98 0.98	$ \begin{array}{c c} (Av^{\cdot}) \\ 0.56 \\ 0.35 \\ 0.63 \\ 0.84 \end{array} $		
A. vinelandii A. chroococcum A. chroococcum (HCM) A. beijerinckii	$     \begin{array}{r}       0.28 \\       0.56 \\       2.66 \\       0.84     \end{array} $	0.42 0.28 lost 2.66	$     \begin{array}{r}       0.35 \\       0.42 \\       2.66 \\       1.75     \end{array} $	$     \begin{array}{r}       0.98 \\       0.98 \\       1.40 \\       0.28     \end{array} $	$0.98 \\ 1.12 \\ 1.40 \\ 0.28$	$     \begin{array}{r}       0.98 \\       1.05 \\       1.40 \\       0.28     \end{array} $		

That these transfers should have been made at longer intervals is evidenced by the fact that tables I and II showed that 12 of the eultures had increased in efficiency after they had been transferred every three days for 36 days. However, during the latter work the organisms did not show any indications of a loss of vitality and the growth at all times was vigorous and rapid. Table III shows a decrease in the nitrogen fixing powers of all the organisms except in the case of A. chroococcum (HCM) which appears to have retained its efficiency thruout the experiment.

# Series 2. To Determine the Effect on the Nitrogen Fixing Power of Transfers Made Each Week in Sand Cultures.

In the following experiment sand was used instead of agar as the basis for the medium. Ground oats straw, ground red clover hay and either the regular dextrose solution, or the dextrose solution containing nitrogen were added. The tests were carried out in tubes arranged as follows: 6.25 gr. sand+2.5 cc N. free dextrose solution.
6.25 gr. sand+2.5 cc dextrose solution containing 0.2 gr. NaN0<sub>3</sub> per liter.
6.25 m sand+2.5 cc N. free destrose solution (0.1 m s)

6.25 gr. sand+3.5 cc N. free dextrose solution+0.1 gr. clover nay. 6.25 gr. sand+3.5 cc N. free dextrose solution+0.1 gr. oats straw.

5.25 gl. salu+5.5 cc N. free dextrose solution+0.1 gr. oats straw

The organisms were transferred directly from the slants into the tubes and there allowed to incubate at room temperature for seven days. A small portion of the sand was then transferred to a fresh tube of the same medium as the original. As this particular experiment did not directly follow the others the efficiency of the organisms was tested before they were inoculated into the sand. Table IV shows the amount of nitrogen fixed by the pure

TABLE IV-THE NITROGEN FIXING POWER OF THE PURE CULTURES IMMEDIATELY BEFORE THE SAND CULTURE EXPERIMENTS.

Organism	[Solution witnout Nitrogen			Solution with Nitrogen		
	(a)	(b)	(Av.)	(a)	(b)	Av.
Lab. No. 4	0.14	0.70	0.42	0.28	1.54	0.91
Lab. No. 22	0.00	0.00	0.00	3.36	3.08	3.22
Lab. No. 26	0.00	0.14	0.07	2.94	2.66	2.80
Lab. No. 27	0.00	0.00	0.00	2.38	2.52	2.45
A. vinelandii	0.00	0.00	0.00	1.40	1.40	1.40
A. chroococcum	0.00	0.14	0.07	2.80	2.66	2.73
A. chroococcum (HCM)	0.00	0.00	0.00	2.52	2.38	2.45
A. beijerinckii	0.98	0.84	0.91	2.94	lost	2.94

eultures at the beginning of this series of incubation, and the same methods as in the previous experiments.

At the end of the fourth transfer period, i. e., four weeks after the start of the experimental series, the organisms were inoculated into dextrose solution and their nitrogen fixing powers determined. After four more weeks of transferring or in all eight weeks the final inoculation into dextrose solution was made. The influence of the oats and elover in the presence of sand on the nitrogen fixing power of the organisms used is shown in tables V and VI, by the fact that both the large celled organisms of the

 TABLE V.—NTTROGEN FIXED BY THE PURE CULTURES AFTER FOUR

 TRANSFERS IN SAND AT PERIODS OF SEVEN DAYS EACH.

	Nitrogen Fixed in Mgs.						
Organism	dex. sol.	${ m dex. \ sol.} + { m N}$	dex. sol.+ oats straw				
Lab. No. 4           Lab. No. 22           Lab. No. 26           Lab. No. 27           A. vinelandii           A. chroococcum           A. chroine (HCM)           A. beiterinekii	$\begin{array}{c} 0.28 \\ 0.07 \\ 0.77 \\ 0.42 \\ 0.14 \\ 0.21 \\ 0.07 \\ 0.14 \end{array}$	$\begin{array}{c} 0.35\\ 0.28\\ 0.07\\ 0.35\\ 0.21\\ 0.21\\ 0.14\\ 0.28\end{array}$	$\begin{array}{c} 0.28 \\ 0.00 \\ 0.14 \\ 1.27 \\ 0.42 \\ 0.07 \\ 1.19 \\ 0.35 \end{array}$	$\begin{array}{c} 0.98\\ 0.35\\ 0.00\\ 0.42\\ 0.28\\ 0.21\\ 0.42\\ 0.28\end{array}$			

	Nitrogen Fixed in Mgs.						
Organism	dex. sol.	$\stackrel{\rm dex. \ sol.}{+\rm N}$	dex. sol.+ oats straw	dex. sol.+ clover hay			
Lab. No. 4	0.30	0.20	0.30	0.40			
Lab. No. 22	0.20	0.20	0.20	2.00			
Lab. No. 26	0.70	0.40	0.00	1.00			
Lab. No. 27	0.20	0.10	0.40	0.50			
A. vinelandii	0.20	0.40	0.40	0.50			
A. chroococcum	0.30	0.10	0.20	lost			
A. chroococcum (HCM)	0.10	0.40	0.40	1.40			
A. beijerinckii	0.00	0.00	0.20	0.50			

TABLE VI-NITROGEN FIXED BY THE PURE CULTURES AFTER EIGHT TRANSFERS AT PERIODS OF SEVEN DAYS EACH.

azotobacter type and the azotobacter themselves, made gains in their nitrogen fixing powers. There was no distinct gain due to any one kind of carbonaceous material. Of the six organisms showing gains A. chroococcum made the most notable, especially in the presence of the oats straw. The nitrogen fixing power of No. 4 appears to be rather constant thruout the series, with no appreciable gain or loss. A. beijerinckii showed a decided loss in its power to fix nitrogen in each of the four media, but gave a slight indication that in the presence of the clover hay it might be slowly regaining its power.

# Series 3. To Determine the Effect of Growing the Organisms on Both Agar and Sand With and Without the Presence of Growing Plants.

The main points considered in this experiment were: An increase in the surface area over which the organism could grow; an increase in the time between transfers and the growing of the organisms in the presence of an undetermined species of algae and with growing oats and red clover plants. Two liter Erlenmeyer flasks were used and arranged in the following manner conforming to the outlines of the experiment:

Flask No. 1. 1000 cc N. free dex. agar+1 gr.  $CaCO_3$  planted to oats. Flask No. 2. 1000 cc N. free dex. agar+1 gr.  $CaCO_3$  planted to red clover.

Flask No. 3. 1000 cc N. free dex. agar+1 gr. CaCO<sub>3</sub> planted with an undetermined species of algæ.

Flask No. 4. 1000 gr. pure quartz sand+180 cc N. free dex. solution neutralized with CaCO<sub>3</sub>, planted with oats.

Flask No. 5. 1000 gr. pure quartz sand +180 cc N. free dex. solution neutralized with CaCO<sub>3</sub>, planted with red clover.

Flask No. 6. 1000 gr. pure quartz sand+180 cc solution without dex. neutralized with CaCO<sub>3</sub>, planted with an undetermined species of algæ.

Check flasks of sand and dextrose agar.

The flasks of agar were sterilized in the autoclave at ten pounds for 30 minutes, but the flasks of sand were sterilized at 15 pounds pressure for four hours once a day for three consecutive days. Bacteriological tests on the sand at the end of that time showed it to be sterile.

The culture of the algae used was so closely associated with a bacterial growth that a separation would have required a long time. For that reason it was not purified but was grown in sterile distilled water, for about three months before inoculation. The inoculation of the algae was made in the flasks about two weeks before the inoculation with the azotobacter cultures in order that the algae might make a sufficient growth to supply the bacterial cultures with the proper amount of carbonaceous material. -Toprevent contamination by the oat and red clover plants, the seeds were soaked three minutes in a 1-500 solution of mercuric chloride, washed in sterile distilled water three times and then planted in sterile agar plates. By this means the seeds were sprouted and those which were contaminated were discovered and rejected. The sprouted seeds were transferred from the plates to the flasks by means of the platinum needle. A block of the agar containing the sprouted seed was cut out and placed in the proper position on the medium in the flask. The flasks were then carefully observed for five days to insure the absence of contamination.

As all the flasks contained growing plants no attempt was made to exclude the light, but neither were they placed in the direct sunlight. They were kept on a table about eight feet from a large window facing the west. All the flasks were plugged with non-absorbent cotton and after inoculation a cap of paraffined paper was placed over the mouth and held in place with a rubber band. While the plants did not develop rapidly the oats grew much faster than the clover for about three weeks, after which time both began to lose chlorophyl and by the end of the five weeks' experimental period, the majority of the plants had died. The oats and clover in the flasks inoculated with A. chroococcum (IICM) and the clover in the flasks inoculated with B. radicicola showed a slight gain in growth and altho far from vigorous at the end of the experiment were still alive and growing slowly.

# ORGANISMS USED.

The organisms used were A. chroococcum (HCM), A. vinelandii, A. beijerinckii and for the purpose of comparison, B. radicicola isolated from the nodules of sweet clover. The latter were isolated and purified especially for this series, and introduced to compare the effects of symbiotic and non-symbiotic organisms on the growth of the plant used. The results secured with it, however, were of no great significance. After the bacteria had remained undisturbed in the flasks for five weeks, transfers were made directly from the flasks into 50 ec. of the nitrogen-free dextrose solution, incubated for three weeks, and the nitrogen fixed determined in the usual manner. The total amount of nitrogen fixed by the bacteria themselves, as well as the amount fixed by the bacteria but due to the stimulative action of the plants on the bacterial activities, is shown in table VII. There was a stimulation of the nitrogen fixing power of the organisms due to the presence of a growing plant, especially noticeable in the case of *A. vinelandii* and *A. chroococcum* (HCM) and to some extent in the case of *A. beijcrinckii*. *A. vinelandii* was stimulated thruout the entire series except when grown in sand in the presence of the algae. The oats and algae showed no

TABLE	VII-THE	EFFECT O	F GRO	WING	PLANTS	ON	THE	NITROGEN	FIXING
		POW	ER OF	PURE	CULTU	RES.			

			[	1					
					Nit	rogen	Fixed	in M <sub>i</sub>	gs.
Inoculum	Medium Plant Used (		(a)	(b)	(Av)	N. in algae check	Total amount due to bact. action.	Nitrogen in sand or agar check.	Total amount due to stim- ulative action of plants.
								1	
Algae	agar	check	0.84		0.84				
Algae Vinelandii	sand	check	1.40	0.98	1.19				
Vinelandii	agar sand	check	$1.12 \\ 1.40$	$0.98 \\ 1.40$	$1.05 \\ 1.40$				
Vinelandii	agar	oats	3.66	3.52	3.59		3.50	1.05	2.54
Vinelandii		red clover	2.10	2.38	2.24		2.24	1.05	1.19
Vinelandii		algae	4.20		4.20	0.84		1.05	2.31
Vinelandii		oats	4.20	4.06	4.13	0.01	4.13	1.40	2.73
Vinelandii		red clover	4.48	4.06	4.27		4.27	1.40	2.87
Vinelandii	and	algae	2.52	2.80	2.66	1.19		1.40	0.07
Chroococcum (HCM)	agar	check	0.00	0.00	0.00				
	sand	check	0.28	0.14	0.21				•
	agar	oats	0.28	0.44	0.36		0.36		0.35
	agar	red clover	1.82	3.22	2.52		2.52	0.00	2.52
Chroococcum (HCM)		algae	3.92	3.50	3.71	0.84		0.00	2.87
	sand	oats	0.28	0.56	0.42		0.42	0.21	0.21
Chroococcum (HCM) Chroococcum (HCM)	sand	red clover	0.28 3.22	lost	$0.28 \\ 4.20$	1 10	$0.28 \\ 3.01$	$0.21 \\ 0.21$	0.07 2.80
Beijerinckii		algae	0.00	$5.18 \\ 0.00$	4.20	1.19	5.01	0.21	2.00
Beijerinckii		check	0.00		$0.00 \\ 0.35$				
Beijerinckii		oats	0.14	0.00	0.07		0.07	0.00	0.07
Beijerinckii		red clover	1.40		1.40		1.40	0.00	1.40
Beijerinckii		algae	0.00		0.00	0.84			
Beijernnckii		oats	1.39	1.39	1.39		1.39	0.35	1.04
Beijerinckii	sand	red clover	1.39	1.39	1.39		1.39	0.35	1.04
Beijerinckii		algae	1.39		1.40	1.19		0.35	
B. rad., S. elo		oats	0.14	0.14	.014		0.14		
B. rad., S. clo.	agar	red clover	0.14		.014		0.14		
	agar	algae			0.56	0.84			
B. rad., S. clo B. rad., S. clo	. sand	oats	$  0.42 \\ 0.14$		$0.21 \\ 0.07$		$0.21 \\ 0.07$		
B. rad., S. clo.			$0.14 \\ 0.42$			1.19			1
1. Iau, D. Co.		uigut	· U•#4	0.00	0.40	1.10			

difference when grown on the agar and in the sand medium the greatest stimulation was produced by the red clover. The activities of A. chroococcum were stimulated to the greatest extent by the presence of algae in both sand and agar, the oats gave a poor stimulation in both cases, and red clover gave good results in the agar but not in the sand.

