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INTRODUCTION

Some YEARS ago anhydrous ammonia (NH_3) was introduced in California as a nitrogen fertilizer. This compound, sold as "Agricultural Ammonia" and "Shell NH_3 ," is available in the form of a liquefied gas compressed in strong steel cylinders that weigh about 150 pounds.

Agricultural Ammonia is applied on the land during irrigation. The cylinders are connected to a simple metering device, and the ammonia is released as a gas into the irrigation water. Thus the ammonia is distributed in a very dilute solution (100 to 500 p.p.m.). In water ammonia (NH_3) forms ammonium hydroxide (NH_4OH) .

Every new method of fertilization unfolds problems for investigation. The mere fact that a new type of fertilizer initially produces good economic returns is no guarantee of lasting efficiency. Since the continued use of a given fertilizer may deteriorate the chemical and physical properties of a soil and impair the crop yields, it is, therefore, of utmost importance to acquire a clear understanding of the reactions that take place and the resulting changes in soil properties.

In this paper are discussed certain fundamental problems of ammonia fertilization, especially those phases that require laboratory investigation. For the purpose of comparison with commonly used nitrogenous fertilizers, all experiments with ammonia (NH_3) were accompanied by corresponding experiments with ammonium sulfate $[(NH_4)_2SO_4]$ or ammonium chloride (NH_4Cl) . The various topics investigated may be conveniently grouped as follows: (1) the extent and chemical nature of ammonia fixation in soils; (2) the availability of ammonia in soils to plants and microörganisms; (3) the influence of ammonia on physical soil characteristics, especially permeability.

FIXATION OF AMMONIUM COMPOUNDS IN SOILS

Experimental Procedure.—To 50-gram portions of air-dry soil were added electrolyte solutions containing 2 to 500 milliequivalents of NH_4OH or $(NH_4)_2SO_4$. All systems were brought to a volume of 1 liter. The suspensions were kept at room temperature for 24 hours and were shaken occasionally. Samples of clear supernatant liquid were obtained by centrifuging or by means of Pasteur-Chamberland filters. According to special tests, no NH_3 was lost under the experimental conditions. The NH_3 was determined by distillation with MgO into standard acid, using methyl red as an indicator.

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Influence of Soil Texture and Acidity.—Of the twenty-three soils investigated, three typical examples are shown in figure 1. Two features are outstanding: First, the total amount of nitrogen taken up varies with the kind of soil; second, the compounds NH_4OH and $(NH_4)_2SO_4$ behave differently in different soils. The Aiken clay loam, for example, fixes larger amounts of nitrogen from NH_4OH than from $(NH_4)_2SO_4$, whereas certain other soils,



Fig. 1.—Typical examples of uptake of NH_4 from ammonium hydroxide and from ammonium sulfate by various soil suspensions. Note the variations in the relative position of NH_4OH and $(NH_4)_2SO_4$.

such as Yolo clay loam, fix larger amounts from $(NH_4)_2SO_4$. A few soils, for instance the Holland sandy loam, behave indifferently towards the two nitrogen compounds. These relations may well be examined in greater detail.

As has long been known, the adsorption of many electrolytes depends on the texture of the soil, particularly its colloid fraction. In this regard NH_4OH and $(NH_4)_2SO_4$ are no exceptions. This fact is clearly brought out in figure 2, which depicts the adsorption of nitrogen as a function of soil texture, the latter being expressed in terms of moisture equivalents. In general, the fine-textured soils exhibit higher nitrogen adsorption than the coarse-textured ones, from NH_4OH as well as from $(NH_4)_2SO_4$; but the correlation is far from

perfect. Judging from the distinct scattering of the data, other factors besides texture influence adsorption.

Inasmuch as solutions of NH_4OH are alkaline, whereas those of $(NH_4)_2SO_4$ are acid, there would probably be a correlation between the pH of the soil and NH_4 adsorption from the two solutions. This situation is well illustrated in



Fig. 2.—Relation between uptake of nitrogen and soil texture, the latter being expressed as moisture equivalents. Data refer to soil suspensions containing 50 grams of soil and 50 milliequivalents of nitrogen in a volume of 1 liter.

figure 3. On the abscissa is plotted the pH value of each soil;⁶ and on the ordinate is indicated the difference between the NH_4 adsorption from $(NH_4)_2SO_4$ and the corresponding NH_4 adsorption from NH_4OH , at a NH_4 concentration of 50 milliequivalents in 1,000 cc of solution (compare fig. 1). As may be clearly seen, the relative NH_4 adsorption from the two electrolyte solutions depends upon the pH of the soil. Acid soils tend to adsorb more nitrogen from NH_4OH solutions, whereas alkaline soils favor $(NH_4)_2SO_4$.

^{*} Soil-water ratio is 1:2; pH measured with a glass electrode.

Again wide differences exist among the soils of each group. Probably the crystalline nature of the colloidal material comes into play. The full significance of the nature of the clay colloids will become apparent later in this discussion.



Fig. 3.—Comparison of nitrogen uptake from $(NH_4)_2SO_4$ and NH_4OH as related to the pH of the soil. On the abscissa are plotted the pH values of the various soils, on the ordinate the difference "nitrogen uptake from $(NH_4)_2SO_4$ minus nitrogen uptake from NH_4OH ."

Effect of Soil-Water Ratio.—The data hitherto presented refer to systems composed of 50 grams of soil in 1,000 cc of solution. Under practical conditions of irrigation the proportion of soil to water differs from that of the systems mentioned previously. The amount of the solid phase exceeds that of the liquid phase. To ascertain how the ratio of soil to water affects NH_4 adsorption,

750 cc of a solution containing 6.67 milliequivalents of electrolyte per liter was added to each of a series of soil samples having different weights. Figures 4 and 5 show the results obtained for Aiken clay loam and Yolo clay loam. The percentage of nitrogen adsorbed is plotted as a function of cubic centimeters of solution per gram of soil. The electrolyte concentration of 6.67 milliequiv-



Fig. 4.—Percentage adsorption of NH_4 as conditioned by the soil-water ratio; 750 cc solution containing 6.67 milliequivalents of NH_4 in equilibrium with various amount of Aiken clay loam.

alents per liter is equal to 114 p.p.m. of NH₃, a value frequently used in irrigation practice.

