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UNIVERSITY OF ALBERTA

THE FATE OF ADDED PHOSPHORUS IN CHERNOZEMIC AND LUVISOLIC SOILS.

BY

THOMAS MESSIER

A THESIS

SUBMITTED TO THE FACULTY OF GRADUATE STUDIES AND RESEARCH IN PARTIAL FULFILLMENT OF THE REQUIREMENTS FOR THE DEGREE OF

IN SOIL FERTILITY

DEPARTMENT OF SOIL SCIENCE

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The undersigned certify that they have read, and recommend to the Faculty of Graduate Studies and Research for acceptance, a thesis entitled THE FATE OF ADDED PHOSPHORUS IN CHERNOZEMIC AND LUVISOLIC SOILS submitted by TOM MESSIER in partial fulfillment of the requirements for the degree of MASTER OF SCIENCE in SOIL FERTILITY.

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DEDICATION

To my mom and dad, Mary-Rose and Raymond Messier, in appreciation of the constant love and support that you have given me.

To my dear wife Helen for your love and commitment.

Abstract

Low efficiency of P fertilizer can lead to excess input costs for the producer and to depletion of the Earth's supply of P are deposits. This study was conducted to determine the face of P fertilizer within the fertilizer zone in the sesson of application. A greenhouse experiment was conducted on samples from the Ap horizons of Breton (Othic Gray Luvisol) and Paddockwood (Dark Gray Chernozem) soils to determine the efficiency of ³²P uptake by barley and the distance of P movement from the fertilizer pollet sits. The uptake of ³²P by barley was not affected by the granule size of monoammonium phosphate (MAP) fertilizer; however, when the granule of MAP fertilizer was placed inside a volume of past or straw the uptake efficiency of ³²P by barley increased. Also, the efficiency of P uptake was greatly affected by the position of the band of MAP fertilizer. Deep bands 10 cm deep had significantly lower uptake of ³²P then bends located 2cm beside and 2cm below the seed row. The soil around the 0.10g granule of MAP fertilizer showed an increase in P to a distance of 1.5cm.

A follow up experiment focused on the zone of soil that interacts with MAP fertilizer. Five mg P/g soil was added to the two soils and equilibrated for 2 months; the soil solution was analysed and the solid P fraction underwent sequential extraction. Phosphorus mineral solubility diagrams indicate that brushite (DCPD), actacalcium phosphota and tricalcium phosphota were the minerals controlling H_2PO_4 ° activity in soil solution. The sequential extraction showed that P added as MAP fertilizer moved from readily available to less available soil P fractions with time. However, most of the added P remains in the readily available Resin-P fraction.

In the soil environment affected by MAP fertiliser added to Broton and Paddeckwood samples, the Ca phechate minerals central the H_2PQ_4 ° anti-thy and the added P remains readily available.

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1.0 Introduction

Phosphorus (P) is an essential element required for plant growth. It is classified, along with nitrogen, potassium, calcium, magnesium and aulfur, as a mecronutient. Macronutrients are required for plant growth. Therefore, without an adequate supply of phosphorus, plant growth and crop production will be limited.

Phosphorus is most readily absorbed from soil solution as orthophosphete. The form of orthophosphete in solution is pH dependent (Figure 1.1). The highest rate of orthophosphete absorption into the plant occurs at pH 5 to 7 and therefore $\rm H_2PO_4^{-2}$ is the most available form at P for the plant.

in addition to inorganic orthophosphate ions, plants may also absorb certain soluble organic phosphates. Nucleic acids were found to be as effective as monopotassium phosphate as a P source for barley growth on phospharus deficient soils. However, the instability of the soluble organic phosphates in sell limits their importance as sources of phospharus for higher plants under field conditions.

The concentration of P in plants varies from 0.1 to 0.4%. So a crop of 1000 lights would contain 1 to 4 kg P/hs. This value is considerably lower than the concentrations of carbon, hydrogen, enygen, nitrogen and peterelium in plants. Dispite its low concentration in plants, P has a vital role in plant growth.