The nitrogen fixing power of A. beijerinckii was retarded by the presence of algae, but was stimulated by red clover in both the agar and sand. The oats stimulated this organism only when grown on the agar. The nitrogen fixing power of B. radicicola was so low thruout the experiment that the results are not considered.

# CONCLUSIONS FROM LABORATORY STUDIES.

1. Transfers made on a nitrogen free dextrose agar more often than once each week were detrimental to the nitrogen fixing power of azotobacter and other large celled nitrogen fixing organisms of the same type.

2. Transfers made once each week into a pure sand medium containing some carbonaceous material were beneficial to the nitrogen fixing power of the azotobacter in general, but the effect on *A. beijerinckii* was detrimental.

3. The nitrogen fixing power of A. vinelandii was stimulated to a marked extent when grown in large flasks for five weeks in the presence of red clover and oats on both agar and sand. It was stimulated by the presence of algae when grown on agar but not when grown on sand.

The nitrogen fixing power of A. chroococcum was stimulated markedly when grown on agar for five weeks in the presence of growing oats and red clover, but to a less extent when grown with the same plants in sand. The greatest stimulation for this organism was produced by growing it in the presence of algae in either sand or agar for the same period of time.

5. The nitrogen fixing power of A. beijerinckii was stimulated by the presence of red clover when grown on either sand or agar, and by oats when grown in sand. Algae in either agar or sand appeared to have a depressing effect on the nitrogen fixing power of this organism.

## GREENHOUSE STUDIES.

At the conclusion of the first experiment the eight organisms used in the laboratory series 1, 2 and 3 were also inoculated into soils in pots in the greenhouse. Ground oats straw or ground clover hay was added to these soils and the nitrogen fixing efficiency of the organisms both in fallow soils and in the presence of growing oats plants determined. Three experiments were carried out in this test, as soon as the soil in which one crop had been grown was sampled, it was immediately reseeded and another erop grown. Striet account was kept of the amount of nitrogen added in the seed and in the organic matter. The dry weight of the crop and the N. content as well as the nitrogen content of the soil was determined at the end of each experiment.

The soil used thruout the experiment was of the type classified by the United States Bureau of Soils as Miami silt loam, and according to tests in the laboratory did not contain azotobacter or any similar organisms. A large amount of this soil was thoroly air dried, sieved and mixed. Ten pounds were placed in each of eighty glazed pots, seventy-two of which were given the following treatment: Half, or thirty-six pots received an application of 22.68 grams ground oats straw, and the other half received an equivalent amount of ground red clover hay. This application (22.68 grams) was equivalent to a five-ton application of this material per acre. The ground material was thoroly incorporated in the soil, which was packed firmly in the pots. The pots used were glazed on the inside and made tight so there was no loss by neaching, neither was there any drainage provided.

# METHODS OF INOCULATION.

The inoculum used was the dextrose solution described above. 1500 e e were placed in each of six 2 L. flasks, inoculated with the organism desired and incubated for seven days. Microscopie examinations were made at the end of the incubation period to insure vigorous growth and the purity of the culture. 150 ee of the solution was used as the inoculum for each pot. This was poured over the surface of the dry soil and washed into it by the addition of sufficient water to bring the moisture content up to the optimum, in this case 25%. The pots were then weighed, covered with a cloth, and allowed to remain undisturbed for three days, in order to permit the moisture to become thoroly distributed thruout the soil. The pots were then arranged in the following manner and seeded to oats.

Thirty grains of Early Champion oats were planted in each pot at each seeding. They were planted at five points. One in the eenter of the pot and the other four were arranged between the eenter and the edge at equal distances apart. Six seeds were planted at each place and when the plants appeared they were thinned out and but one plant left in each place. The discarded plants were allowed to remain and decay on the soil in the pot from which they were drawn.

The length of the growing period was determined by the appearance of the seed-bearing spike. This period varied slightly in each of the series, the first closed in sixty-three days, the second in sixty-nine days, and the third in seventy days after planting.

#### PLAN OF EXPERIMENT

Pot. No.		Treatment	Inoculation
1	Fallow	Oats straw	A. chroococcum (HCM)
2-4	Cropped	Oats straw	A. chroococcum (HOM)
5-7	Fallow	Clover hay	A. chroococcum (HCM)
6-8	Cropped	Clover hay	A. chroococcum (HCM)
9-11	Fallow	Oats straw	A. chroococcum
10-12	Cropped	Oats straw	A. chroococcum
13-15	Fallow	Clover hay	A. chroococcum
14-16	Cropped	Clover hay	A. chroococcum
17-19	Fallow	Oats straw	<ol> <li>beijerinckii</li> </ol>
18-20-	Cropped	Oats straw	A. beijerinckii
21-23	Fallow	Clover hay	A. beijerinckii
22-24	Cropped	Clover hay	A. beijerinckii
25-27	Fallow	Oats straw	A. vinelandii
26-28	Cropped	Oats straw	A, vinelandii
29-31	Fallow	Clover hay	A. vinelandii
30-32	Cropped	Clover hay	A. vinelandii
33-35	Fallow	Oats straw	26 D.
34-36	Cropped	Oats straw	26 D.
37-39	Fallow	Clover hay	26 D
38-40-	Cropped	Clover hay	26 D.
41-43	Fallow	Oats straw	27 D.
42-44	Cropped	Oats straw	27 D.
45-47	Fallow	Clover hay	27 D.
46-48	Cropped	Clover hay	27 D.
49-51	Fallow	Oats straw	22 D.
50-52	Cropped	Oats straw	22 D.
53-55	Fallow	Clover hay	22 D.
54-56	Cropped	Clover hay	22 D.
57-59	Fallow	Oats straw	4 D.
58-60	Cropped	Oats straw	4 D.
61-63	Fallow	Clover hay	4 D.
62-64	Cropped	Clover hay	4 D.
65-67	Fallow	Oats straw	Mixed culture
66-68	Cropped	Oats 'straw	Mixed culture
69-71	Fallow	Clover hay	Mixed culture
70-72	Cropped	Clover hay	Mixed culture
73-74	Fallow	Oats straw	Check
75-76	Cropped	Oats straw	Check
77-78	Fallow	Clover hay	Cheek
79-80	Cropped		Check

The pots were watered with tap water about every other day and were weighed weekly. The loss in weight was replaced with water in order to keep the moisture content at the optimum. The growth of the plants was carefully noted and recorded by means of photographs at different periods. The harvested plants were dried, weighed, and the total nitrogen content determined by the Kjeldahl method.

At the close of each series of experiments the soils were removed from the pots, placed on a sterile oil cloth, thoroly mixed, sampled and returned to the original pot. The sample taken at this time approximated 500 grams dry weight. The pots were seeded again as soon as possible and the experiment continued. During the short period between sampling and reseeding the moisture content was kept at the optimum. The preliminary analyses, showing the nitrogen content of the original air dried soil, and of the same soil mixed with the ground oats or clover are as follows:

22.68 grs. ground oats straw co	ntained0.1416 grs. N. av. 6 dets.
22.68 grs. clover hay contained.	
10 lbs. original soil + oats stray	v contained2.4910 grs. N. av. 6 dets.
10 lbs. original soil + clover ha	y contained2.7647 grs. N. av. 6 dets.

## ACTION OF DENITRIFYING BACTERIA.

Some of the plants were very much stunted in their growth and an experiment was conducted to determine whether this was due to action by the denitrifying organisms. Samples weighing about six or eight grams were drawn from near the center of each pot by means of a sterile corkborer and placed immediately in sterile tubes. Sterile water was added and a soil suspension made from which inoculations were made into Giltay's denitrifying solution. The solution was incubated three weeks and the amount of nitrate nitrogen as well as the total nitrogen determined, the first by the aluminum reduction method of Potter and Snyder, and the second by the official method. The aluminum reduction was carried out by aeration, thus leaving the original solution available for analysis for total nitrogen. The results given in table VIII show that the denitrifying organisms were not the limiting factor in the growth of plants. Only the five soils in pots Nos. 29, 45, 62, 64 and 66 show any great loss in nitrogen and some of the pots show an actual gain in total nitrogen content. This gain is particularly noticeable in the soils inoculated with A. beijerinckii, No. 26 and in the cheek pots.

Pot	Nitrate N. mgs.	N. mgs.	Total N. mgs.	Check	Amt. denitri- fied
1	$\begin{array}{c} 0.70\\ 1.05\\ 0.98\\ 0.91\\ 0.90\\ 0.56\\ 0.70\\ 0.79\\ 0.70\\ 0.42\\ 0.70\\ 0.84\end{array}$	Iost 6.02 5.60 4.20 8.12 7.84 8.14 7.07 7.07 8.96	7.07 6.51 5.10 8.68 8.54 8.93 7.77 7.49	8.23 8.23 8.23 8.23 8.23 8.23 8.23 8.23	1.16 1.72 3.13 *: : 0.46 0.74
13 14 15	0.86 0.65 0.56	6.16 7.56	7.02 8.21	8.23 8.23 8.23	1.21 0.02

TABLE VIII—THE ACTIVITIES OF THE DENITRIFYING BACTERIA IN THE SOILS THREE WEEKS AFTER THE START OF THE EXPERIMENT

					-
	Nitrate N.				Amt. denitri-
Pot	mgs.	N. mgs.	Total N. mgs.	Check	fied
					nea
		1			
	]	1			
16	1.71	6.16	7.87	8.23	0.36
17 18	$0.56 \\ 0.96$	$7.56 \\ 7.70$	8.12	8.23	0.11
19	0.42	8.61	8.66 9.03	$8.23 \\ 8.23$	
20	0.84	8.40	9.24	8.23	:
21	0.84	7.91	8.75	8.23	-
22	0.56	9.38	9.94	8.23	
23	0.70			8.23	
24	0.78	7.21	7.98	8.23	0.25
25	0.86	6.44	7.30	8.23	0.93
26	0.06 0.58	7.42 8.04	7.48	8.23	0.75
27	0.58	7.28	$\frac{8.62}{8.15}$	$8.23 \\ 8.23$	0.08
29	0.56	5.04	5.60	8.23	2.63
30		6.30	0.00	8.23	2.00
31	1.54	5.32	6.86	8.23	1.37
32	0.43	9.10	9.53	8.23	:
33	0.70			8.23	
34	0.56	8.54	9.10	8.23	:
35	0.98	7.56	8.54	8.23	
36	0.79 0.86	8.68 7.00	$9.47 \\ 7.86$	$8.23 \\ 8.23$	0.37
38	0.90	6.44	7.34	8.23	0.89
39	1.54	6.58	8.12	8.23	0.11
40	0.63			8.23	
41	0.63	7.50	8.19	8.23	0.04
42	0.56	4.90	5.46	8.23	2.77
43	0.70			8,23	
44	0.98	6.86	7.84	8.23	0.39
45	0.77 1.05	$2.80 \\ 6.30$	3.57 7.35	$\frac{8.23}{8.23}$	$4.66 \\ 0.88$
47	6.77	6.16	7.35	8.23	0.88
48	0.56	6.02	6.58	8.23	1.65
49,	0.94	7.28	8.32	8.23	:
50	0.91			8.23	
51	0.81	7.77	8.58	8.23	:
52 53	0.50	7.70	8.20	8.23	0.03
53	$0.77 \\ 0.49$	$8.12 \\ 7.56$	8,89 8,05	$8.23 \\ 8.23$	; 0.18
55	0.85	6.44	7.30	8.23	0.18
56	1.00	6.09	7.09	8.23	1.14
57	0.86	5.46	6.32	8.23	1.81
58	0.86	7.70	8.56	8.23	:
59	0.86	5.60	6.46	8.23	1.77
60	1.33			8.23	
61	0.91	1.00	2.74	8.23 8.23	5.49
62 63	0.78 1.26	$1.96 \\ 5.60$	6.86	8.23	5.49 1.37
64	1.31	4.76	6.07	8.23	2.16
65	1.31	7.28	8.54	8.23	:
66	0.93	1.54	2.52	8.23	5.71
67	0.14	7.28	7.42	8.23	0.81
68	0.70	6.16	6.86	8.23	1.37
69	0.56	6.72	7.28	8.23	0.95
70	0.84	4.62 8.40	5.46	$\frac{8.23}{8.23}$	2.77
71	0.49	8.40	8.89 8.65	8.23	:
72	0.87	6.02	6.89	8.23	1.34
74	1.05	7.70	8.75	8.23	:
75	0.65	9.80	10.43	8.23	:
76	0.51	8.12	8.63	8.23	:

TABLE VIII-CONTINUED.