Both figures 4 and 5 reveal a pronounced uptake of nitrogen at higher soil contents. Aiken clay loam maintains its preference for nitrogen from NH_4OH throughout the range of soil-water ratios investigated. With Yolo clay loam, the NH_4OH and $(NH_4)_2SO_4$ curves cross each other.

The soil-water ratios selected for these measurements pertain to systems

that range from suspensions to very thick muds. It is difficult if not impq sible to measure NH_4 adsorption at such low moisture values as prevail under field conditions. Technical difficulties of extracting the soil solutions are greatly increased by such phenomena as hydration effects and perhaps Donn4 equilibria.



Fig. 5.—Percentage adsorption of NH₄ as conditioned by the soil-water ratio; 750 cc solution containing 6.67 milliequivalents of NH₄ in equilibrium with various amounts of Yolo clay loam.

To show the approximate intensity of nitrogen fixation at moisture percentages corresponding to field conditions, figure 6 was constructed. It is based on the fact (to be demonstrated in subsequent sections) that the adsorption of NH_4 ions involves a base-exchange reaction, which some investigators $(7, 9)^7$ have described quantitatively by an equation. With the assumption that the base exchange equation obtains throughout the moisture range under consideration, figure 6 plots the calculated fixation of NH_4 by a clay soil

⁷ Italic numbers in parentheses refer to "Literature Cited" at the end of this paper.

having a base-exchange capacity of 30 milliequivalents per 100 grams and by a sandy soil with a capacity of 3 milliequivalents per 100 grams. On the abscissa are indicated the moisture percentages (on dry bases). By division by 100, these may be readily converted to the scales given in figures 4 and 5. The ordinate indicates the percentage of NH_4 adsorbed from 6.67 milliequivalents of $(NH_4)_2SO_4$. The curve calculated for the clay soil agrees well with the



Fig. 6.—Extension of soil-water relations shown in figure 5 to field conditions (small values of soil-water ratios).

extrapolated curve of figure 5. The curve for the sandy soil is considerably below that of the clay soil, a feature in harmony with the adsorption data presented in figure 1. The graph also lists the approximate moisture percentages for the field capacity and for the soil system saturated with water. The latter corresponds to the water conditions during irrigation; the former reflects the approximate degree of moisture a few days thereafter.

From the experimental as well as the theoretical evidence, one may safely conclude that under field conditions the percentage fixation of NH_4 assumes very high magnitudes.

Distribution of NH_4 in Relation to Depth.—Many farmers firmly believe that NH_3 dissolved in irrigation water distributes itself in the soil in ac-

cordance with the depth of penetration of water. Although such behavior is characteristic of nitrates (NO_3) , it need not occur with NH_3 and with NH_4 compounds.

To obtain a quantitative estimate of the distribution of ammonium fertilizers in relation to depth, brass cylinders 1 inch in diameter and 6 to 8 inches in length were filled with various types of soils (400 grams). The brass cylinders consisted of sections of short cylinders, each 1 inch long—an arrange-



Fig. 7.—Amount of nitrogen found in soil sections in percentage of nitrogen added.

ment that permitted the entire soil column to be sliced into 1-inch segments whenever desired. Through these columns was passed 500 cc of NH_4OH or $(NH_4)_2SO_4$ solution containing 7.5 milliequivalents of nitrogen (127 p.p.m. NH_3). This volume corresponds to an irrigation of 10 acre-inches. After the experiment the column was taken apart, and the 1-inch segments were analyzed individually.

The soils investigated included textures from sandy to clay. The following twelve types were examined: Oakley sand, Goldridge fine sandy loam, Hanford fine sandy loam, Fresno fine sandy loam, Yolo-sandy loam, Vina silt loam, Yolo silt loam, Ramona loam, Yolo clay loam, Aiken clay loam, Aiken clay, and Altamont adobe clay. The results are summarized in the characteristic patterns given by figure 7. Nitrogen penetration is evidently deep in light soils and shallow in heavy ones. In sands (fig. 7, A), nitrogen distributes itself rather evenly throughout the 7-inch column; and, furthermore, the percolating solution contains appreciable nitrogen. In medium-textured soils, as illustrated by Yolo sandy loam (fig. 7, B), the bulk of the nitrogen, in the form of NH₄OH or (NH₄)₂SO₄, is held in the first 3 inches of soil. The percolating solution is almost free of nitrogen. This behavior is especially characteristic of heavy-textured soils such as Yolo clay loam (fig. 7, C), in which nitrogen penetration is restricted to the first 2 inches. The lower portions of the column receive no measurable amounts of nitrogen. These observations agree with the previously mentioned studies which demonstrated that the fixation of NH₄ is directly related to the colloid content of the soil.

The Aiken series, which has an acid reaction and contains kaolinitic rather than montmorillonitic clay particles, deviates from the behavior of the neutral Yolo soils in one important respect. NH_4OH and $(NH_4)_2SO_4$ penetrate at widely different rates. The major fraction of the NH_4OH is held in the surface inch of soil, whereas in the columns receiving $(NH_4)_2SO_4$, the second inch contains nearly as much nitrogen as the first. Moreover, $(NH_4)_2SO_4$ percolates much deeper in Aiken clay loam than in any other soil of similar colloid content. These findings harmonize with figure 3, which emphasizes the role played by soil acidity and, perhaps, by the crystalline nature of the clay minerals.

The question arose whether subsequent leachings with distilled water would release the adsorbed NH_4 and carry it deeper. To examine this possibility, ammoniated soils were prepared by leaching 1-inch soil columns (Yolo clay loam and Aiken clay loam) with 250 cc of NH_4OH or $(NH_4)_2SO_4$ containing 12 milliequivalents of nitrogen per liter. For either electrolyte, over 98 per cent of the nitrogen applied was held by these 1-inch columns. The only exception was Aiken clay loam, which retained only 62.6 per cent of the NH_4 applied as $(NH_4)_2SO_4$.

Immediately after percolation had ceased, 250 cc of distilled water was passed through the column, and the new percolate was analyzed for nitrogen. Since in Yolo clay loam only about 6 per cent of the nitrogen was removed by this treatment, evidently the adsorbed NH_4 ions were rather tightly held. The leachates of Aiken clay loam contained even less nitrogen—1.6 per cent only.