The most important role of P in plant growth is energy storage and transfer. Phosphete compounds store energy that has been derived from photosynthesis and the metabolism of carbonases. This energy is stored in adenasine di- and triphosphetes (ADP and ATP), as high energy bonds between phosphete melecules. Almost every metabolic reaction in plants proceeds with energy derived from those phosphete derivatives.

in addition to this metabolic role, phosphorus is an important structural component of a wide variety of biochemicals. These biochemicals include nucleic solds, coonsymes, nucleotides, phosphoproteins, phospholipids, and sugar phosphotes. The reproductive primards of a plant requires adequate phosphote early on in development. P is assential for

FIGURE 1.1 - Removed because of copyright restriction.

SOLUTION PH

Figure 1.1. The effect of solution pH on ionic forms of descrived phosphorus (Barber, 1984).

seed formation as it is found in large quantities in seeds and fruit. P is also involved in the structural framework of the protoplesm and thus occurs as part of the chloroplest structure.

Since P is important for the growth of plants, if plants are unable to attain adequate P from the soil, plant growth will be restricted. Without adequate P a plant may suffer from insufficient root growth and seed development. Resistance to diseases such as root rot will decrease. The plant will also be prone to lodging due to low straw strength.

The A horizons of Western Canadian soils contain significant amount of P (500 to 900 up P/g soil). Dispite this high level of total P in many soils, available P tends to be low. This may be because P reacts with soil components to form relatively insoluble compounds, many of which have limited availability to plants. The major factor affecting the supply of phosphate to plants is the relationship between the amount of soil P (quantity) and the concentration of soil solution P (intensity). The quantity of total soil P (Pt) is much greater, in most soils, then the more insertent available P.

Fifty to one hundred years of cultivation under wheat-fallow systems have decreased the organic matter content of Western Canadian soils substantially. Mineralization of P during cultivation has increased its rate of removal from sell by crops. However, as the amount of organic matter decreases, the contribution that mineralization has for increasing the systems P pool decreases. Therefore, with cultivation, increasing amounts of fertilizer additions are required to maintain crops production.

The addition of phosphate fortillaers to sells increase the plant available P content of the sell. Sells that are low in plant available P respond the best to fortillaer P additions. Upon addition to the sell, phosphate fortillaer is quietly integrated into the sell P cycle. Therefore, sells that have consistently received P fortillaers have experienced a buildup of sell P; with a relative increase in plant available P.

Understanding sell P and its response to fartillier addition requires an understanding of the sell P cycle. P is present in this cycle as inergenic (PI) and organic (Po) forms. The surface horizons of Western Canadian sells centain 25 to 55% Po. This Po is found in

humus and other organic materials. The Pi content of these soils is frequently higher than that of Po. Pi occurs in combinations with elements and clay minerals. Pi forms sparingly soluble to very insoluble complexes with iron, sluminum, and calcium. And the clay phosphate complexes are generally insoluble. The major processes of the P cycle are adsorption onto inorganic minerals, uptake by plants and microbial turnover through immobilization and mineralization. This transformation of soil P forms occurs under the belance of steady state conditions.

In this thesis the natural processes of the P cycle were studied to develop an understanding of the fate of fertilizer P once added to the soil. By studying both soil solution and solid phase characteristics resulting from the reactions of soils with P fertilizers, the nature of the fate of added P could be determined.