\*No denitrification is shown by :

# PRELIMINARY TESTS FOR NITROGEN FIXATION.

To discover the action of the bacteria in the incculated soils samples were taken from the fallow pots four weeks after the start of the experiment and their total nitrogen content determined. Table IX shows a gain in the nitrogen content over the original soil and the check soils but the actual gain due to the action of the bacteria introduced was very slight. Organisms 22, 4 and the mixed cultures showed no gain whatever, and the others showed only a slight gain in those soils to which clover had been

TABLE 1X—THE ACTIVITY OF THE BACTERIA IN THE INOCULATED FALLOW SOILS AFTER FOUR WEEKS.

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$\begin{array}{c c c c c c c c c c c c c c c c c c c $		Gran	os N.		rogen t of	in.	cks at in id hay	e to etivi-
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	Pot	st	2nd	Av.	Orig. nit conten soil	Nitrogen crease soils	N. in che and thi oats an clover	Gain due bact. a ties
$\begin{array}{cccccccccccccccccccccccccccccccccccc$					-			
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	1							
$\begin{array}{cccccccccccccccccccccccccccccccccccc$								
$\begin{array}{c c c c c c c c c c c c c c c c c c c $							1.0025	
$\begin{array}{c c c c c c c c c c c c c c c c c c c $								0.1270
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	9							0.0000
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$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	13							
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	15							
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	17							0.0083
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$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	33							0.0083
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$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	37							
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	39							
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	41							0.0221
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	43							0.1000
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$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	59							
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	61							
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	63							
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$ \begin{array}{cccccccccccccccccccccccccccccccccccc$								
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	69							
$\begin{array}{cccccccccccccccccccccccccccccccccccc$		3.6069	3.4431	3.5250	2.7047	0.7603	1.0025	
$\begin{array}{cccccccccccccccccccccccccccccccccccc$			0.0000	0.0570	0.0101	0.9010		
$\begin{array}{cccccccccccccccccccccccccccccccccccc$								
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$								
Average four checks								
Average four checks	78	3.0643	3.2239	3.1441	2.3494	0.7947		
Average four checks0.08/2 + N in oats 0.1310 $= 0.128$					0.000	D I NI in	anta a l	160 7999
	Average four check	KS			0.087	2 + N m q + N m	olo 0.14	53-1 0025
					0.387	4 T IN III	0.0.41	00-1.0020

Grams of Nitrogen Per 10 Pounds of Soil.

added. This difference may have been due to variation in the rate of decomposition between the elover and the straw.

At the end of the three growing periods the soil in each pot was sampled and the total nitrogen content of both the soil and the entire crop determined. The amount of nitrogen found in the determinations and its relation to the total amount due to the bacterial activities is given in three separate tables, one for each growing period. From these complete tables three condensed tables have been made as follows: For the first growing period, table X, for the second growing period, table XI, for the third growing period, table XII, and a recapitulation table XIII.

TABLE X-THE NITROGEN FIXED B	<b>ACTERIA</b> —FIRST PERIOD
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(Condensed from Appendix Table I.)

	rt.
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	pa
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	
9-11.       F oats       A. chrooc. $3.232$ $3.2532$ $3.6232$ 10-12       Cloats       A. chrooc. $3.5380$ $3.2532$ $3.6232$ 10-12       Cloats       A. chrooc. $3.5380$ $3.5380$ $3.7300$ 13-15       F clover       A. chrooc. $4.7726$ $4.6183$ $3.8063$ $0$ 14-16       C clover       A. chrooc. $4.7726$ $4.6183$ $3.8063$ $0$ 17-19       F oats       A. beyer. $5.1701$ $5.2218$ $3.6232$ $1$ 18-20       C oats       A. beyer. $5.1701$ $5.2243$ $3.7309$ $1$ 22-24       C clover       A. beyer. $4.4394$ $4.4827$ $3.863$ $0$ 22-24       C clover       A. beyer. $4.4394$ $4.4827$ $3.8063$ $0$ 22-24       C clover       A. vine. $3.9267$ $3.9959$ $3.6232$ $0$ 22-31       F clover       A. vine. $2.4572$ $2.4817$ $3.8063$ $0$ 23-35       F oats       A. vine. $2.4572$ <td>.4525</td>	.4525
$\begin{array}{c c c c c c c c c c c c c c c c c c c $	.2638
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	7220
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	.7220 .5741
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$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	.5515
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	.5864
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	.6567
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	3427
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$	4485
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	.3411
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	.0131
37-39_         F clover         No. 26         4.4823         4.5271         3.8963         0           38-40_         C clover         No. 26         4.5722         4.6833         4.0046         0	
38-40. C clover No. 26 4.5722 4.6833 4.0046 0	6308
41-42 F 0ats No 97 2 6150 2 6000 0 6000 0	6787
	0590
42-44_ C oats No. 27 3.5720 3.6217 3.7309	
	6510
	4149
49-51. F oats	
50-52- C oats No. 22 3.5323 3.5519 3.7209	
	5478
	3820
	0193
58-60_ C oats No_ 4 3.5006 3.5323 3.7309	
61-63- F clover No. 4	6113
	3919
65-67. F oats Mixed culture 3.3865 3.4203 3.6232	
66-68_ C cats Mixed culture 3.6028 3.6021 3.7309 69-71_ F clover Mixed culture 4.3787 4.4123 3.8963 0.	F1.00
	5160
interest int	5748
75-76_ C. nothing Check 3.5085 3.5689 77-78_ F nothing Check 3.3002 3.3332	
79-80. C. nothing Check	

#### FIRST GROWING PERIOD.

The determinations for this period are shown in appendix table I and in condensed table X. As indicated by table IX, there was a steady increase in the total amount of nitrogen fixed in all the soils. This increase is still more marked if the last columns of tables IX and X are compared. The bacteria were increasingly active in fixing the free atmospheric nitrogen and in practically every case the total amount fixed due to the bacterial solution was more than doubled during the latter five weeks of this series.

These activities may be divided into two elasses, as the bacteria were more markedly affected by the presence of elover hay or of oats straw. In the first class A. chroococcum, A. chroococcum (HCM), No. 26, No. 22 and the Mixed Culture stood out prominently. None of these four organisms showed any fixation due to the presence of the decaying oats straw, but they did show appreciable gains due to the presence of the clover hay. The presence of the oats straw had apparently either inhibited the activities of the organisms or increased the activities of the other forms that are ineapable of fixing nitrogen for their own use and have utilized that fixed by the inoculating organisms. Organisms 4 and 27 showed a decided stimulation due to the clover hay and were able to utilize the oats straw as a source of energy.

A beijerincki and A. vinelandii were more markedly affected by the presence of oats straw. The stimulation of the activities of the former due to the presence of the decaying clover was parallel to that of the other organisms, and in addition the presence of the decaying oats straw stimulated its nitrogen fixing powers to over 250% of that of any other organism in the series with the single exception of A. vinelandii. On the other hand, A. vinelandii, while showing a marked stimulation due to the presence of the oats, also showed that the clover hay affected its activities much the same as the oats straw affected the other organisms, that is, the presence of the decaying clover hay in the cropped soils, deereased its nitrogen-fixing power, and in the fallow soils, completely inhibited it.

# SECOND GROWING PERIOD.

The results for this period are shown in appendix table II, the more important parts of which are repeated in condensed table XI. In comparison with the first growing period the results for the second period are decidedly lower thruout the second series. Not only are the total amounts of nitrogen found lower, but also the total amount of dry matter produced in the erop, indicating a possible direct relation between bacterial action and erop yields. These low results are explained by the fact that this series as grown during the hottest part of the summer, the pots being planted in the latter part of June and harvested during the earlier part of August. The results confirm those given in Table X except that in this series the only organism stimulated by the presence of the decaying oats straw was organism 27 in the cropped pots. Each of the inoculated organisms showed a direct stimulation due to the presence of the clover hay.

The organisms may be divided into two classes according as their activities are stimulated or retarded by the presence of

			Gram	s Nitrogen	per 10 lbs	s. soil.
Dupli- cate Pots	Tieatment	Bacterial inoculum used.	Nitrogen found.	Nitrogen in seed + amount removed by the growing crop.	Nitrogen in check and in oats and clover.	Nitrogen fixed by bacteria.
		1				
1- 3	F. oats	A. chroococcum (HCM)	2.7340	3,0128	3.3370	
2-4	C. oats	A. chroococcum (HCM)	2.8218	3.0916	3.1866	
5-7	F. clover	A. chroococcum (HCM)	3.3205	3.6592	3,6107	0.0485
6- 8	C. clover	A. chroococcum (HCM)	3.3123	3.6402	3.4603	0.1799
9-11	F. oats	A. chroococcum	2.6867	2.9607	3,3370	
10-12	C. oats	A. chroococcum	2.5479	2.7947	3.1866	
13-15	F. clover	A. chroococcum	3.5242	3.8836	3.6107	0.2729
1416	C. clover	A. chroococcum	3.1813	3.4901	3,4603	0.0298
17-19	F. oats	A. Beijerinckii	2,7155	2.9924	3.3370	
18-20	C. oats	A. Beijerinckii	2.6134	2.8626	3.1866	
21-23	F. clover	A. Beijerinckii	3.3343	3.6743	3.6107	0.0636
22-24 2527		A. Beijerinckii	3.2681	3.5857	3.4603	0.1254
26-28	F. oats	A. vinelandij	2.8235	3.1115	3.3370	
29-31	C. oats F. clover		2.6936	2.9763	3.1866	0.0201
30-32	F. clover	A. vinelandij	3.3205	3.6591	3.6107	0.0384 0.1719
33-35	F. oats	No. 26	3.4143	3.6322 3.0356	$3.4603 \\ 3.3370$	
34-36	C. oats	No. 26	$2.7547 \\ 2.8912$	3.0556	3.1866	
37-39		No. 26	3.4921	3.8492	3.6107	0.2385
38-40		No. 26	3.4921 3.0648	3.3747	3.4603	0.2000
41-43		No. 27	2,7015	2.9770	3.3370	
42		No. 27	3,0503	3.3429	3.1866	0.1563
45-47		No. 27	3.4487	3.8004	3.6107	0.1897
46-48		No. 27	3.5089	3.8613	3.4603	0.4010
49-51		No. 22	2.7083	2.9845	3.3370	0.10
50-52		No. 22	2,8984	3.1700	3.1866	
53-55	F. clover	No. 22	3.3977	3.7442	3.6107	0.1335
54-56	C. clover	No. 22	3.2667	3.5925	3.4603	0.1322
57-59	F. oats		2.6146	2.8812	3.3370	
58-60	C. oats		2,8246	3.1262	3.1866	
61-63		No. 4		3.7465	3.6107	0.1358
62-64	C. clover		3.1740	3.4913	3.4603	0.0310
65-67	F. oats		2.5210	2.7781	3.3370	
66-68		Mixed culture	2.6935	2.9310	3.1866	
69-71			3.2696	3.6031	3.6107	
70-72			3.3852	3.7159	3,4603	0.2556
73-74			2.8235	3.1115		
75-76			2.7518	3.0537		
77-78			2,9758	3,2793		
79-80	C. nothing	Check	2.7736	3.0364		

 TABLE
 XI-THE
 NITROGEN
 FIXED
 BY
 BACTERIA-SECOND
 PERIOD.

 (Condensed from appendix Table 2.)
 (Condense from appen

growing plants such as clover. In the first class are included A. chroococcum (HCM), A. beijerinckii, A. vinelandii, No. 27 and the mixed cultures. The first three organisms have had their nitrogen-fixing powers stimulated by the presence of the plants in practically the same ratio and have fixed similar amounts in both the fallow and cropped soil. A. beijerinckii showed the highest fixation of any of the eight for this series. The mixed cultures showed no fixation whatever in the fallow soils, but quite an appreciable amount in the cropped soils. No. 26, on the contrary, fixed an appreciable amount of nitrogen in the fallow soils but none at all in the presence of the growing oats plants. A. chroococcum and No. 4 showed the same stimulation under practically the same conditions, namely, that they possess a greater nitrogen-fixing power in the presence of decaying clover if no crop is grown upon the soil, while No. 22 was apparently neither stimulated nor retarded by either fallow or cropped conditions, but was affected by the presence of the decaying oats straw.