In medium- and fine-textured soils, then, NH_4 apparently does not follow the movement of the irrigation water, but accumulates in a relatively shallow surface zone. After a time, fortunately, nitrification takes place; and part of the adsorbed nitrogen is converted to NO_3 , which may be carried readily into deeper horizons.

Often the irrigation water contains dissolved salts, which theoretically might hinder the adsorption of NH_4 by way of cation competition. Tests conducted with $(NH_4)_2SO_4$ and NH_4OH in presence of $CaCl_2$ and $NaHCO_3$ (5 to 15 milliequivalents per liter of solution) did not, however, reveal any significant effect on the distribution of NH_4 in the soil columns.

MECHANISM OF FIXATION OF AMMONIUM COMPOUNDS

Because of the close correlation between soil texture and the adsorption of cations from electrolyte solutions, soil chemists have intensively investigated the base-exchange properties of the colloidal fraction of the soil. They have shown repeatedly that the colloidal clay and humus particles govern the fixation processes of ammonium ions. It will prove advantageous to discuss separately the behavior of NH_4OH and that of its salts, the sulfates and the chlorides.

BEHAVIOR OF AMMONIUM SALTS

Since the days of Thomas Way (1850) chemists have known that the fixation process of $(NH_4)_2SO_4$ and NH_4Cl involves the cation (NH_4) only. The anion does not participate. The mechanism involves an exchange process. For a given amount of NH_4 taken up, an equivalent amount of other cations is released by the colloidal particle. A typical reaction may be formulated as follows:



The extent of the reaction (from left to right) depends on numerous factors such as salt concentration, clay-water ratio, and temperature. Quantitative descriptions have been formulated by various investigators (7, 9).

For illustration we may present the data obtained with Yolo colloid and Aiken soil. The colloidal fraction $(\langle 2\mu \rangle)$ of Yolo clay loam and the samples

(Total volume of systems = 500 cc) Nature of system and Exchange-Hydrolysis Electrolyte NH₄ K adsorbed released amount of clay or soil used able K added m.e. m.e. m.e. m.e. m.e. K-Yolo clay,* grams: 0.41 NH₄Cl 4.50 2.07 8.29..... 4.50 1.89 8.29..... 4.50 0.41 NH4OH 4.50 1.75 0.68 K-Aiken soil,* grams: 1.07 (NH₄)₂SO₄ 5.11 1.84 1.99 5.11 5.11..... 1.07 NH4OH 5.11 1.69 5.11..... 5.110.61 K-Aiken soil, † grams: (NH₄)₂SO₄ 5.18 1.35 5.18 1 66 1.81 25.00..... 5.18 1.35 NH4OH 5.18 2.12 0.62 25.00....

TABLE 1 Cation Exchange between K Systems and Ammonium Compounds

* Prepared by leaching with KCl.

† Prepared by leaching with potassium acetate.

of Aiken clay loam were transformed into K systems—that is, systems containing only exchangeable potassium ions. These K-clays hydrolyze to the extent shown in table 1. After the addition of NH_4Cl or $(NH_4)_2SO_4$ or NH_4OH , the reaction was allowed to proceed for several days at constant temperature. The solid particles were then separated by ultrafiltration or centrifugation, and potassium and NH_4 were determined in the ultrafiltrate or in the supernatant liquid. Table 1 gives the amounts of NH_4 taken up and of potassium released (corrected for hydrolysis).

Focusing attention on NH_4Cl and $(NH_4)_2SO_4$, we see that the exchange is practically stoichiometric. For every NH_4 ion taken up, a potassium ion has been released from the clay particles. This holds true both for the Yolo colloid and for the Aiken soil.

Calcium clays behave likewise. In an experiment by W. P. Kelley,^{*} 10 grams of Yolo soil possessing 2.94 milliequivalents of exchangeable calcium ions (but



Fig. 8.—Amounts of NH, adsorption from NH₄Cl in three clay systems. The fixation of NH, depends on the nature of the cation on the colloidal clay.

no other exchangeable cations) reacted with 2.94 milliequivalents of NH_4Cl in 100 cc volume. The NH_4 adsorbed amounted to 0.97 milliequivalent; the calcium released to 1.01. Again the exchange took place in stoichiometric proportions. For every divalent calcium ion released, two monovalent NH_4 ions were taken up by the soil.

As will be noted, the percentage adsorption of NH_4 is higher in the K system than in the Ca system—namely, 46 per cent as compared with 33. In other words, NH_4 fixation depends on the nature of the exchangeable ion on

⁸ Kelley, W. P. Personal communications.

the clay. Figure 8 well illustrates this fact. It refers to pure K-clay, Ca-clay, and H-clay systems (500 cc), each containing 7.5 grams of clay (montmorillonite type) and 4.50 milliequivalents of exchangeable cations. H-clay, which corresponds to an acid soil, fixes the smallest amount of NH_4 ion from NH_4Cl as well as from $(NH_4)_2SO_4$. In the farming regions of California, where soil leaching is not pronounced, the curve depicting the reaction with Ca-clay may be taken as the most appropriate example.

Importance of the Degree of Saturation, with Special Reference to Potassium.—From the viewpoint of fertilizer practice, it is interesting to examine more closely certain aspects of the liberation of adsorbed cations such as potassium by additions of ammonium fertilizers.

As shown in figure 8, NH₄ salts added to a pure K-clay release considerable potassium. The liberation may be materially reduced if, besides potassium, other cations also are present on the clay particles. The proportion of exchangeable potassium in relation to the other exchangeable cations is known as the degree of saturation of potassium. It is usually expressed on a percentage basis. Thus if a given soil contains 20 milliequivalents of exchangeable cations, and if 5 milliequivalents of these are potassium, then the degree of saturation of potassium is $\frac{5}{20} \times 100 = 25$ per cent. The remaining 75 per cent

of cations may be designated as complementary exchangeable ions.

To understand fully how the degree of saturation will affect plant nutrition and fertilizer applications, we must consider two separate aspects. These will be clearer if we restrict the discussion to hypothetical examples, assuming that all ions participating in the exchange have equal adsorption energies—that is, are held to the surface of the clay particles with equal forces.