2. LITERATURE REVIEW

2.1 Phosphorus cycle in the soil

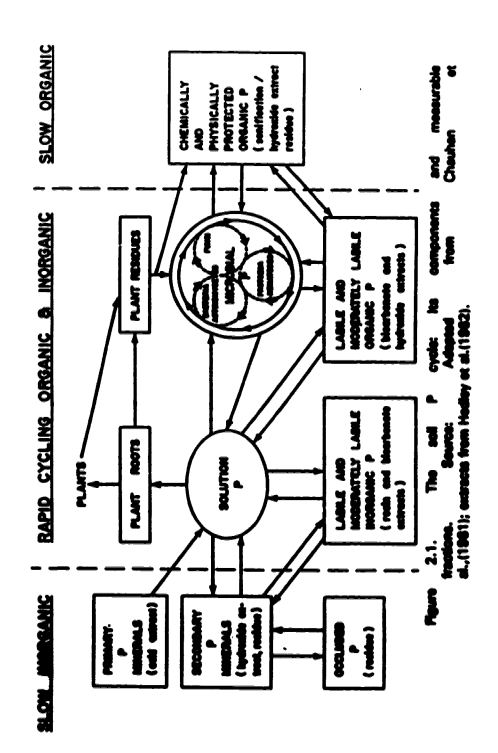
2.1.1. Introduction

Phosphorus (P) occurs in soil in many different forms. These different forms of soil P are components of the soil P cycle; depictions of the soil P cycle such as Figure 2.1 simply model the transformations of soil P between these different forms. In order to better understand the soil P cycle, the forms of soil P will be discussed along with possible transformation mechanisms.

2.1.2. Solid phone P

2.1.2.1. Ornenic P

The surface horizons of notive Albertan soils contain 80-130 μ g Po g⁻¹ soil, while cultivated Albertan soils contain 70-80 μ g Po g⁻¹ soil (McKensie, 1986). Only 90 - 70% of the Po forms have been identified (McGIII and Cala, 1981; Stewart and McKersher, 1982). Featers complicating Po



identification include high chemical complexity of soil Po, strong suseptibility of some Po compounds to hydrolysis during extraction, strong scription of Po by clays and insoluble solt formation with metal cations such as AI and Fo. Most Po has been identified as phosphate esters (Anderson, 1980) with a significant portion of polyphosphate compounds containing P-C, P-N and P-S linkages (McGill and Cole, 1981). The remaining Po probably occurs as insoluble complexes with clay and organic matter (Tate, 1985). Microbial P (Pm) represents a small but important table.

pool of soil P (Jenkinson and Ladd, 1961) that is responsible for the redistribution of P through mineralization and immobilization.

make up most of the Pe in sell (Black, 1988). Since the plant does not affect phosphate, this ion enters into organic combination largely unaltered. Thus the P in IP, phosphate/and nucleic acids is found as phosphate (Alexander, 1977). IP comprising up to 60% of total Pe are homocyclic compounds which can form a series of phosphate esters ranging from monophosphates to homophosphates. Penta-(IPS) and homophosphates (IPS) such as physic acid are the predominant ester forms of this group found in sells (Ficietzed and McKercher, 1975). These higher inceltal phosphates (IPS and IPS) are more prevalent than the lower ester forms in more weathered sells (Helessed and Anderson, 1970).

Insolated allowing them to accumulate to 25% of Pt. They are stabilised by reacting with Fe and AI under sold conditions and with Ca under alkaline conditions to form very insoluble solts. Sall IP contains are found to be significantly correlated (r=0.91) with sold encions actuable iron but not with organic corbon (Alexander et al, 1974). These complexes are more resistant to enzyme attack than more ashable ester solts. Clay minorale and finally divided acceptantions will evengly adopt homophosphate (Alexander et al, 1974). Phytose annumes cause IP breakdown through dephasphorylation by attacking a phosphate group adjacent to

the free hydroxyl provided that it is not sterically attached as an axial group (Irving and Coagrove, 1971). The activity of phytose is reduced by IP retention by Fe and Al complexes in acid conditions and retention by clays in neutral conditions (Alexander, 1977).

RNA and DNA occur in all fiving organisms and consist of a chain of sugar units, joined by phosphate ester bridges. A nitrogen base is then attached to each sugar molecule. The evidence for the existence of nucleic acids in soil is indirect. The presence of nucleic acid bases in a bound form in humic acid fractions, and the proportion in which they occur are indicative of DNA derived polynucleotides of microbial origin (Halateed and McKercher, 1975). Upon residue decomposition these nucleic acids are quickly released into soil, accounting for 5 - 10% of Po, and subsequently broken down. Mineralization proceeds by an initial depolymenization of nucleic acids and then subsequent cleavage of the phosphate from the products generated by the depolymentaling enzymes (Alexander, 1977).