# THIRD GROWING PERIOD.

The results for this period are shown in appendix table III and condensed table XII. The total nitrogen content of the soil according to tables IX, X and XI, increased steadily through-

-			Gram	s Nitrogen	per 10	Pounds	Soil
Pots	Treatment	Bacterial Inoculum Used	N. found	N. in seed+ amt. re- moved by erop	N. in check and in oats and clover	N. fixed by bacteria	Av. N. fixed by bacteria
1	F cats	A. chrooc. (HCM)	2.3626	2.8648	2.8502	0.0146	
3	F oats	A. chrooc. (HCM)	2.7342	3.3165	2.8502	0.4663	0.2495
2	C oats	A. chrooc. (HCM)	2.3643	2,9647	3.3284		
4	C oats	A. chrooc. (HCM)	2.1930	2.6754	3.3284		
5	F clover	A. chrooc. (HCM)	2.9255	3.5486	3.1239	0.4247	
7	F clover	A. chrooc. (HCM)	2.8274	3.4296	3.1239	0.3057	0.3652
ů.	C clover	A. chrooc. (HCM)	3.2852	4.0691	3.6021	0.4670	
8	C clover	A. chrooc. (HCM)	2.8688	3.6142	3.6021	0.0121	0.2395
9	F oats	A. Chroococcum	2.7489	3,3344	2.8502	0.4842	
11	F oats	A. Chroococcum	2.3103	2.8024	2.8502		0.2421
10	O oats	A. Chroococcum	2.3652	2,9149	3.3284		
12	O oats	A. Chroococcum	2.3785	2.9537	3.3284	0.0050	
13	F clover	A. Chroococcum	2.8928	3.5089	3.1239	0.3850	0.5438
15	F clover	A. Chroococcum	3.1546	3.8265	3.1239 3.6021	0.7026	0.0438
14	C clover	A. Chroococcum A. Chroococcum	2.8489	$3.5649 \\ 3.4326$	3.6021 3.6021		
$\frac{16}{17}$	C Clover	A. beijerinckii	$2.7501 \\ 2.5789$	3.4320 3.1279	2.85021	0.2777	
19		A. beijerinckii	2.5789 2.4805	3.1279	2.8502 2.8502	0.1586	0.2182
19	F oats	A. beijerinckii	2.4805 2.2943	2.8469	3.3284	0.1550	0.2102
20	C cats	A. beijerinckii	2.2943 2.0737	2.5764	3.3284		
21	F clover	A. beijerinckii	3.5203	4.2705	3.1239	1.1466	
23	F clover	A. beijerinckii	3.0368	3.6836	3.1239	0.5597	0.8532

TABLE XII-THE NITROGEN FIXATION BY BACTERIA-THIRD PERIOD (Condensed from Appendix Table III),

TABLE XII-CONTINUED

			Gran	ns Nitrogen	Per 10	Pounds S	oil
Pots	Treatment	Bacterial Inoculum Used	N. found	N. in seed amt. re- moved by crop?	N. in check and in oits and clover	N. fixed by bacteria	Av. N. fixed by bacteria
22	C clover	A. beijerinckii	2.7429	3.4256	3.6021		
24	C clover	A. beijerinckii A. vinelandii	2.7694 2.5431	$3.4584 \\ 3.0847$	3.6021 2.8502	0.2345	
25 27	F oats	A. vinelandii A. vinelandii	2.3451	2.9691	2.8502 2.8502		0.1767
26	C oats	A. vinelandii	2.3917	2.9781	3.3284		
28	C oats	A. vinelandii	2.3491	2.9494	$3.3284 \\ 3.1239$		
29 31	F clover	A. vinelandii A. vinelandii	3.0172 3.0711	3.6598 3.7252	3,1239		
30	C clover	A. vinelandii	2.7429	3.4284	3.6021		
32	C clover	A. vinelandii	2.7363	3.3959	3.6021		
33 35	F oats	No. 26	2.4609 2.3823	2.9850 2.8887	2.8502 2.8502		0.0862
34 34	C oats	No. 26	2.3823	3.0673	3.3284		0.0002
36	C oats	No. 26	2.3321	2.9746	3.3284		
37	F clover	No. 26	2.9895	3.6162	3.1239		
39 38	F clover	No. 26 No. 26	2.0164 2.7749	2.4459 3.4477	$3.1239 \\ 3.6021$		0.2462
40	C clover	No. 26	2.8877	3.5888	3.6021		
41	F oats	No. 27	2.4478	2,9691	2.8502		
$\frac{43}{42}$	F Oats	No. 27 No. 27	2.4871 2.4583	3.0169 3.0736	2.8502 3.3284		0.1428
44	C oats	No. 27	2.4917	3.1595	3.3284		
45	F clover	No. 27	2.8143	3.4137	3.1239	0.2898	
47	F clover	No. 27 No. 27	3.0564	3.7074	3.1239		
$\frac{46}{48}$	C clover	No. 27 No. 27	$3.0181 \\ 2.8621$	$3.7421 \\ 3.5807$	3.6021 3.6021		0.0700
49	F oats	No. 22	2.4961	3.0278	2.8502		
51	F oats	No. 22	2.2186	2.6911	2.8502		
$\frac{50}{52}$	C oats	No 22 No. 22	2.5580 2.4451	$3.1751 \\ 3.0758$	3.3284 3.3284		
53	F clover	No. 22	2.7230	3.3029	3.1239		
55	F clover	No. 22	2.8667	3.4773	3.1239		
54 56	C clover	No. 22 No. 22	2.7893 2.6236	3.5060 3.2953	$3.6021 \\ 3.6021$		
57	C clover F oats	No. 22 No. 4	2.3430	2.8321	2.85021		
59	F cats	No. 4	2.4094	2.9126	2.8502	0.0624	0.0312
58	C oats	No. 4	2.3170	2.8086	3.3284		
60 61	C cats F clover	No. 4 No. 4	2.6037 3.0173	$3.3103 \\ 3.6599$	3.3284 3.1239		
63	F clover	No. 4	2.8208	3.4216	3.1239	0.2977	0.4168
62	C clover	NO. 4	2.8877	3.6172	3.6021		
$\frac{64}{65}$	C Clover F oats	No. 4 Mixed culture	2.8224 2.3496	$3.5098 \\ 2.8501$			
67	F oats	Mixed culture	2.3234	2.8183			
66	C oats	Mixed culture	2.4381	3.0435	3.3284		
68 69	C oats	Mixed culture	$2.3586 \\ 3.0761$	$2.8937 \\ 3.7313$	$3.3284 \\ 3.1239$		
71	F clover F clover	Mixed culture	2.7831	3,3759	3.1239		
70	C clover	Mixed culture	2,8366	3.5867	3.6021		
72	C clover	Mixed cilture	3.0306	$3.7664 \\ 2.8161$	$^{3.6021}_{*}$	0.1643	0.0822
$\frac{73}{74}$	F nothing F. nothing	Check	$2.3216 \\ 2.0944$	2.8161 2.5405	*	*	*
77	F nothing	Check	2.1074	2.5563	*	*	*
78	F nothing	Check	2.4085	2.9215	*	*	*
75 76	C nothing	Check	$2.7760 \\ 2.5375$	$3.3649 \\ 3.1041$	(† †	Ť	ŧ
79	C nothing	Check	2.4911	3.1522	+	+	t .
80	C nothing	Check	2.4901	3.1362	+	+	

\*Average four fallow checks 2.7086. †Average four cropped checks 3.1868.

out the first growing period, declined somewhat during the second, and according to table XII, there was a pronounced tendency to increase again during the third period. The crop response of this last period of growth confirms the results of the determinations, the amount of dry matter produced being practically midway between the production of the first and second growing periods. Figs. 1-6, which show the growth of oats in representative pots for the three periods, show that the first crop when ready to harvest was in the majority of cases leafy and heavy and showed a decided tendency to lodge; the second crop in the same stage of growth was somewhat dwarfed in appearance and with no indication of leafiness or weakness of stem: the third crop, while not as heavy as the first, showed all of its characteristics except that as a whole the production was more uniform and did not show the variation in the total dry weight of the harvested The bacterial activities, which are plotted in the tables erop. shown in fig. 7, varied in the same proportion as the crop response of the treated soils, being practically parallel with the production of the dried weight of the crop. The activities increased during the first growing period, declined thruout the second, but increased again during the third. The discussion of the third and last period of growth will be a combination of the activities of the inoculated bacteria as discussed in the first and second growing periods.

The last column in table XII indicates that each inoculated bacterial culture acted without exception in the same general manner instead of showing the expected variations. All of the inoculated bacteria fixed greater amounts of nitrogen in the soils to which clover hay was added as organic matter than in soils that were treated with the same amount of oats straw, and the growing crop on these soils reduced the nitrogen-fixing power of each and every one of these organisms. The activities of any one of the eight organisms used during the third period of growth would be an accurate measure for the activities of any of the others, a fact not even indicated in the other periods of growth.

Conclusion: Table XIII, recapitulating tables X, XI and XII, shows that inoculation, especially in fallow soils to which clover hay or oats straw was added, is not only possible but practical. The amounts of nitrogen shown in these tables are the actual amounts fixed by the organisms in ten pounds of soil and if these amounts are calculated on a 2,000,000 pound acre basis, the result is distinctly profitable. With proper soil conditions the greenhouse experiments can be duplicated in the field.

All of the organisms have shown an appreciable fixation of nitrogen but A. beijerinckii and A. vinelandii have been decidedly the most active. This finding confirms the suggestion of

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$\mathbf{BA}$	
TABLE XIH-NITROGEN CONTENT OF SOLLS AT END OF EACH GROWING PERIOD AND GAIN DUE TO THE INOCULATED BAC	
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DUE	
GAIN	
AND	
PERIOD .	OF SOIL.)
GROWING	0 POUNDS
EAOH	PER 1
ΟF	SIM
END	GRA
AT	IN.
T OF SOILS	TERIA. (RESULTS EXPRESSED IN GRAMS PER 10 POUNDS OF SOIL.)
EN CONTEN	(RESULTS
IH-NITROGI	TERIA. (RESULTIS EXPRESSED IN GRAMS FER 10 POUNDS OF SOIL.)
TABLE X	

Pots	Trea	Treatment '		Bacterial inoculum used	Original Nitrogen content	Nitrogen fo + amoun	Nitrogen found — that in seed + amount removed by grow- ing crop	at in seed by grow-	Nitrogen Average	Nitrogen fixed by bacteria Average of duplicate pots	bacteria ate pots
					+ oats or clover	First Period	Second Period	Third Period	First Period	Second Period	Third Period
1	F. oats -		Ā.		2.4910	3.3833	3.0128	3.0906			0.2405
2- 4	C. oats -		A.	<u> </u>	2.4910	2.9870	3.0916	2.8200			
	F. clover		Α.	~	2.7647	4.5488	3.6592	3.4891	0.4525	0.0485	0.3652
6-8	C. clover		¥.	chroococcum (HCM)	2.7647	4.2684	3.6402	3.8416 9.0604	0.2638	0.1799	0.2395
10 10 10 10 10 10 10 10 10 10 10 10 10 1	F. 0ats -		4 -	enroococcum	0.164.42	0.2002	1000.7	4000.0			U.2421
13-15	U. Oats -		Υ.	enroococam ehroococam	2.7647	o. eoo	3.8836	2.8043	0.7220	0.2729	0.5438
14-16	C. clover		Ā	chroococcum	2.7647	4.5787	3.4901	3.4987	0.5741	0.0298	
17-19	F. oats -		Ā	beijerinckii	2.4910	5.2218	2.9924	3.0683	1.5986		0.2182
18-20	C. oats -		A	beijerinckii	2.4910	5.2842	2.8626	2.6616	1.5515		
21-23	F. clover		A.	beijerinckii	2.7647	4.4827	3.6748	3.8770	0.5864	0.0536	0.8532
2224	C. clover		Ą.	beijerinckii	2.7647	4.6613	3.5857	3.4420	0.6567	0.1254	*********
25-27	F. oats		Ŀ.	vinelandii	2.4910	3.9659	3.1155	3.0269	0.3427		0.1767
26-28	C. oats -		Ą.	vinelandii	2.4910	5.1794	2.9763	2.9637	1.4485		
29-31	F. clover		A.	vinelandii	2.7647	2.4817	3.6591	3.6925		0.03S4	0.5686
30-32	C'. clover		A.	vinelandii	2.7647	4.3457	3.6322	3.4172	0.3411	0.1719	
33-35	F. oats		No.	26	2.4910	3.5235	3.0356	2.9368			0.0862
34-36	C. oats		No.	26	2.4910	3.5133	3.1730	3.0209			
37-39	F. clover	*******	No.	26	2.7647	4.5271	3.8492	3.0310	0.630S	0.2385	0.2472
38-40	C. clover		No.	20	2.7647	4.6833	3.3747	3.5382	0.6787		
41-43	F. oats -		No.	27	2.4910	3.6822	2.9770	2.9930	0.0590		0.1428
42-44	C. oats		No.	27	2.4910	3.6217	3.3429	3.1165		0.1563	
45-47	F. clover		N0.	22	2.7647	4.5473	3.8004	3.5605	0.6510	0.1897	0.4366
46-48	C. clover		No.	27	2.7647	4.4195	3.8613	3.6614	0.4149	0.4010	0.0700
49-01	F. oats		N0.		2.4910	3.4442	2.9845	2.8094			0.0555
60-92	C. oats		NO.		2.4910	3.0019	3.1700	3.1294	0.440	1005	0.0200
00000	r. clover			57	1101.7	1 9000	2002 0	1020.6	0.94/0	0.1200	700710
5750	C. CLUVEL		No.	T	0101 6	3 6495	6188 6	0001E.0	0.0193	0.1040	0.0312
58-60	C oats		S N	4	0101-6	3 3393	3 1969	3.0594			
61-63	F. Clover		No.		2.7647	4.5076	3.7465	3.5407	0.6113	0.1358	0.4168
62-64	O. clover		Nov	Ŧ	2.7647	4.3965	3.4913	3.5635	0.3919	0.0310	0.6076
65-67	F. oats		MIX	Mixed culture	2.4910	3.4203	2.7781	2.8342			
66-68	C. oats		Mix	Mixed culture	2.4910	3.6021	2.9310	2.9736			
	F. clover		Mix	Mixed culture	2.7647	4.4123	3.6031	3.5536	0.5160		0.4297
70-72	C. clover		MIX	Mixed culture	2.7647	4.5794	3.7159	3.6765	0.574S	0.2556	0.0822
73-74	F. nothing	20	Che	"heck	2.3494	3.6301	3.1115	2.6783			
75-76	O. nothing	B	Cheek	ek	2.3494	3.5689	3.0537	3.2395			
77-78	F. nothing		Check	ek	2.3494	3.3332	3.2793	2.7389			
79 80	C. nothing	g g	れしいけい		2.3404	3.6097	3.0364	3.1442			