Case I: Release of potassium when NH_4Cl is added in proportion to the base-exchange capacity of the soil. Suppose we consider the release of potassium upon addition of NH_4Cl in amounts equal to the base-exchange capacity. Calculations indicate that under such conditions the replacement of potassium is always 50 per cent of the amount of potassium on the clay, no matter what the degree of saturation. This relation is illustrated in figure 9 by a straight line parallel to the horizontal axis.

Case II: Release of potassium when NH_4Cl is added in proportion to the amount of exchangeable potassium in the soil. Suppose we select the specific case in which the amount of NH_4Cl is equal to the amount of exchangeable potassium. According to calculations as well as experiments, under these conditions the replacement of potassium strongly declines as the degree of saturation becomes less (curve II in fig. 9). In most soils the degree of saturation of potassium is very low—considerably under 10 per cent. In consequence, the liberation of adsorbed potassium upon addition of NH_4 fertilizers is apt to be of a small magnitude.

BEHAVIOR OF AMMONIA (NH₄OH, NH₃)

Unlike NH_4Cl and $(NH_4)_2SO_4$, which are strong electrolytes, NH_4OH is a weak base. An aqueous solution of ammonia contains NH_4 ions as well as NH_3 molecules. Since, moreover, the anion is OH, additional complications are introduced in the adsorption mechanism.

According to the exchange experiments reported in table 1, the amount of potassium released by NH_4OH is much smaller than in the corresponding studies with NH_4Cl or $(NH_4)_2SO_4$. Presumably the small exchange of potassium is due to the limited number of NH_4 ions in the hydroxide solution. In contrast to the NH_4 salt solutions, the amount of nitrogen taken up from NH_4OH greatly exceeds the amount of potassium replaced. The excess of nitrogen adsorbed over potassium released presumably results from NH_3 ad-



Fig. 9.—Release of adsorbed potassium by ammonium compounds as influenced by the degree of saturation of potassium.

sorption and from the reaction of NH_4OH with hydrogen ions of the clay. These two processes were studied with the aid of titration curves and by exposing clays to NH_3 vapor.

Titration Curves.—Colloidal clays extracted from soils of the Yolo and the Aiken series were converted into H-clays by electrodialysis. Suspensions containing 1 gram of clay in 100 cc of solution gave a pH reading (glass electrode) of 4.30 for Yolo colloid and 6.67 for Aiken colloid. To these acid systems were slowly added increasing amounts of NH_4OH . The pH values obtained as a function of NH_4OH added are illustrated by the titration curves of figure 10. In comparison with distilled water and washed quartz sand, the clay sys-

tems are strongly buffered against NH_*OH —a feature very pronounced in the Yolo colloid. The reaction taking place may be formulated as follows:

$$\begin{array}{|c|c|c|c|} \hline Clay & H + NH_4OH & \hline Clay & NH_4 + HOH \\ \hline \end{array}$$

Judging from the shape and position of the curves, H-Aiken colloid acts as a much weaker acidoid than H-Yolo colloid.



Fig. 10.—Titration curves of electrodialyzed colloidal clays with NH₄OH (1 per cent suspensions).

Under conditions of soil formation in California the colloids of the Yolo type are completely saturated with bases, and those of the Aiken type are nearly so. Naturally, therefore, the titration curves of the natural soils will differ markedly from those of their electrodialyzed colloids. As figure 11 shows, the addition of small amounts of NH_4OH to 5 per cent suspensions of natural soils increases the soil alkalinity to values exceeding pH = 10. In many soils (for example, Ramona and Yolo) the pH resulting from addition of NH_4OH is higher than the pH value obtained by adding NH_4OH to corresponding amounts of distilled water. This increase in pH may be explained

by the release of exchangeable calcium ions by NH_4 ions, resulting in the formation of $Ca(OH)_2$, a stronger base than NH_4OH .

The increase of alkalinity upon the addition of NH_4OH may temporarily upset the physiological balance of the plants growing in the soil. In most soils, however, NH_4 is readily nitrified, so that the reaction is shifted toward the neutral point.

Fixation of NH_4 as a Function of the pH of the Solution.—As may be seen in figure 11, the Aiken clay is strongly buffered at high pH values. Con-



Fig. 11.—Titration curves of natural soils with NH₄OH; milliequivalents per 100 grams of soil.

ceivably, then, the OH ions of the crystal lattice of the kaolinitic clays of the Aiken soils may participate in the reaction. To test this possibility, the base-exchange capacity of Aiken colloid and Yolo colloid for NH_4 was ascertained as a function of pH. Solutions of varying pH were prepared by adding NH_4OH or acetic acid to normal ammonium acetate. From 1 to 2 grams of the colloidal material was mixed with 100 cc of ammonium acetate solution. When the mixture had been digested on the water bath for 2 hours at 65° C, the samples were transferred to a Büchner funnel and leached with 1,000 cc of the same solution. After suction had been applied, the excess ammonium acetate was removed by leaching the clays with 50 cc of methyl alcohol. Dilute HCl was immediately added to the sample in order to prevent losses of NH_3

by evaporation. The amount of nitrogen fixed by the clay was determined by distillation.

In figure 12 appear the results for various colloids and soils. The striking difference between kaolinitic and montmorillonitic (bentonitic) material is noteworthy. Clays of the montmorillonite type such as bentonite and Yolo colloid adsorb nearly constant amounts of NH_4 , irrespective of the pH of



Fig. 12.—Adsorption capacity of various colloidal clays and soils for NH₄ as conditioned by the pH of the solution.

the solution. On the other hand, the kaolinitic materials such as kaolinite, halloysite, and Aiken clay fix much more nitrogen at high than at low pH values. This behavior is undoubtedly associated with the crystalline structure of the materials. Unlike the montmorillonitic, the kaolinitic clays contain sheets of OH ions as an essential part of the crystal lattice. With increasing pH, evidently, the exposed OH groups react progressively with the NH₄ ions of the solution; hence the adsorption capacity varies as a function of pH (5).