Phospholipids are esters of fatty acids containing P and comprise 1% of Po. The concentrations of fipid P were highly correlated with Pt and soil organic carbon in a study in Saskatchewan Blaisteed and McKercher, 1975). It has been shown that actinomycetes are the major organisms responsible for decomposition of phospholipids in soil organic matter. Phosphate is cleaved from the organic compound, and it is then assimilated by responsible populations (Alexander, 1977). However, phospholipids may also be adsorbed to soil constituents (McGIII and Cola, 1981). The importance of this group of P compounds in supplying P to plants cannot be everloched since their synthesis and degradation may be fairly repld in the soil system.

Other exters that originate from the breakdown of bectarial cell walls tend to be stable in sells. In addition, other as yet unidentified components may be formed by microbial action (Alexander, 1977).

Studies of Po minorelization in sell are not numerous because of the uncertainty of identifying the origin of Fi, since both Po and Fi forms occur simultaneously. Easily actuals Po can be converted to Fi relatively quietly by phosphatoco enzymos (Colo et al., 1977).

These enzymes are released into the soil by microbes and plant roots (Tarafder and Jungit, 1987; Tarafder and Classon, 1988). The observation that Po is more variable in soil organic matter than are N and S (Delel, 1977) suggests that Po may be stabilized by mechanisms other than those stabilizing organic matter in general. The biological activity that regulates the simultaneous mineralization and immobilization of P is affected by available C, available nutrients, temperature, water content and soil chemical factors such as pH.

Carbon from growing roots and dead plants is a major source of energy for microbes. The belance between mineralization and immebilisation of P during organic matter breakdown depends on the C:P ratio of the substrate being utilized by the microorganisms (White, 1981). Cheuhan et al (1979,1981) determined that the rate of residue decomposition affects the rate of P immobilization according to the energy released and the C:P ratio of the residue.

The amount and type of carbon source in the sell affects the rate of P immobilization. Collulose produces a lower amount of energy when decomposing and therefore will decompose at a slower rate then simple sugars. However, collulose decomposition has a higher capacity to immobilize P (Singh et al, 1988); because fungel flora, which are the more active organisms in collulose degradation (Alexander, 1977), assimilate 15 - 20% more carbon than bectaris and actinomycoses and therefore use more N and P. Nevertheless, slower decomposition of collulose means a slower rate of P immobilization thus leaving larger amounts of available P during initial growth when plants need very high amounts for their proper growth (Singh et al, 1988). Readily enidicable carbonaceous compounds proved harmful to the growing plants because they rapidly immobilize very high amounts of available P during initial stages of growth. Callulose and crude fiber, both easily available to farmers in the form of crop wester, initiate a slow rate of P immobilization but the process continues for a long period of time.

When fortilleer P is added with estitions, 58 - 89% of the added P was found in Fi terms (Chauten et al, 1981). The addition of P fortilleer without a carbon substrate

followed by incubation does not significantly after the Po content of soil and the added P remains in inorganic forms. Readily plant available P was greatly increased by addition of fertilizer P. But when carbon was added with the fertilizer P, the increase in table P indices were as great as when P fertilizer was added alone (Chauhan et al., 1981).

Mechanisms involved in the interaction of the major nutrients are very complex, making predictions of P transformations based on such interactions difficult (McGill and Cole, 1981). The Po level is known to be directly related to the concentration of other humas constituents. The P content of humas is 0.3 to 1.0% and 5 to 20% of the C and N concentration, respectively (Alexander, 1977). However, Spiers and McGill (1979) detected an increase in phosphatase activity with the addition of N, S and C but not with the addition of P.