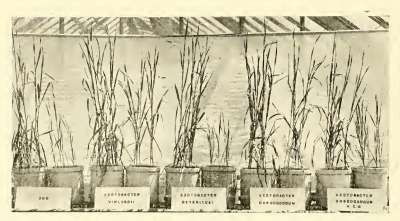


FIG. 1. Oats at end of first growing period, immediately before harvest; in pors 2, 6, 10, 14, 18, 22 26, 30, 24, 38

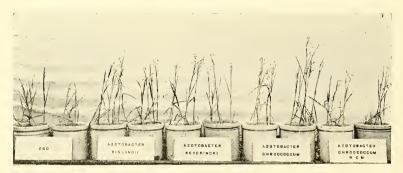


FIG. 2. Oats at end of first growing period, mmediately before harvest; pots 44, 48, 52, 56, 60, 64, 68, 72, 76, 80

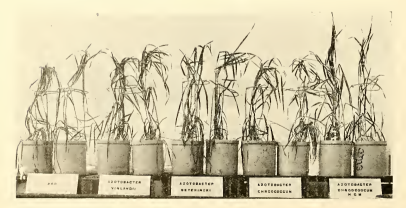


FIG 3. Oats at end of second growing period, immediately before harvest; pots 2, 6, 10, 14, 18, 22, 26 30, 34, 38

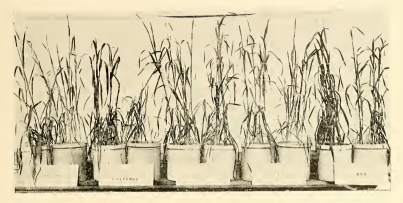


FIG. 4. Oats at end of second growing period, immediately before narvest; pots 42, 46, 50, 54, 58, 62, 66, 70, 75, 79



1G. 5. Oats at end of third growing period, mmediately before harvest; pots 4, 8, 12, 16, 20 24, 28, 32, 36, 40

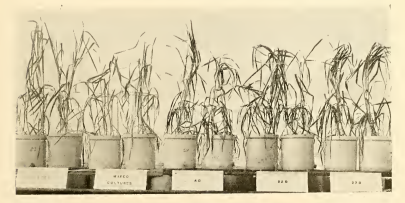


FIG. 6. Oats at end of third growing period, immediately before harvest; pots 44 48, 52, 56 60 64, 68, 72, 76, 80

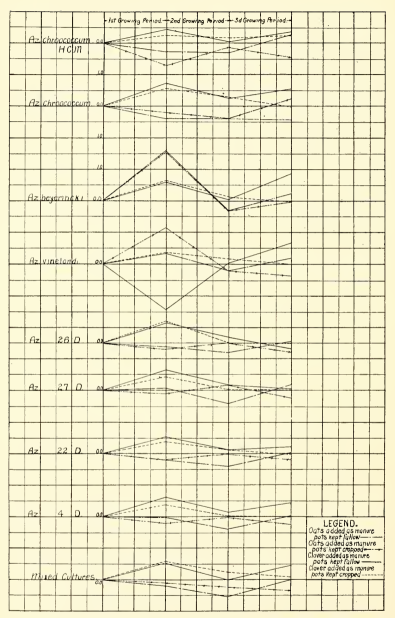


FIG. 7. This graph shows the variation in bacterial activity in the different growing periods

Lipman and Brown (41) by proving definitely that these organisms are capable of being profitably inoculated into field soils, provided that organic matter, carrying a sufficient amount of nitrogen as a stimulus, is supplied.

#### Summary.

1. When three crops of oats were grown continuously on this soil the nitrogen content of the soil increased during the first cropping period, decreased during the second, and increased slightly again during the third.

2. The nitrogen-fixing powers of the bacteria and the erop response were parallel with the total nitrogen content of the soil.

3. The nitrogen-fixing powers of some types of azotobacter and other large celled organisms of the same general character, were stimulated to a greater extent by the presence of decaying clover hay than of decaying oats straw.

4. The nitrogen-fixing powers of A. beijerinckii and A. vinelandii were stimulated to a greater extent by decaying oats straw than by clover hay, especially during the earlier stages of decomposition.

5. The nitrogen-fixing powers of the azotobacter and other large celled organisms of the same general type eventually became greater in fallow than in cropped soils.

6. The non-symbiotic nitrogen-fixing organisms of the azotobacter group were all eventually influenced in their activities in the same manner and by the same materials.

7. Soils may be profitably inoculated by azotobacter and other large celled organisms of the same type, the best effects being secured in this work by an inoculation with A. beijerinckii or A, vinelandii.

8. The conditions necessary for the greatest fixation are: Good environmental factors such as tillage, drainage, etc.; the presence of a rapidly decaying organic matter carrying a small nitrogen content, and freedom from growing plants.

# ACID EXTRACT, AMINO, NON-PROTEIN AND POLYPEP-TID NITROGEN CONTENT OF THE POT SOILS.

Introduction: The nitrogen of the soil is found in many complex combinations, in the determination of which the Bureau of Soils has isolated a large number of nitrogenous compounds and many different forms have been discovered. In investigating methods for the determination of amino acids and nitrates in a limed and unlimed soil, both with and without heavy applications of manures, Potter and Snyder (53) have found that they could accurately measure the amino nitrogen by a modification of the method devised by Kober and Sugiura (32). They discovered no tendency for the amino acid to accumulate under the conditions of the experiment. Accordingly in the present investigation determinations were made of the acid extract, non-protein, amino, and polypeptid nitrogen of some of the soils inoculated with the azotobacter cultures used in the greenhouse experiments, in order to prove this point and also to discover if the bacterial action had any effect on the accumulation or disappearance of these nitrogenous forms.

Soils used: Only the three soils inoculated with A. chroococcum, A. beijerinckü and A. vinelandii were analyzed.

# METHODS.

Acid extract: Place 166 gr. of air dried soil on a wetted double filter paper in a Buchner funnel and extract with 600 c. c. of a 1% HCl solution using gentle suction. Keep the soil barely covered with the solution and when extracted, wash with 200 to 300 c. c. of pure distilled water. Dry as quickly as possible, and determine the nitrogen content of the filtrate by the official salicylic acid method.

Alkali extract: The non-protein, amino, and polypeptid nitrogen determinations are based on the amounts extracted by a 1.5% NaOH solution. Shake 150 gr. of the air dried acid extracted soil with 600 c. c. of the NaOH solution and centrifuge to a clear solution. At least 210 c. c. of the clear solution must be obtained.

Non-protein nitrogen: Pipette off 25 c. c. of the alkali extract, neutralize with a sulphuric acid solution and add sufficient trichloracetic acid to make a 2.5% solution. To do this use 4.3 c. c., of a 1 3/10 N.  $H_2SO_4$  solution and 0.75 c. c. of a saturated trichloracetic solution. This method precipitates the proteins which are removed by filtering. Pipette 10 c. c. of the clear filtrate into large test tubes, add a couple glass beads, 2 drops of a 5% CuSO<sub>4</sub> solution, 1 c. c. C. P.  $H_2SO_4$ , and approximately 1 gr. C. P. potassium sulfate. Digest and distil as in the regular Kjeldahl method determining the ammonia colorimetrically.

Amino acid nitrogen: Pipette 80 c. c. of the alkali extract into 100 c. c. measuring flasks, neutralize with strong HC1 until neutral to litmus, add 7 c. c. saturated lead acetate solution, fill the flask to the mark with concentrated  $NH_4OH$  and shake vigorouly. Allow to settle for a few minutes then pass through double filter, using gentle suction and obtain at least 80 c. c. of the filtrate. Measure off 75 c. c. of this filtrate, add 25 c. c. saturated  $Ba(OH)_2$  and phenolphthalein as indicator and distill over steam bath under reduced pressure until there remains a volume of about 25 or 30 c. e. It is important that the reaction of the solution throughout this distillation should be at all times alkaline. Discard the distillate, wash residue into 100 c. c. graduate, cool, make up to 75 c. c., filter quickly to remove all carbonates, pipette 50 c. c. into 100 c. c. measuring flasks, make approximately neutral with N/10 HC1 and add 40 c. c. of buffer solution, stopper tightly and keep in cool place, if possible, on ice. (The buffer is made by dissolving 0.2 gr. molecules of boric acid in water, adding 100 c. c. of  $CO_2$  free N/10 NaOH solution and making up to 1000 c. c. with pure  $CO_2$  free water. Three volumes of this mixed with one volume of O/1/N HC1 makes the desired solution.)

Use pure water as cold as possible to prepare fresh the following solution: Place 10-20% copper chloride solution in 20-30 volumes cold water, add a few drops phenolphthalein and a saturated solution  $Ba(OH)_2$  until the purple color just forms. Centrifuge, decant off the clear liquid, wash with cold water and recentrifuge, repeating until there is no pink color formed by the addition of phenolphthalein in the wash water. Suspend the copper hydroxide in about 100 c. c. cold water and add approximately 1 c. c. to the cool flasks, shake vigorously, make up to the mark, and allow to warm up to the room temperature. Filter through No. 589 blue ribbon filter, pipette off 50 c. c. of the filtrate and determine the copper complex present as shown below as a measure of the amino nitrogen. Pipette off 40 c. c. of the filtrate for the determination of the polypeptid nitrogen.

Polypeptid nitrogen: Hydrolize the polypeptids into amino acids by adding approximately 5 c. c. concentrated  $H_2SO_4$  to the 40 c. c. and placing under a steam pressure of 8-10 pounds for 10-12 hours. Remove the excess acid with a saturated solution Ba(OH)<sub>2</sub> keeping the solution slightly alkaline to phenolphthalein, filter and wash with carbonated water at least three times. Evaporate the filtrate to about 35 or 40 c. c., place in 100 c. c. measuring flasks, neutralize with N/10 HC1, add 40 c. c. buffer solution, 1 c. c. of the copper hydroxide solution in the cold water as for the amino determinations and determine the copper present in the same manner.

Copper determination: Place the beakers containing the 50 e. c. on the hot plate, heat to boiling and neutralize with dilute  $HNO_3$ . Boil down to about one-half and add bromine water until a decided bromine color appears, evaporate to about 10-15 e. c., add 20-30 e. c. pure water and a little more bromine water and evaporate down again to 10-15 e. e. Cool, add 2-3 c. e. glacial acetic acid, a few crystals potassium iodide, a few drops of starch solution and titrate immediately with .001/N sodium thio-sulfate until the blue color disappears. Each c. c. of the .001/N thio-sulfate solution is the equivalent of 0.000028 gr, amino acid nitrogen. Preliminary determinations: In addition to the work on the soils, an unsuccessful attempt was made to determine the amount of non-protein and amino acid nitrogen fixed by the bacteria inoculated into the dextrose solution used in the other experiments. 250 c. c. of the dextrose solution was inoculated with the organisms indicated in Table 14 and incubated three weeks at room temperature. Enough e. p. sodium hydroxide was added directly to the solution to make 1.5% and the determination earried out in the above manner. A slight trace was the greatest amount found.

This table shows a decided increase in the soils under field conditions over the same soils in the dry state, the greatest increase taking place during the earlier periods of growth. The results of these determinations are grouped in three tables, each showing the amount of the different nitrogenous forms found at the end of each growing period.