The question of the adsorption of NH_3 molecules was investigated by exposing dry H-clay to vapors containing variable proportions of NH_3 and H_2O molecules. The reactions to be expected may be shown by the following scheme :

Under the experimental conditions one would expect a molecular adsorption of NH_3 as well as a chemical reaction of NH_3 on the exposed OH ions of the clay lattice (4).⁹

The H-Yolo clays used have a base-exchange capacity of 64 milliequivalents per 100 grams as determined by the ammonium acetate method. Leaching of H Yolo with KCl and titration of the filtrate with NaOH give a capacity of 60 milliequivalents per 100 grams. H-Yolo clay samples were dried in the oven at 80° C and then placed in desiccators that contained NH_4OH solutions of various concentrations. After one week the samples were removed, and the



total nitrogen adsorbed was determined by distillation. The results are portrayed in figure 13 by the solid curve, which has the characteristics of an adsorption isotherm. In a parallel experiment the samples, after removal from the desiccators containing ammonia solutions, were transferred to aerators; and a current of dry, filtered air was passed through the apparatus for 4 days. Subsequently, the NH₃ remaining on the clay was determined by distillation. According to the dashed curve in figure 13, the nitrogen content of the aerated clay samples is independent of the NH₃ concentration in the vapor phase. The amount of NH₃ fixed corresponds closely, moreover, to the saturation capacity of the clay particles. Evidently we are dealing here with the reaction



The difference between the two curves in figure 13 is probably a measure of

⁹ These aspects have since been studied in considerable detail in the University of California Division of Soils laboratory.

the NH_3 molecules held loosely on the surface of the clay particles. These molecules may be readily removed by a current of air, whereas the NH_3 molecules combined with the hydrogen ions of the lattice adhere much more tightly to the surface. According to computations involving a comparison between the proportions of NH_3 and H_2O molecules in the vapor phase and on the clay particles, the NH_3 molecules at low partial pressures of ammonia have little chance to remain on the clay surface, because of competition with H_2O molecules. This probably explains why, in dilute clay suspensions, NH_3 molecules are not adsorbed by the clay in large amounts.

According to these experiments, NH_3 gas could be utilized for fertilizing soils in nonirrigated areas. A tank of NH_3 could be placed on a plow or disk, and the gas could be led through a nozzle into the soil, several inches below the surface.¹⁰ Fixation would immediately take place in accordance with the chemical equation given on page 444.

BIOLOGICAL EXPERIMENTS

Although the relation of nitrogen fertilization to crop yields is beyond the scope of this investigation, certain aspects pertaining to availability of adsorbed NH_4 and its nitrification by microörganisms lend themselves to laboratory tests.

Availability of Adsorbed NH_4 to Plants.—All data presented in this study indicate that nitrogen, in the form of $(NH_4)_2SO_4$ or NH_4OH , is held in the soil primarily as an NH_4 ion. In contrast to nitrate nitrogen, which is a mobile constituent of the soil solution, NH_4 is largely immobilized. Although plants readily adsorb nitrates from solution, there are no reliable data on the availability of NH_4 that is adsorbed on clay particles.

This problem was examined by adapting the culture-solution method of Hoagland and Broyer (6) to the requirements of colloidal-clay research. Barley plants 3 weeks old were decapitated, and the roots were placed in flasks containing the test solution. After 10 hours the roots were removed, washed, dried, and analyzed. According to Hoagland and Broyer, freshly excised barley roots are in an active state of metabolism and adsorb nutrients as readily as roots not severed from the shoots.

In each flask 100 grams (fresh weight) of excised roots were immersed in 3 liters of test solution. Two corresponding series of test solutions were used the first a series of $(NH_4)_2SO_4$ solutions, and the second a series of clay suspensions that contained NH_4 in the adsorbed form. The total nitrogen in the media varied from 0.75 to 27 milliequivalents. Besides varying amounts of NH_4 , all solutions contained a constant amount (3 milliequivalents) of potassium, either as K_2SO_4 or as K-clay.

Figure 14 shows graphically the intake of NH_4 , as computed from total nitrogen determinations of the roots before and after experimentation. The higher the amount of NH_4 in the nutrient medium, the more nitrogen the roots absorb. The differences between the two curves are probably not significant and are accounted for by root variability and experimental errors. There exists little doubt that in this experiment the roots accumulate nitrogen with equal ease from NH_4 Cl and NH_4 -clay.

¹⁰ A patent has since been obtained for this process (F. H. Leavitt, No. 2,285,932).

Throughout the experiment the roots also absorbed potassium. This intake was greatest at low concentrations of NH_4 and decreased as the amount of NH_4 in the solution and suspensions increased. In these experiments, therefore, large amounts of NH_4 tended to repress the accumulation of potassium by the roots. Probably the degree of saturation or complementary-ion principle outlined on page 440 was operating.

Studies on Nitrification.—Under normal field conditions NH_4 compounds in the soil are changed to nitrates by the activities of microörganisms. To



Fig. 14.—Uptake of NH₄ from NH₄-clay and NH₄Cl by excised plant roots. This graph demonstrates a high degree of availability of adsorbed NH₄.

obtain some information regarding the speed of nitrification and the individual behavior of NH_4OH and $(NH_4)_2SO_4$ during the process, laboratory experiments were conducted with Aiken clay loam and Yolo clay loam. The soils were ground and sieved (1-mm mesh), and 100-gram portions were placed in glass tumblers. Each soil was subjected to three series of treatments. To each sample was added, in the first series, 25 cc of distilled water; in the

second series, 25 cc of NH_4OH solution containing 5 mg of nitrogen; in the third series, 25 cc of $(NH_4)_2SO_4$ solution containing 5 mg of nitrogen.

The transformation of NH_4 into nitrate was measured weekly by the phenoldisulphonic acid method. The data plotted in figure 15 show the amounts of nitrate—expressed as nitrogen—that were found in the treated soils in excess of the amounts in the checks (soils receiving distilled water). Under the experimental conditions, 3 to 4 weeks were required to convert half the ammonia nitrogen into nitrate nitrogen. In the field, oxidation may proceed



Fig. 15.—Nitrification of NH₄OH and (NH₄)₂SO₄ added to Yolo clay and Aiken clay loam. The curves show the nitrate nitrogen in the treated soil in excess of the checks.

faster or slower than in this example, depending on moisture, aeration, temperature, and other conditions. No significant difference, apparently, exists between the behaviors of $\rm NH_4OH$ and $(\rm NH_4)_2SO_4$ at the low concentration employed.