The plant availability of P in crap recidue is governed by warmer temperatures and adequate available moisture levels (Abbott, 1978). Seasonal fluctuations of Po have been observed in cultivated sells (Dormaer, 1972), where sell Po increases over winter and decreases in apring. Campbell et al (1994) observed several significant flucuations of Pi during the growing season, which are correlated to air temperature and sell moisture.

Soil organisms function over a wide range of water contents (Wilson and Griffin, 1975), therefore decomposition of organic residues occurs over a similarly wide range of water contents. Campbell (1973, 1976) found peaks in microbial activity after rainfall, indicating that westing and drying affects both mineralization and immobilization.

Po is less stable in neutral or alkaline soils than in acid soils because microbial diversity decreases with decreasing pH (Tate, 1986). Mineralization of Po increased with rises in sell pH but mineralization of organic cerbon and total N does not. These same pH effects on mineralization in the field are indicated by the fact that ratios of total organic cerbon and total N to total Po increase with pH in the same sell servates.

P occurs in sells in a number of inergenic and organic forms. The complex

transformations of these forms are affected by a combination of physical, chemical and biological reactions. C and N levels help the microbial population to redistribute P through mineralization and immobilization.

The P of this organic pool in virgin soils is derived from primary P minerals. For a better understanding of the P cycle the forms and solubilities of these minerals should be examined.

2.1.2.2. Phosphate minerals

P is pentavalent in nearly all naturally occurring phosphases. The crystal structures of P minerals have the P atom surrounded by four atoms forming an approximate tetrahedral arrangement. The formation of stable atomic structures containing PO₄ tetrahedra is accomplished through the high affinity of PO₄ for cations exhibiting 8-fold coordination. This results in mineral species that can be classified by a system similar to that used for allicate minerals - ie) framework, chain, layer phosphates. The most common naturally occurring phosphate minerals are the Berlinite, Variecite, Apathe and Crandallite groups.

The Berlinite group is homostructural with quartz. This framework group has PO₄ and AIO_A tetrahedra linking and alternating along verticles.

The Variacite group has a framework structure and is widely distributed in nature. Variacite and strengths are end members of this isomorphous series, the composition of which varies from AI(PO₄)01₂O1₂ to Fe(PO₄)01₂O1₂, respectively (Lindsey and Visk, 1977).

The Apades group has an insular structure consisting of columns of Ca and O atoms parallel to the unique axis forming trigonal priems. A detailed description of the structure of apades is given by Key et al (1984). From seructural and compositional studies of natural materials it has been found that many substitutions are possible in the general formula (Ca₂Ca₃(FO₄)₃(OH,F)) of species. Only OH and F show partiest isomorphism in the anionic substitution of Ca(II) trigonal priems resulting in the end members of the isomorphism series. An extensive listing of species subgroups is given by Kraidler (1987).

Plorepodie is the accessory mineral of ignoous racks and is the primary enriched

phase of carbonattes. Francoite is the primary marine phosphase in sedimentary rock. Delhite makes up the mineral metter of fossil bone; and hydroxyapatite is found in guano-altered limestone (Lindsay and Viek, 1977).

The Crandellite group has insular structure with P occupying a tetrahedra position and Fe or AI an octahedral position. The third position is filled from a group of a large number of cations. Gorceixite and crandellite are common forms of soil phosphate as they are present in the clay fraction of a veriety of soils (Lindsay and Viek, 1977).

The persistence of spetite in soils depends on the intensity of weathering by geochemical and biological processes. In neutral to alkaline environments of high Ca concentration, spetite is stable and weathering is probably limited to the somewhat acid environment adjacent to plant roots. In more humid regions, where acidity is a consequence of soil formation and development, spetites dissolve rapidly, releasing their phosphates into soil solution (Tiessen and Stewert, 1983). Most of the P enters Fe and Al minerals by substituting for allicate. However, Lajtha and Schleeinger (1988) found in their study of a chronosequence a decrease in Ca-P but did not find a corresponding increase in Al-P or Fe-P with increasing soil age. In soil, P ions can also react with solution Al or Fe to form stable secondary minerals. These minerals are comparatively stable under acid conditions. The low solubility of these minerals not only limits the leaching loss of P but also its availability to plants.