Discussion of results: A comparison of the results given in Tables XVI, XVII and XVIII, shows that there was a definite variation of the nitrogenous forms with the length of the time of cropping. In almost every case the amount extracted by the acid varied with the length of time that the soil had been cropped, growing smaller and smaller, and the amino and polypeptid nitrogen gave similar results. The amount of these nitrogenous

TABLE XIV—AMINO ACID AND NON-PROTEIN NITROGEN FIXED BY THE PURE CULTURES IN SOLUTION.

Inoculum	Non-protein N. Amino Acid N.
1 A. chroococcum (HCM)	
2       A. chroococcum (HCM)         3       A. chroococcum (HCM)	
4 No. 26	trace
5 No. 26 6 No. 26	
7 A. chroococcum (HCM) and No. 26	trace trace
8 A. chroococcum (HCM) and No. 26	trace
9 A. ehroococeum (HCM) and No. 26	

TABLE XV—THE AMOUNT OF DIFFERENT NITROGENOUS FORMS IN THE SOIL AT THE BEGINNING OF THE EXPERIMENTS, ALSO THE SAME SOIL PLUS THE FQUIVALENT OF FIVE TONS GROUND OATS, STRAW OR GROUND CLOVER HAY ADDED TO THE SAMPLE. DETERMINATIONS BASED ON THE AMOUNT IN 25 GR. OF THE SAMPLE AND RESULTS FXPRESSED IN MG. NITROGEN AND IN PER CENT OF THE TOTAL NITROGEN.

Soil	Original Nitrogen content	Acid ex- N. mg. found	Per cent	Non-pro- tein N. mg. found	Per cent	Amino acid N. mg. found	Per cent	Poly-pep- tid N. mg. found	Per cent
Original Original + oats Original + red clover hay	12.95 13.75 15.40	$1.1242 \\ 1.1666 \\ 1.4424$	$8.7 \\ 8.5 \\ 9.3$	2.2275 2.3390 2.3475	$17.2 \\ 17.0 \\ 15.2$	$0.6840 \\ 0.1025 \\ 0.1050$	$0.7 \\ 0.7 \\ 0.7 \\ 0.7$	$0.2100 \\ 0.2550 \\ 0.2625$	$1.6 \\ 1.8 \\ 1.7$

TABLE XVI—AMOUNTS OF THE DIFFERENT FORMS OF NITROGEN IN THE CROPPED AND FALLOW INOCULATED SOLLS AT THE END OF THE FIRST PERIOD OF GROWTH. RESULTS EXPRESSED IN MG. NITROGEN FOUND AND IN PER CENT OF THE TOTAL NITROGEN CONTENT, BASED ON 25 GR. SAMPLE.

Pots	Total N. content 25 gr.	Acid ex- tract mg. found	Fer cent	Non-pro- tein mg. found	Per cent	Amiro acid mg. found	Per cent	Poly-pep- trd mg. tound	Per cent
$\begin{array}{c} 9-11\\ 10-12\\ 13-15\\ 14-16\\ 17-19\\ 21-23\\ 22-24\\ 25-27\\ 26-28\\ 29-31\\ 30-32\end{array}$	$\begin{array}{c} 14.35\\ 15.60\\ 20.39\\ 19.81\\ 23.03\\ 23.09\\ 19.77\\ 20.19\\ 17.50\\ 22.54\\ 20.94\\ 18.69\end{array}$	$\begin{array}{c} 2.0242\\ 1.9181\\ 2.2151\\ 1.9818\\ 2.2939\\ 1.1121\\ 2.3424\\ 0.9848\\ 2.4030\\ 1.9878\\ 2.3848\\ 0.7515\end{array}$	$\begin{array}{c} 14.1+\\ 12.3-\\ 10.8+\\ 10.0+\\ 9.9+\\ 4.7+\\ 11.8+\\ 4.8+\\ 13.7+\\ 13.7+\\ 13.8+\\ 11.4-\\ 4.1-\\ \end{array}$	$\begin{array}{c} 3.1675\\ 2.6650\\ 2.3690\\ 3.0650\\ 3.5000\\ 2.8500\\ 2.7175\\ 2.7325\\ 2.6950\\ 2.7425\\ 2.5150\\ 2.5875\end{array}$	$\begin{array}{c} 22.0\\ 17.0+\\ 11.6+\\ 15.4+\\ 15.2+\\ 13.8-\\ 13.5+\\ 15.4\\ 12.1+\\ 12.0+\\ 13.8+\\ \end{array}$	$\begin{array}{c} 0.1260\\ 0.1505\\ 0.1400\\ 0.1085\\ 0.1015\\ 0.1680\\ 0.2310\\ 0.1400\\ 0.1820\\ 0.2110\\ 0.1750\\ 0.0980 \end{array}$	$\begin{array}{c} 0.8+\\ 0.9+\\ 0.7-\\ 0.5+\\ 0.4+\\ 0.7+\\ 1.1+\\ 0.7-\\ 1.0+\\ 0.9+\\ 0.5+\\ \end{array}$	$\begin{array}{c} 0.4725\\ 0.2975\\ 0.4550\\ 0.2000\\ 0.2625\\ 0.2100\\ 0.4200\\ 0.3500\\ 0.4375\\ 0.3150\\ 0.2100\\ 0.3200 \end{array}$	3.2+ 1.9 2.2+ 1.0+ 1.1+ 0.9+ 2.1+ 1.7+ 2.5 1.4- 1.0 1.7+

TABLE XVII—AMOUNTS OF THE DIFFERENT FORMS OF NITROGEN IN THE FALLOW AND CROPPED INOCULATED SOLLS AT THE END OF THE SECOND GROWING PERIOD. RESULTS EXPRESSED IN MG. N. FOUND AND IN PER CENT OF THE TOTAL N. CONTENT BASED ON 25 GRAM SAMPLE.

Pots	Total N. content 25 gr.	Acid ex- tract mg. found	Per cent	Non-pro- tein mg. found	Per cent	Amino acid N. mg.found	Per cent	Polypep- tid N. mg.found	Per cent
9-11 10-12 13-15 14-16 17-19 18-20 21-23 22-24 25-27 26-28 29-31	$\begin{array}{c} 16.45\\ 15.44\\ 21.48\\ 19.32\\ 16.62\\ 15.84\\ 19.74\\ 19.78\\ 17.28\\ 16.32\\ 20.31\\ \end{array}$	$\begin{array}{c} 1.6515\\ lost\\ 2.1896\\ 0.7257\\ 0.9212\\ 0.7257\\ 1.7895\\ 0.7500\\ 1.4727\\ 0.6030\\ 2.1363\end{array}$	$ \begin{array}{r} 10.0 \\ \hline 10.2 \\ 3.7+ \\ 5.5+ \\ 4.6- \\ 9.1- \\ 3.8- \\ 8.5+ \\ 3.7- \\ 10.5+ \\ \end{array} $	$\begin{array}{c} 1.8375\\ 3.4150\\ 2.0700\\ 2.9925\\ 2.6825\\ 3.4075\\ 3.5000\\ 3.3350\\ 2.6370\\ 2.6370\\ 2.6000\\ 2.8350\end{array}$	$\begin{array}{c} 11.1+\\ 22.1+\\ 9.6+\\ 15.5-\\ 16.1+\\ 21.5+\\ 17.8-\\ 16.8+\\ 15.3-\\ 15.9+\\ 13.9+\\ 18.9+\\ \end{array}$	$\begin{array}{c} 0.0700\\ 0.1110\\ 0.1260\\ 0.1400\\ 0.4750\\ 0.0201\\ 0.1190\\ 0.0630\\ 0.2660\\ 0.0630\\ 0.0910\\ \end{array}$	$\begin{array}{c} 0.4+\\ 0.7+\\ 0.6-\\ 0.7+\\ 2.8+\\ 0.1+\\ 0.3+\\ 1.5+\\ 0.4-\\ 0.4+\\ \end{array}$	$\begin{array}{c} 0.2800\\ 0.3675\\ 0.1600\\ 0.1225\\ 0.1750\\ 0.3500\\ 0.2100\\ 0.2800\\ 0.2100\\ 0.2975\\ 0.2800 \end{array}$	$\begin{array}{c} 1.7+\\ 2.4-\\ 0.7+\\ 0.6+\\ 1.1-\\ 2.2+\\ 1.0+\\ 1.4+\\ 1.2+\\ 1.8+\\ 1.8+\\ 1.8+\\ \end{array}$

TABLE XVIII—AMOUNTS OF THE DIFFERENT FORMS OF NITROGEN IN THE FALLOW AND CROPPED INOCULATED SOLLS AT THE FND OF THE THIRD AND LAST GROWING PERIOD. RESULTS BASED ON 25 GRAM SAMPLE, EXPRESSED IN MG. N. FOUND AND IN PER CENT OF TOTAL N. CONTENT.

Pots	Total N. content 25 gr.	Acid ex- tract mg. found	Per cent	Non-pro- tein N. mg.found	Per cent	Amino acid N. mg.found	Per cent	Polypep- tid mg. found	Per cent
			1	1					
9-11	16.93	1.2833	7.5+	2.1475	12.7-	0.0420	0.2 +	0.2275	1.3 +
10-12	15.65	0.5773	3.7-	3.2850	20.9 +	0.0490	0.5 +	0.2100	1.3 +
13-15	20.23	0.7485	3.7-	2.3675	$11.2 \pm$	0.0420	0.2 +	0.2450	$1.2 \pm$
14-16	18.19	0.6306	3.4 +	3.3005	18.1 +	0.0910	0.4 -	0.2100	1.1 +
17-19	16.90	1.1773	6.9+	2.7025	15.9+	0.0210	0.1 +	0.2625	1.6-
18-20	14.94	0.5560	3.7+	3.6825	24.7—	0.0630	0.4 +	0.1400	0.9 +
21-23	22.03	1.7727	8.1-	2.8560	12.9+	0.0420	0.2-	0.2800	1.3-
22-24	18.21	0.7000	3.8 +	3.0300	16.5 +	0.3240	1.8-	0.1750	0.9+
25-27	16.71	1.5060	9.0 +	2.6925	16.1+	0.0350	0.2+	0.5950	3.6 -
26-28	15.66	0.7212	4.6+	3.6350	23.2+	0.0420	0.3-	0.3150	2.0+
29-31	20.36	2.2060	10.8 +	2.7400	13.4+	0.0560	0.4+	0.7075	3.4 +
30-32	18.09	0.5929	3.3	3.4500	11.3—	0.0770	0.4+	0.2800	1.5+

compounds became smaller, as decomposition of the organic matter proceeded, at a slightly faster rate than the total nitrogen content of the soil became depleted. The non-protein nitrogen also varied considerably, altho not in the marked degree shown by the other forms. Neither the oats straw or the red clover hav. added as manures to the pots, showed any effect on the forms of nitrogen determined, further than the small amount shown in the preliminary determinations. If there was a difference in the soil under field conditions it evidently was too small to be measured by these methods. It is entirely possible that the amounts of these complex nitrogenous compounds are rapidly changing into other forms and that the per cent they bear to the total nitrogen content remains somewhat constant, varying only with the amount of organic matter present in the beginning, then as decomposition proceeds and the more complex combinations are broken up, this percentage relation becomes smaller and smaller until it reaches a constant.

Once decomposition had begun in the soil there was absolutely no tendency for the more complex nitrogenous forms to accumulate under conditions approximating those in the field. Instead of an accumulation there was a steady reduction. How closely this reduction is coupled with the decay of the organic matter and what would be the final equilibrium between the total nitrogen content and the nitrogenous compounds are questions for further study.

## Summary.

1. The acid extracted, non-protein, amino and polypeptid nitrogen changed into other forms with the advance of decomposition much faster than the total nitrogen contents of the soils in question decreased.

2. Oats straw and clover hay added as manures at the rate of five tons per acre had little effect in influencing this change.

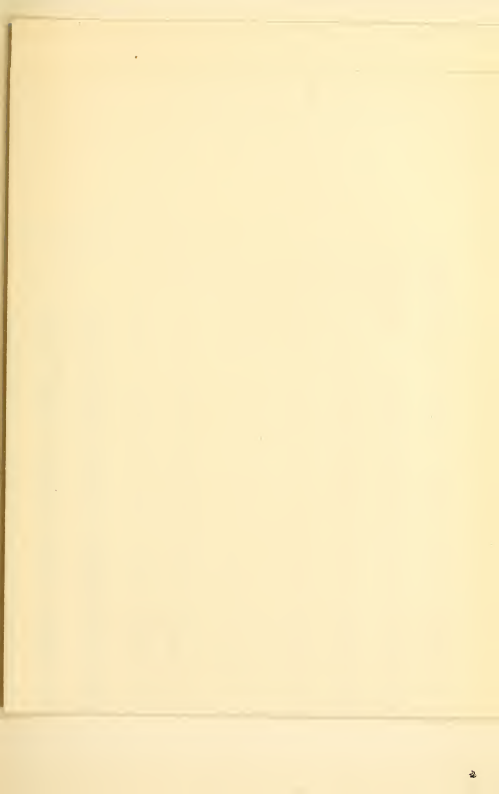
3. The amounts of non-protein and amino acid nitrogen fixed by bacterial cultures in solution were negligible.