Nitrification was also studied with NH_4 in the adsorbed state. An artificial soil was made up by mixing silica sand and Yolo clay containing adsorbed ammonium, potassium, calcium, and magnesium ions. To 50 grams of this material a small amount (0.5 gram) of fresh garden soil was added in order to provide the necessary microbiological population. The results resembled those shown in figure 15, except that the rate of nitrification was somewhat smaller at the beginning of the experiment than that observed with the natural soils. Evidently, adsorbed NH_4 may be readily oxidized to nitrates (1).

INFLUENCE OF NH₄ COMPOUNDS ON WATER PENETRATION

In irrigation agriculture, the rate at which water penetrates the soil is important. Accurate information on the effect of ammoniated irrigation water on soil permeability would therefore be interesting and significant. Such a study is also pertinent in view of the many contradictory claims advanced regarding the specific influence of NH_3 on water penetration.

Description of Method.—Soil samples were crushed, passed through a 1-mm sieve, and packed in metal tubes 2 inches in diameter and 8 inches in length. Successive 100-gram portions were placed in the tubes and subjected to ten



Fig. 16.—Entry of distilled water into soil columns, as influenced by soil type.

compacts with a standard compactor instrument. About four 100-gram portions were necessary to fill each tube. Metal screens placed at the bottom and top of the soil column provided support and stability and reduced possible disturbances of the surface of the soil. A 2-cm head of solution above the surface was maintained throughout the experiment. The volume of solution entering the soil column (infiltration) was calculated from the amount of liquid necessary to maintain constant head. The solution leaving the bottom of the column (percolation) was also measured frequently. All experiments were run in triplicate or quadruplicate in a room of nearly constant temperature. Over a dozen soils, comprising a wide variety of textures, were investigated.

Infiltration and Percolation of Distilled Water.—Curves in figure 16 depict the entry of distilled water into dry soil columns. The vertical axis indicates, in cc, the accumulative amount of water that has entered the soil column during a given period. The time is plotted on the horizontal axis. The

dominating influence of texture on water infiltration is an impressive feature of the graph. The time required for the dry soil column to take up 100 cc of water is 3 minutes for Oakley sand, 150 minutes for Vina silt loam, and 660 minutes (11 hours) for Altamont adobe clay. Data illustrating the exit of water are plotted in figure 17. Here the textural differences are even more outstanding. Generally speaking the heavier the texture of the soil—the higher its content in clay—the more slowly the water penetrates. This is, however,



Fig.17.—Percolation of distilled water in relation to soil texture.

a general rule rather than a strict relation, for the Aiken soils are outstanding exceptions. Aiken clay (not shown in graph) behaves much like Hanford fine sandy loam, and Aiken clay loam like Fresno fine sandy loam. In all probability the chemical nature of the Aiken clay (kaolinitic clay minerals) and the relative coarseness of the colloidal particles account for this behavior.

Infiltration and Percolation of NH_4 Solutions.—It would consume too much space to present the corresponding graphs for all the solutions of NH_4OH and $(NH_4)_2SO_4$ that were investigated. One example may suffice to bring out the pertinent relations. Figure 18 shows the entry of $(NH_4)_2SO_4$ and NH_4OH solutions of various strengths into a Yolo clay loam. As compared with distilled water alone, the presence of $(NH_4)_2SO_4$ markedly increases the rate of penetration. This beneficial behavior is the more pronounced at the higher concentrations of $(NH_4)_2SO_4$. Ammonia, on the other hand, does not produce such striking effects. The rate of entry of NH_4OH solutions is slightly less than that of distilled water. Very similar relations were obtained for the percolation rate.

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Figure 18, of course, applies to a specific soil only—namely, a representative of Yolo clay loam. Other soils behave differently. With Hanford fine sandy loam, all curves lie close together; this indicates that both NH_4OH and $(NH_4)_2SO_4$ exert a negligible influence on infiltration and percolation of water. The Aiken series suffers a marked reduction in permeability in the presence of NH_4OH .

These studies are summarized in table 2 and in figure 19. Table 2 shows (in cc) the volume of solution that percolates through a tube in 10 hours. For



into Yolo clay loam.

each soil the beginning of this 10-hour period was made to coincide with the emergence of the first drop of percolate at the bottom of the tube. According to figure 17 the percolation during the first day is very nearly a straight-line function of the time, so that the values in table 2 may be used to calculate hourly percolation rates. Table 2 also contains the percolation rates of the solutions relative to distilled water. In figure 19 the various soil types are arranged according to decreasing rates of NH₄OH percolation. As may be readily seen, the rate of percolation of the (NH₄)₂SO₄ solution exceeds that of distilled water in all soils except Aiken clay loam. This improvement in penetration is especially conspicuous for the heavier soils of the montmorillonitic clay group, a behavior that might be anticipated on theoretical grounds. As is well known, $(NH_4)_2SO_4$ has high flocculating powers and thereby increases the state of aggregation of the colloidal clay particles. In consequence

TABLE 2

VOLUME OF OUTGOING SOLUTION DURING A PERIOD OF 10 HOURS

Soil type	Distilled water	(NH4)2SO4 (15 milli- equivalents per liter)	NH4OH (15 milli- equivalents per liter)	Percolation relative to distilled water	
				(NH4)2SO4	NH4OH
	cc	cc	cc	ratio	ratio
Oakley sand	5,600	6,700	6.100	1.20	1.09
Goldridge fine sandy loam	684	716	596	1.05	0.87
Hanford fine sandy loam	606	680	606	1.12	1.00
Aiken clay	590	650	220	1.10	0.37
Yolo sandy loam	396	480	360	1.21	0.91
Aiken clay loam	150	145	133	0.97	0.89
Fresno fine sandy loam	140	190	150	1.36	1.07
Ramona loam	140	180	140	1.29	1.00
Yolo silt loam	75	125	80	1.67	1.07
Vina silt loam	68	88	78	1.29	1.15
Yolo clay loam	25	48	25	1.92	1.00
Altamont adobe clay	5	19	6	3.80	1.20



(NH4)2 SO4

Fig. 19.—Relative percolation of NH_4OH and $(NH_4)_2SO_4$ in comparison with distilled water. The bars that extend beyond 1.0 indicate soils in which the solution penetrated better than distilled water.

the pore space of the soil tends to become larger, and the passage of the water molecules easier. The results obtained with NH_4OH solutions present a more complicated picture. The soils on the upper portion of the vertical axis exhibit slight improvement in penetration when NH_4OH is applied, whereas the last four show a diminishing effect as contrasted with distilled water. The latter feature is most conspicuously manifested in Aiken clay.