Insoluble inorganic phosphases are known to be solubilized by different genera of bacteria, fungi and actinomycoses (Hayman, 1976). Phosphase solubilizing microbes have been used extensively in Eastern Europe to increase availability of native sell P (Michaelin and Naumova, 1982). The important machanisms involved in the dissolution of insoluble phosphases in sell include formation of organic solds and choisting substances. The increase in microbial activity as a result of composting produces these organic solds and choisting substances and has been shown to increase the calubilization of reck phosphase (Mishre and Bangen, 1986). The rate of P mineral dissolution is influenced by the

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conceneration of calcium and P in the soil solution (MacKay et al. 1986).

The use of phosphate solubility diagrams has become well established for describing P mineral solubility in the soil environment (Aslying, 1954; Lindsay et al, 1959; Lindsay and Moreno, 1960; Rai and Lindsay, 1975). Solubility diagrams are constructed by including ion activities in a minerals' equilibium reaction. The isolines are plotted as the relationship between pH₂PO₄ and pH.

Solubility isolines are particularly useful for determining relative stabilities of phosphete compounds in soils at various pH values. A superssturated solution indicates precipitation and an underesturated solution indicates dissolution. These solubility isolines have been quite useful in indicating the solid phase calcium phosphetes that control soluble P levels.

However, the thermodynamically generated chemical transformations characterized by under- or supersaturated conditions take place very slowly. Soil scientists have tended to think of crystalline phases in soil as pure stoichiometric crystals, when in fact such purity is difficult to obtain in an environment as heterogeneous as soil. Isomorphous substitution is a wide occurrence in soil minerals whether of geologic or pedogenic origin. Solubility product isotherms represent and members in an isomorphic substitution series of minerals. For the purpose of assessing soil P, solubility product isolines can be excellent tools in determining the kind of P compound that influences P in the soil solution and possibly delineating the path by which P is supplied to a growing plant. However, solubility product leotherms cannot measure the quantity of such compounds, and this is a serious limitation.

Pertilizers can be supplied to the sell in mineral form; largely as monoammentum phosphate (MAP) and as monocolclum phosphate (MCP). As dissolution of water-actuate fertilizers precede in sells, conditions often fover in situ precipitation of P compounds. The best documented examples of such reaction products are the Brushite (CaHPO $_4$ '2H $_2$ O) recidues remaining at application situs of MCP fertilizers. Laivr at al (1960) placed MCP granules in five sells at two mainture contents and found that from 20 - 34% of the applied

P remained as brushles. With time these reaction products of brushles change to octacalcium phosphete (OCP) (Lehr and Brown, 1958; Bell and Black, 1970b). The soil property most widely used to predict the ultimate resection products is soil pH, recognizing that pH is itself a reflection of degree of weathering, minerals present, exchangeable cations, and a multitude of other factors.

The solubilities of P fertilizer reaction products have also been described by solubility diagrams. These crystalline minerals are generally much more soluble then the primary P minerals. The more soluble reaction products are more likely to be a source of P for plants then selectively adsorbed P. Since labile phosphete and crystalline phosphete coexist in equilibrium: Labile $P \rightarrow$ Solution $P \rightarrow$ Crystalline P.

2.1.2.3. Adequated P

P sorption occurs on the surfaces of soil constituents of either constant charge or variable charge. Iron and aluminum oxides, amorphous clay minerals and organic matter all have surfaces that allow H + and OH to determine the surface charge. Whereas, crystalline clay minerals have largely a permanent negative charge through isomorphous substitution. These secondary clay minerals interact through cations held tightly to pleasilite surfaces.