4. Bacterial inoculation had apparently no effect on the amounts of non-protein, amino or polypeptid nitrogen in the soil.

5. There was no tendency for the above forms of nitrogen to accumulate in the soil under conditions approximating those in the field.

Acknowledgments: I wish to express my thanks to Dr. P. E. Brown for his help and suggestions thruout this work and to Dr. R. S. Potter for his suggestions in the determinations of the complex forms of nitrogen.

1.



#### APPENDIX TABLE 1.

			Actual on I	Basis of 449	Found in . 1 gr. in Fi cropped	Pot Soils, C illow and 4 -Pot	alculated 536 gr.			pot at ed in by	of ,530	hus con- ats or ts	cn- the
Inoculum Used and		Kind of Crop Grown	D	eterminatio	ons	css amt. N. added in sced	.в	crop	content grams	Total N. content of p sampling—N adder seed+N. removed b crop	on basis o of soil, 4,	Less check to which hus been added the N. con- tent of the dried oats or elover hay that was added to off-or pots	Grams N. gain due o tirely to action of inceulation
Pot No.	nt'	Grown				1.7	z	5	con	Log and	e e	the fldee hay	n ac
	tmc			ć		ldec	e g	wt.	I N.	Nigh a	ante z. v	che t of ver	N to the
	Treatment,		Orig	Dup.	Av.	Less	To <sup>4</sup> al pot	Dry wt.	Total N. o	Total san see cry	N, content orig. vol. gr.	Less bcc ten clo	Gran fire inc
A, chrooc. (HCM)		0-44	3.6309	0.3630	3,3499		3,5490						
1. 8	F	Oats	2.7658	3.2079	2.9883	0.0136	2.9747	0.8350	0.0123	3.3409 2.9570	3.3833 2.9870	3.6232 3.7309	
2- 4	F	Clover	4.2744	4.3373	4.3058		4.3055			4.3053	1.3485	3.8963	0.4525
6- 8	Ø	Olover	4.1071	4.2047	4.1809		4.1073	7.1150	0.1011	4.2684	4.2684	4.0046	0.2638
n 11	F	Oats	2.8601	3,5620 3.6310	3,2210		3.2210			3.2210	3.2532	3.6232	
	O F	Oats	3.4450 4.5663	4.5798	3.5380 4.5726	0.0136	3.5244	3.7400	0.0564	3.5808	3.5S0S	3.7309	
13-15	ô	Clover	4.4770	4.5087	4.4928	0.0136	4.1752	6.7600	0.025	4.5726 4.5787	4.6183 4.5787	3.8963	0.7220 0.5741
A hencildchil	F	Oats	5,1230	5,2173	5.1701		5.1701						
17-19	ō	Oats	5.1392	5.3343	6.2807	0.0136	5.2751	0.5250	0.003	6.1701 5.2824	5.2218 5.2824	3.6232 3.7309	1.5986 1.6515
91.28	F	Clover	4.5102 4.5881	4.3657	4.4394 4.5801	0.0136	4.4394 4.5665			4.4304	4.4827	3.8963	0.5564
20-24 A. vinc.anars	С	Clover				0.0130	4-5000	7.0100	0,0948	4.6013	4.6613	4.0046	0.0567
05.97.	F	Oats	8.9601	3.5973	3.9267		3.0267			3.9267	3.9659	3.6032	0.3427
96-28	0 F	Oats	5.1755 2.4051	2.4494	5.1120	0.0136	5.0984 2.4572	6,0050	0.0810	5.1794	ŏ.1794	3.750-7	1.4485
29-31	ĉ	Clover	4.2865	4.1912	4.2355	0.0136	4.2252	8.2950	0.1205	2.4572 4.3457	2.4817 4.3457	3.8563	0.3411
Az. 25D	F	Outs	3.4557	3.4857	3,4857		3.4887						010111
.43. 95	E C	Oats	3,4927	3.5463	3.5170	0.0136	3.5034	0.4900	0.0009	3.4897 3.5133	3.6235 \$.5133	3.6232	
37-39	F	Clover	4.4703	4.4944	4.4823		4.4823			4,4823	4.5271	3.8963	0.6305
38-40	C	Olover	4.6040	4.5405	4.5722	0.0136	4.5556	7.9650	0.1247	4.6833	4.6833	4.0046	0.6/87
Az. 27D	F	Oats	3,5830	3.7087	3.0458		3.6458			3.6458	3.6822	3.6232	0.0590
42-44	O F	Oats	3.5562 4.4473	3.5879 4.5573	3.5720 4.5023	0.0136	3.0581	3.7500	0.0633	3.6217	3.6217	3.7309	
45-47	F O	Clover	4.3.00	4.2729	4.3114	0.0130	4.5023 4.2978	9.0450	0.1217	4.5023 4.4195	$\frac{4.5173}{4.4195}$	3.8963	0.6510 0.4149
Az. 23D									0.1517				011110
49-51	FC	Oats	3.3630 3.4768	3,4573 3,5879	8.41(1 3.5323	0.0136	3.4101 2.5187	2.1850	0.0332	3.4101 3.5519	3.1442 3.5510	3.6232	
50+52	F	Clover	4.4310	4.3687	4.4001		6.4031	2.1850	0.0332	4.4001	4.4441	3.8963	0.5478
54-56	C	Clover	4.2729	4.3500	1.3114	0.0126	4.2978	6.7850	0.0858	4.3866	4 3866	4.0046	0.3520
Az. 4D 57-59	F	Oats	3.6144	3,5087	3,6065		3,6065			3.6065	3.6425	3.6232	0.0:93
58-60	õ	Oats	3,4927	3.5085	3.5006	0.0186	5.4876	3.8650	0.0453	3.5323	3.5323	3.736'1	
61-63	F	Clover	4.4316 4.2230	4.4944 4.3817	4.4630 4.3023	0.0136	4.2587			4.4630	4.5076	3.8963	U.8113
62-64 Mixed cultures	0	Ci0ver		areart	410020	0.0130	1.2001	7.1400	0.1073	4.3965	4 3965	4.0015	i <sup>1</sup> .3919
65-67	F	Ogts	3.3315 3.5562	3.4415 3.6514	3,3865		3.3865			3.3865	3.4203	3.6232	
60-68	O F	Oats	3.5562 4.3687	3.6514	3.6038 4.3687	0.0136	3.5902 4.3687	1.0450	0.0119	3.6021 4.3687	3.6021 4.4123	3.7309 3.8563	0.5160
70-72	ô	Clover	4.4135	4.5405	4.4720	0.0136	4.4584	9.5300	5.1210	4.57114	4.5794	4.0646	0.5748
Check	F	Nothing	3,6359	3,5515	3.5942		3,5943			3,5942	3,6301	Average 3.4816	
73-74	Č.	Nothing	3.5562	3.4609	3,5085	0.0136	3.5943	\$.5650	0.0740	3.5689	3.56SD	3.4816	
77-78	F	Nothing	3,3002	3.3002	3,3002		8.3002			3.3002	3.3332		
79-80	0	Nothing	3.4609	3.6197	3.5403	0.0136	3.6267	6.7700	0.0830	3.6097	3.6097		

#### APPENDIX TABLE II.

Inceulum Used and Pot No.			Actual G on J	Basis 4116	ound in F gr. in Fall op; ed.—F	ot Scils, Ca ow, 4161 g ots	alculated r. in	âms	of crop	lien sum- arnt. N. arnt. N. grow-	tume	o which has d the N, con- dried onts or added to	e en- af the
		Kind of Crop Grown	De	terminatio	)n			erop grams	content <b>jo</b> f	seed-	expres rig. vo 336 gr.	od the e dried / addee	sin cu c' ion t
	Treatment	Clown	Orig.	Dup.	Av.	Amt. added i seed	Total gr. N. in pot	Dry. wt. of	Total N co grams	Total N. in pot when sum- ple was taken-ant. N. ad-ied in seed-ant. N. removed by the grow- ing crop	Column 11 expresses of basis of orig. volume of soil, 4536 gr.	Less check to v been addied t tent of the di clover bay ac other pots	Grams N. gain cue en- tirely to act ion uf the inoculum
a. chrooc. (HCM)			0.5051	0.5000	0.0210		2.7340			2.7310	2 0100	0.0250	
1- 3 2- 4	P	Oats	2.7371 2.8537	2.7309 2.7900	2.7340	0.0136	2.8082	1.4995	0.0282	2.8364	3.0128 3.0916	3.3370 3.1866	
- 7	F	Olover	3,3854	3,2557	3.3205		3.3205			3.3205	3.6592	3.6107	0.0485
. ohroococcum	C	Clover	3.1070	3.2177	3.3123	0.0136	3.2987	2.1740	0.0410	3.3397	3.6402	3.4603	0.1709
0-11	P	Oats	2.6795	2,6939	2.6867		2,3567			2.6867	2.9607	3.3370	
)-12	0	Oats	2.6188	2.5771	2.5479	0.0136	2.5343	2.7877	0.0297	2.5640	2.7947	3.1866	
3-15	F	Clovel	3.6191	\$,5294	3.5242	0.0100	3.5242	1.9253	0.0240	3.5242 3.2020	3.8836 3.4901	3.6107	0.2729
-16	0	Clovet	3,1886	3.1740	3.1813	0.0136	3.1677	1.0200	0.0343	0.2020	5.2001	3.4603	0.0298
7-10	F	Oats	2.7227	2,7083	2.7155		2.7155	1		2.7155	2.9924	3.3370	
5-20_	Ø	Oats	2.6353	2.5916	2.6134	0.0136	2.5958	1.8895	0.0265	2.6263	2.8626 3.6743	3.1866 3.6107	0.0536
1-23 2-24	F	Olover	\$.2269 3.2322	3,4318 3,3051	3.8343 3.2681	0.0136	3.3313 3.2445	1.8340	0.0452	3.3343	3.5857	3.4603	0.1254
		C'over	0.2022	0.0001	012001	0.0400	014110	1.0010	010104	012001	010001	011000	0.1001
5-27	F	Oats	2.8379	2,8091	2.8235		2.8235			2.8235	3.1115	3.3370	
3-28	0	0ats	2.6936	2.6036	2.6936	0.0136	2.6800	2.0178	0.0506	2.7306 3.3205	2.9763 3.6591	3.1866 3.6107	0.0384
3-32	F	Clover	3.3277 3.4070	3,8183 3,4216	3.3205 3.4143	0.0136	3,4007	3.1785	0,0629	3.4636	3.6322	3.4603	0.1719
z. 26D		010 901	0.4010	014210	011130	010100		1	0.0020				1
3-35	F	Oats	2.7517	2.7577	2.7547		2.7547			2.7547	3.0356	3.3370	
4-36 7-39	F	Oats	2.8537	2.9255	2.8912 3.4921	0.0136	2.8776 3.4921	2.2545	0.0344	2.9120 3.4921	3.1730 3.8492	3.1866 3.6107	0.2385
8-40	ΰ	Clover	3.5443 3.0867	3.4400 3.0489	3 064S	0.0136	3.0512	2.2800	0.6149	3.0061	3.3747	8.4603	
12, 27D	1							1				1	1
2-44	F	Oats	2.6795	2,8235	2.7015		2.7015	1 0000		2.7015	2.9770	8.3370	0,1563
5-47	10	Outs Clover	2.9411 3.4574	3,1595 3 4400	3.0503	0.0136	3.0367 8,4487	1,6860	0.0302	3.0669	\$.3429 3.5004	3.1866 3.6107	0.1897
0-48-	- õ	Clover	3.5089	3.5089	3,5089	0.0136	3,4953	1.8225	0.0472	3.5425	3.8613	3.4003	0.4010
4z. 22D 9-51	1												
		Oats	2.7227	2.6939	2.7083	0.0136	2.7088 2.8743	1.8025	0.0005	2.7053	2.9845 3.1700	3.3370 3.1566	
	1 10	Clover	2.8683	2.9286 3.3977	2.8954 3.8977	0.0150	3.3977	1.0020	0.0335	\$.3977	3.7142	3,6107	0.1335
4.56. Az. 4D	- 0	Clover	3.3051	3.2323	3.2607	0.0136	3.2531	2.5103	0.0428	3.2959	3,5925	8.4603	0.1322
7-59	F	Octo					0 0110			0.0740	2.8812	3.8270	
		Oats	2.6218 2.8537	2.6074 2.7955	2.6146 2.8246	0.0136	2.6146 2.8110	3.0305	0.0571	2.0146 2.8681	3.1262	3.1665	
1-63. 2-64	- F	Clover	3.3854	3.4142	3.3998		3.3998		0.0571	3.3998	3.7465	3.6107	0.1358
		Clover	3.1740	3.1740	3.1740	0.0136	3.1601	2.2090	0.0427	3.2031	3.4013	3.4603	0.0310
55-67	F	Oats	2,5066	2,5354	2.5210		2,5210			2.5210	2.7781	3.3370	
		Oats	2.7081	2,5354	2.6935	0.0156	2.6709	2.2315	0.0422	2.7221	2.9310	3.1565	
19-71 70-72	- F	Clover	3,9696	3.2696	8.2896		3.2696			3.2696	3.6031	8.65 /7	
		Clover	3.4216	3.3188	3.3852	0.0136	3.3716	1.5933	0.0375	8.4091	3.7159	3.4 4 3	0.2556
13-74	F	Nothing	2.8091	2,8379	2.8235		2.8235			2,8235	3.1115	Average	
75-78		Nothing	2.7518	2.7518	2.7518	0.0136	2.7352	3.0903	0.0469	2.8285	3.0537	1	+
79-50	F	Nothing	2.9696	2,9820	2.9758		2.9758			2.9758	3.2793	1	1
	- 0	Nothing	2.7955	2.7518	2.7736	0.0136	2.7600	1.0500	0.0257	2.7857	8.0364	1	8