In contrast to $(NH_4)_2SO_4$, which is a strongly flocculating agent, NH_4OH may also act as a dispersing medium. This latter quality is especially conspicuous in acid soils. Addition of NH₄OH to such soils neutralizes the acidity, as indicated on page 442. The resulting NH₄-clay possesses a higher zeta potential than the original H-clay, and dispersion of the clay particles takes place. The effective pore size tends to be reduced, and the percolation of water is slowed.

Ammoniated and Nonam	MONIATED	SOILS		
Qu'l turn	Soil treatments before measurement			
Son type	H ₂ O	(NH4)2SO4	NH4OH	
Yolo clay loam Vina silty loam Aiken clay	1.00 1.00 1.00	0.96 .67 .61	1.17 1.10 0.34	
Ramona loam Yolo silty clay loam	1.00 1.00	.37 0.31	1.29 1.04	

TABLE 3

RELATIVE PERCOLATION RATES OF DISTILLED WATER INTO

Water Penetration Subsequent to Ammonia Treatments.-Irrigating a soil that has been previously treated with NH₄ solutions will remove free electrolytes, and changes in rates of water percolation are likely to ensue. These were studied in conjunction with the percolation experiments already mentioned.

Soil columns were leached with $(NH_4)_3SO_4$ and NH_4OH solutions in concentrations and amounts used in irrigation practice. Corresponding sets of columns were leached with distilled water only. After treatment all soils were allowed to drain for 3 days. Infiltration and percolation rates of distilled water were then measured over a period of 200 hours. During the first few hours the infiltration rates closely corresponded to those of the NH_4 solutions; but as leaching continued, marked changes in rates were observed.

Table 3 summarizes the findings. It contains the percolation rates of distilled water into soils previously leached with NH₄ solution or with distilled water. Surprising is the fact that in all soils, except Aiken clay, the $(NH_4)_2SO_4$ treated soil has the poorest water penetration, whereas before leaching it had the best. (Compare fig. 19.) This suggests that removal of free electrolyte may produce significant dispersion of the $(NH_4)_2SO_4$ -treated soil.

The time interval of 3 days between the end of the NH_4 treatment and the first measurements of water percolation was too short to permit significant nitrification. To ascertain the effect of nitrification on permeability, 700-gram lots of Yolo clay loam and Aiken clay loam were placed in shallow pans and moistened. A little water extract of a fertile garden soil was added to insure the presence of nitrifying bacteria. Two thirds of the pans each received 7.5

milliequivalents of $(NH_4)_2SO_4$ or NH_4OH . All soils were kept moist for 4 weeks and were then dried and screened through a 2-mm sieve. Soil columns 2 inches thick were prepared, and the rates of infiltration and percolation of distilled water were determined in the usual manner. Only traces of nitrate nitrogen were found in the leachate, although the ammonia nitrogen content of the soil was reduced by 50 per cent. Presumably this chemical had been consumed by microörganisms.

In the Aiken clay loam the rates of entry and percolation for 5 to 60 hours were highest for $(NH_4)_2SO_4$ - and lowest for NH_4OH -treated soil. The sample not receiving nitrogen occupied an intermediate position. In the Yolo clay loam a similar sequence was obtained for the initial infiltration period, but after 10 to 40 hours all NH_4 -treated soils had only about one half the percolation velocity of the untreated soil.

To summarize, these experiments indicate that soils fertilized with ammonium compounds may eventually suffer a reduction in rates of water percolation. Many field observations on heavy applications of ammonium sulfate support these laboratory findings.

GENERAL DISCUSSION

Considerable controversy exists regarding the comparative merits of Agricultural Ammonia and $(NH_4)_2SO_4$ as a nitrogen fertilizer. The farmer usually asks, "Which of the two fertilizers is better?" His inquiry can be answered only by long-time experiments involving crop yields and economic considerations; the problem lies beyond the scope of the present investigation. Several controversial questions relating to the indirect effects on crop production may, however, well be discussed in the light of these studies (8).

Promoters of Agricultural Ammonia emphasize that their product does not contain anions, such as sulfate or chloride, which might remain in the soil solution as a residue. Theoretically, continued use of large amounts of $(NH_4)_2SO_4$ might ultimately increase the sulfate and sulfuric acid concentration of the soil solution to such an extent as to injure plant growth. In practice, however, such conditions are not very common. Winter rains and occasional heavy irrigations will displace the concentrated soil solutions to greater depth, beyond the reach of most roots. Since, moreover, soils usually contain some exchangeable calcium, the $(NH_4)_2SO_4$ will be converted into NH_4 -clay and $CaSO_4$. The latter salt is sparingly soluble, yielding (at saturation) a sulfate concentration of 30 to 35 milliequivalents per liter, which is not known to be toxic to common crops.

Ammonia dissolved in water becomes NH_4OH . It is a weak base and therefore slightly alkaline, having a theoretical pH value of about 10.8 at a concentration of 10 milliequivalents of NH_4OH per liter (170 p. p. m. NH_3). Under field conditions, because of the pressure of CO_2 , the reaction is less alkaline. In soils containing exchangeable calcium the addition of NH_4OH will produce NH_4 -clay and $Ca(OH)_2$, resulting in a further increase in pH. Locally, unfavorable effects may occur—for example, the precipitation of phosphates and of micronutrients such as zinc. As soon as the ammonia is nitrified, the alkaline reaction will disappear; but not necessarily all the precipitates, since these may be irreversible. According to some claims, ammonia liberates from the soil substantial amounts of plant foods, especially potash and phosphate. As far as exchangeable potassium is concerned, this claim is well substantiated by the present study. However, $(NH_4)_2SO_4$ is more effective in this respect than Agricultural Ammonia. Since potassium held on the clay in exchangeable form is readily taken up by roots, the release of exchangeable potassium by NH_4 is probably less important in plant nutrition than has hitherto been assumed. According to some growers, NH_4 liberates not only exchangeable potassium, but also a "nonexchangeable" kind, which presumably is locked up in the interior of mineral particles. Such an assertion is not easy to verify.