13.0450 + wt. Oats N. = 3.1860 13.1854 + clover N. = 3.6107 #3.0450 + clover N. + 3.4603

### APPENDIX TABLE III.

			Actual C on 1	rams N. F Basis of 37 gr. i	ound in P 41 gr. in Fi n Cropped	ot Soils, Ca allow and 3 Pots	doulated 786	Ę	_	seed"	ou	. con-	to ac-
Inoculum Used and Pot No.	nt	Kind of Crop Grown	De	terminatio	DS	d in	gr. N. in	of crop in	L content of n gr.	Total N. in pot at sampling-N. in se +N. removed by c	Total N. expressed o basis of crip. vol. c soil, 4536 gr/ ms	Less creek to which has been added the N. con- tent of the oats or clover added as manure	entirely loculum
	Treatment		Orig.	Dup.	Av.	N. added in seed	Total g	Dry wt.	Total N. c crop in g	Total N sampli +N.r	Total N basis o soil, 40	Less cue been a tent of clover manur	Gain due o tion of it nressed s
z. chrooc. (HCM)								)					
A	F	Oais	2.8300	2.3953 2.3852	2.3626 2.3643	0.0136	2.3626 2.3507	3,4020	0.1135	2.3626 2.4645	2.8648 2.9647	2.8502 3.3284	0.0146
23	F	Oats	2.5434 2.7619 2.1864	2.6965	2.7342	0.0136	2.7342	2.5110	0.0147	2.7842 2.2241	3.3165	2.8502	0.4063
5	O F	Clover	2.8928	2.1996 2.9583	2.1930 2.9255		2.1784 2.9255			2.9255	3.5486	3.8284 3.1239	0.4247
6	0 F		3.2094 2.8143	$3.2711 \\ 2.8405$	3.2852 2.8274	0.0136	3.2716 2.8274	3.0650	0.1109	3.3525 2.8274	4.0691 3.4246	8.6021 3.1239	0.4670
8 1z. chroococcum	Ŭ.	Olover	2.9417	2,7959	2.8658	0.0136	2.8552	8.4310	0.1492	3,0044	3.6142	8.6021	0.0121
1z. chroococcum 9	P	Oats	2.7489	2.7489	2.7489		2.7459			2.7459	3.8344	2.8502	0.4542
0	ō		2.3321	2.3984	2.3652	0.0136	2.3516	2.5500	0.0715	2.4231 2.8103	2.9149 2.8024 2.9537	3.8284	
0	F	Oats	2.3038 2.3586	$2.3169 \\ 2.3984$	2.3103 2.3785	0.0130	2.3103 2.3649	2.7790	0.090±	2.4553	2.9537	2.8502 3.8284	
3	FC	Clover	2.9059 2.8754	2.8798 2.8224	2.8928 2.8489	0.0136	2.8928 2.8353	3.7675	0.1251	2.8928	3.6089	3.1239 3.6021	0.3850
ō	F	Clover Clover Clover	3.1608	3.1285	3.1546		3.1546	3.3540	0.1173	3.1546 2.8534	3.8205	3.1230	0.7020
z. beijormehn	0		2.5971	2.7032	2.7501	0.0136	2.7361	0.0010	0.1115		3.4326	\$.6021	
4 5	FC	Oats	2.5918 2.2660	2.5656 2.3227	2.5787 2.2943	0.0136	2.5787 2.2807	3.7790	0.0858	2.5787 2.3665	3.1270 2.8469	2.8502 8.3284	0.2777
9	F	Oats Oats Oats Clover Clover Clover Clover	2.5162	2.4478	2.4805		2.4805	3.2680	0.0816	2.3665 2.4805 2.1417	3.0088 2.5764	2.8502 3.3284	0.158
0	UF	Oats	2.0804	2.0670 3.5203	2.0737	0.0136	2.0601 3.5203			3.5203	4.2705	3.1239	1.146
2	Ĉ	Clover	2,7429	2.7429	3.5203	0.0136	2,7293	3.4570	0.1183	2.8476 3.028	3.4256 3.6856	3.6021 3.1239	0.559
d	Č.	Clover	2.9976 2.7694	$3.0761 \\ 2.7694$	3.0368 2.7694	0.0136	3.0368 2.7553	3.1290	0.1191	2.8749	3.4085	3.6021	0.059
2 3	F	Oote	2.5132	2,5731	2.5431		2 5421			2.5431	3.0847	2.8302	0.234
	0	Oats	2,3851	2.3983	2.3917	0.0136	2.5431 2.3781	3.1550	0.0975	2.4756 2.4478	2.9781	3.3284	
8	F	Oats	2.4478 2.8264	2.4478 2.3719	2.4478 2.3491	U.0136	2.4478 2.3351	3.1330	0.1166	2.4517	2.9494	2.8502 3.3284	0.118
3 9 0	F	Clover	3.0107	3.0237 2.7429	3.0172 2.7429	0.0130	3.0172	3.8030	0.1289	3.0172	3.6598 3.4384	3.1239 3.6021	0.535
I	F	Oats Clover Clover Clover	2.7429 3.0499	3.0923	3.0711		3.0711	3.1575	0.1002	2.8582 8.0711	3.7252 3.3959	3.1239	0.601
2 Az. 26D	0	Clover	2.7297	2.7429	2.7363	0.0136	2.7227	0.107.0	0.1002	2.8229		3.6021	
12. 200 5. 14. 15. 16.	F	Oats	2.4478 2.4653	2.4740 2.5044	2.4609 2.4848	0.0136	2.4609 2.4712	8,4015	0.0785	2.4609 2.5497	2.9850 3.0673	2.8502 3.3284	0.134
ō	म	Oats Oats	2,3692	2.3954	2.3823		2,38:33	9.2160	0.4542	2.3523	2.8887	2.8302	0.038
16 7	ÔF	Oats Clover	2.\$586 2.9978	2.3056 2.9714	2.3321 2.9895	0.0136	2.3185 2.9895			$2.4727 \\ 2.9805$	2.9746 3.6162	3.3284 3.1239	0.492
18 19	C	Clover	2.7429 2.0102	2.7969 2.0027	2.7749 2.0164	0.0136	2.7113	3.9045	0.1296	2.8909 2.0164	3.4777 2.4459	3.6021 3.1239	
10 Az, 27D	F	Clover	2.9152	2.8602	2.8877	0.0136	2.0164 2.8741	2.9730	0,1091	2.9832	8,5888	3.6021	
11	F	Oats	2.4478	2.4478	2.4478		2.4478			2.4478	2.9691	2.8502	0.118
12 13 14 15	0 F		2.4513 2.4871	2.4653 2.4871	2,4583 2,4871	0.0136	2.4447	3.6430	0.1103	2.5550 2.4871	3.0736 3.0169	3,3284 2,8502	0.166
И	0	Oats	2.5189	2,4646	2,4917	0.0136	2.4871 2.4781	4.0130	0.1483	2.6264	8.1595	3.3284	
ίδ 46	F	Clover	2.7881 3.0814	2.8405 2.9548	2.8143 3.0181	0.0136	2.8143 3.0045	3.3770	0.1061	2.8143 3.1106	3.4137 3.7421	3.1239 3.6021	0.280
17 18	O F O	Clover	3.0499 2.9019	\$.0630 2.8224	3.0564 2.5021	0.0136	3.0564	\$,8060	0.1280	3.0564 2.9765	3.7074 3.5807	3.1239 3.6021	0.583
Az. 22D		Clover				0.0136	2.8435						
49	FO	Oats	2.5494 2.5448	2.4478	2.4961 2.5580	0.0136	2.4961 2.5144	2.9430	0.0940	2.4961 2.6393 2.2186	3.027S 3.1751	2.8502 3.3284	0.177
51	F	Oats Oats Oats	2.2383 2.4110	2.5713 2.1901	2.5580 2.2186	0.0136	2.5444 2.2186	8.7480	0.1258	2.2486 2.5568	2.6911 3.0758	2.8502 3.8284	
60. 51	F	Clover	2,7032 2,7827	2.4786 2.7429 2.7959	$2.4451 \\ 2.7230$		2.4915 2.7230	4.0300	0.1387	2.7230	3.3029	3.1239	0.179
55	F	Clover	2.7827 2.8067	2,7959 2,8667	2.7893	0.0136	2.7757 2.8607			2.9144 2.8667	3.5060 3.4773	3.6021 3.1239	0.353
54	i õ	Clover	2.6767	2.5706	2.6236	0.0136	2.6100	3.6470	0.1292	2,7392	3,2953	8.6021	
57	F	Oats	2.8109	2.3692	2.3430		2.3430	2.3930		2.3430	2.8321	2.8502	
58 50	OP	Oats Oats Oats	2.3000 2.3602	2.3321 2.4497	$2.3160 \\ 2.4094$	0.0136	2.3024 2.4094		0.0426	2.8450 2.4094	2.8086 2.9126	3.3284 2.8502	0.062
60	. Û	Oats	2,6104	2.5971	2.6037	0.0136	2,5901	4.4220	0.4616	2.7517 3.0173	3.3103	8.3284	
62	P	Clover	2,8602	3.0237 2.9152 2.8374	3.0173 2.8877	0.0136	3.0173 2.8741	4,3300	0.1327	3.0068	3.6599 3.6172	3.1289 3.6021	0.5360
6564	FC	Clover Clover	2.8140 2.7959	2.8274 2.8489	2.8208 2.8224	0.0136	2,8208	3.1685	0.1087	2.8208 2.9175	3.4216 3.6008	3,1239 3,6021	0.2977
97 59 50 60 61 62 62 63 64 Mixed cultures 65	1 1		2.0431			0.0130					1		
66	ÎÔ	Oats	2.646	2.3562 2.4116 	2.2496 2.4381	0.0136	2.3496	3.4770	0.4055	2.3496 2.5300	2.8501 3.0435	2.8502 3.3284	
65 66 67 68 60	F	Oats Oats	2,3189	2.3944	2.3234 2.3586	0.0136	2.4245 2.3234 2.3450	2.9495	0.0604	2.3234 2.4054	2.8183 2.8937	2.8503 3.3284	
60	F	Clover	3.0199		8.0761		3.0761			3.0761	3,7313	8,1239	0.6074
70	F	Clover	2.8773 2.7459	2,7959 2.8274	2.8366	0.0136	2.8230 2.7831	4.3356	0.1385	2.9815 2.7831	3.5807 3,3759	3.6021 3.1239	0,2520
70 71 72 Check	Ô	Olover	2.8947	3.0606	3.0806	0.0136	3.0170	2.9190	0.11:8	3,1208	3,7061	3,6021	0.4643
		Nothing	2.3189	2.3244	2.3216		2.3216			2,8216	2.8161 2.5405		
74 76 76 77	FC	Nothing	2.0944 2.7691	2.0044 2.7827	2.0944 2.7760	0.0136	2 0044	1.6555	0.0339	2.0944 2.7963	2.5405 S.2649		;
76	O P		2.5574 2.1071	2.7827 2.5176 2.1074	2.5375	0.0136	2.7624 2.5239	2.8740	0.0339	2.5803	3.1041		
77	F	Nothing Nothing	2.4216	2.3954	2.1074 2.4085		$2.1074 \\ 2.4085$			2.1074 2.4085	2.5563 2.9215		
	0			2.4646	2,4911	0.0186	2.4775	3,9540	0.1426	2.6211	3,1522		

Av. ck, fallow 2.7086 + N, content of cats ground  $0.1415{=}2.8502$  Av. ck, fallow 2.7086 + N, content ground clover  $0.4153{=}3.1230$ . Av. ck, cropped 3.1888 + N, content ground cats  $0.1416{=}3.2834$ . Av. ck, cropped 3.1868 + N, content ground clover  $0.4153{=}3.5621$ 

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