Ammonia, owing to its alkaline reaction, may possibly release phosphate from those lateritic soils (for example, the Aiken series) in which PO₄ occurs as phosphated kaolinitic clays, or as iron and aluminum phosphate. In many California soils, on the other hand, phosphate exists as calcium phosphate, whose solubility becomes less as the alkalinity becomes greater. In these soils NH₃ would, temporarily at least, adversely affect the availability of phosphate. In contrast to NH₄OH, the reaction of $(NH_4)_2SO_4$ is slightly acid (pH = 5.7 for a concentration of 10 milliequivalents per liter); and its effect on phosphate availability should therefore be opposite to that of ammonia. One would expect $(NH_4)_2SO_4$ to lower the availability of phosphate in lateritic soils and to increase it in soils containing calcium phosphate.

"Wherever the water goes the ammonia goes" is another claim frequently advertised. The present study conclusively shows, however, that water and ammonia may move independently of each other. Yet the counterstatement that NH_3 is always fixed in the first 2 or 3 inches of soil is likewise erroneous. Actually, the depth of NH_3 penetration is conditioned to a high degree by soil texture. In coarse-textured soils (sands) NH_3 , in the amounts used in irrigation agriculture, distributes itself rather uniformly through a depth exceeding 6 inches, whereas in fine-textured soils (clay loam and clays) practically all the NH_3 is fixed in the top 2 inches. With respect to penetration, NH_3 and $(NH_4)_2SO_4$ behave very much alike, though in acid soils $(NH_4)_2SO_4$ penetrates deeper than NH_4OH .

The soil structure must also be considered. Fine-textured soils of the adobe type form wide and deep cracks upon drying. Conceivably, irrigation water that fills these cracks and crevices may carry some of the dissolved NH_3 to considerable depth.

"Ammonia has a very marked effect in causing soil colloids to coagulate so that heavy soils which allow only little penetration of water gradually become lighter and allow better penetration." Again, this statement is true in part. First of all, the coagulating effect of hydroxides is manifested primarily in base-saturated soils—that is, soils having neutral or alkaline reaction. Second, in acid soils monovalent hydroxides tend to have a dispersing or deflocculating effect. The data obtained in this study fully confirm both the coagulating and the dispersing effect for ammonia. NH_4OH increases water penetration in some soils, decreases it in others, as compared with distilled water. In all tests performed, $(NH_4)_2SO_4$ proved to be a superior coagulator. Fortunately, owing to the ultimate utilization of nitrogen by plants, the reduction in penetration observed in some soils need not be cumulative and permanent.

Generally speaking, in their principal behavior in the soil, NH_3 and $(NH_4)_2SO_4$ are similar. Each has certain advantages and disadvantages; but as far as the soil is concerned, apparently neither is more beneficial than the other.

SUMMARY

Laboratory studies on the behavior of NH_3 (or NH_4OH in water) and $(NH_4)_2SO_4$ in soils revealed the following facts:

1. The uptake of nitrogen in soil suspensions containing NH_3 and $(NH_4)_2SO_4$ is, broadly speaking, a function of soil texture. Fine-textured soils adsorb more nitrogen than coarse-textured ones. Numerous exceptions, however, exist, being conditioned by soil acidity and by other factors such as the nature of the clay minerals.

2. Under comparable conditions, acid soils tend to adsorb more nitrogen from NH_4OH than from $(NH_4)_2SO_4$. Alkaline soils, as a rule, adsorb more nitrogen from $(NH_4)_2SO_4$ than from NH_4OH .

3. Determinations of the depth to which NH_4 compounds penetrate in artificially prepared soil columns for a 10-inch irrigation containing 127 p.p.m. of nitrogen indicate that (a) in Oakley sand, NH_4OH and $(NH_4)_2SO_4$ readily penetrate below a depth of 6 inches; (b) in Yolo sandy loam, penetration is restricted to a depth of 4 inches; (c) in Yolo clay loam, all nitrogen is held in the first 2 inches; and (d) in Aiken clay loam, $(NH_4)_2SO_4$ penetrates to greater depth (4 inches) than NH_4OH (2 inches).

4. The process of nitrogen adsorption is governed by base-exchange reactions. With $(NH_4)_2SO_4$ and NH_4Cl the exchange is equivalent. For every NH_4 ion adsorbed, a corresponding amount of cation is released. In NH_4OH the amount of NH_4 adsorbed greatly exceeds the number of bases liberated. Ammonia may react with hydroxyl ions of the clay lattice.

5. Ammonium ions readily displace exchangeable potassium ions from clay particles. The extent of this reaction is greatly influenced, however, by the degree of potassium saturation of the clay. For most soils the release of potassium by NH_4 fertilizers is probably of minor significance.

6. Dry clays readily adsorb NH_3 from vapors containing NH_3 molecules. These molecules are held loosely and may be easily removed by aeration.

7. Judging from experiments with excised plant roots, NH_4 adsorbed on clays can be readily utilized. Likewise nitrification of adsorbed NH_4 is easily accomplished.

8. The influence of $\rm NH_4$ on water penetration in artificially prepared soil columns varies so greatly among different soils that no generalizations appear possible. The following trends, however, were observed : In 11 out of 12 soils, ranging in texture from sands to clays, the percolation rates of $(\rm NH_4)_2\rm SO_4$ solutions were higher than those of distilled water. The influence of $\rm NH_4OH$ solutions was much more variable. In 5 soils $\rm NH_4OH$ improved water percolation as compared with distilled water; in 3 soils $\rm NH_4OH$ had no effect; and in 4 soils it lowered the percolation rates. This decrease was most pronounced in Aiken clay, an acid soil containing kaolinitic clay minerals.

9. In soils that have been treated with NH_4 compounds, the subsequent penetration rate of distilled water may suffer a reduction, especially in the case of $(NH_4)_2SO_4$.

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