Hydrexyexides of Fe and AI are amphotoric, having either negative, neutral or positive charge depending upon pH. As the pH in the soil decreases, the edge hydroxyl groups become HOH groups. By calculating the number of hydroxyls released through exchange, Rejan et al (1974) determined the adsorption of HPO₄ by hydrous AI enides to be due to a binuclear coordination of phosphate ions with two AI stems. Therefore it is well established that figured exchange is the most likely mechanism of the phosphate - surface hydroxyl reaction (Perfitt et al, 1975; Goldberg and Species, 1986). The result of this exchange reaction is the formation of an inner-ophere surface complex (Species, 1981b) between an e-phosphate group and a motal cation which is bound to the hydroxyl group.

Barrow (1986) expanded earlier models of adsorption ento variable charged surfaces

with a four-lever model. The innermost lever contains adsorbed H * and CH ions. These are responsible for the charge development. The next lever contains adsorbed ions. The ions of this second lever differ not only in their affinity for the surface but in their mean position when adsorbed. The closer the anion adsorption plane is to the surface plane the greater the effect of decreasing anion adsorption with increasing pH. Thus, in this model the location of the adsorption plane determines the steepness of the effect of pH and also the magnitude of the charge conveyed to the surface. The third layer consists of electrolyte ions (es: Ne *, Cl*) bound to surface groups. The fourth lever is the diffuse layer.

Hemsites may absorb widely different amounts of phosphate according, not only to their specific surface but to their shape; because shape is related to the relative extent of the different adsorbing and non-adsorbing faces (Barron et al. 1988). A theoretical mean adsorption value of phosphate on gosthites is about 2.5 umol P m⁻² (Goldberg and Sposito, 1984b), while hemsites generally sorb two and a half times less phosphate (Schwertmann, 1987).

Pleastides et al (1900) determined that clays saturated with divelent ions adsorbed more P than clays asturated with monovelent ions. This indicates the thickness of the diffuse double layer may be involved in controlling how much P is adsorbed at the surface of the clay. Hingston et al (1972) postulated that the edge faces of kaclinite behave similarly to the hydrous exides in anion adsorption. The adsorption of low phosphate concentration onto kaclinite is a result of the replacement of specifically adsorbed water molecules at the broken edge of the clay lattice. Kue and Lates (1972) believe that phosphate replaced a water molecule rather than a hydroxyl group, since the pH of the solution did not significantly rise during the reaction.

Adversalen of P by sells is influenced by the species and concentrations of castens in the system. Divolent castens enhance P serption relative to menovalent castens. For example calcium esturated clays have more accessible positively charged edge sizes. And 1 mag of exchangeable AI/100g sell when completely hydrolyced may fix up to 102 gpm of P

in solution. Also both inorganic and organic anions can compete with P for sorption sites.

Organic anions which form stable complexes with Fe and AI of soil components are paticularly effective in reducing P sorption.

P adeorption has been found to correlate with the organic carbon content of soils. This correlation results from the exchange of P anions with hydroxil groups in organic matter and from the effect of P adeorption by AI and Fe associated with organic colloids. Since humas is normally negatively charged, it is not thought to retain much P by itself in soils. However, significant amounts of P are retained by organic matter that is associated with Ca, Fe and AI. For example, when manure was added over a long period of time, Weir and Soper (1983) detected an increase in Fe associated P. Also humic acid reacts with AI from soil minerals to form hydraxyl-AI-humic acid complexes which give rise to new surfaces for P adeorption (Appelt et al., 1976).

Calcium carbonate develops a negative charge due to the greater tendency for Calions than CO3 ions to go into solution. Phosphate ions then become chemisorbed at the surface. This reaction of P with calcite surfaces involves absorption of small amounts of P followed by precipitation of calcium phosphate (Griffin and Jurinak, 1973, 1974). The initial adsorption is thought to eccur at certain alses where phosphate ions form clusters which act as nuclei for subsequent crystal growth. Impure calcites and those of high specific surface result in more adsorption of P and more rapid formation of calcium phosphate precipitates. Therefore, phosphate activity will be lower in those soils that have a high calcium activity, a large amount of calcium carbonate and/or a large amount of calcium saturated clay.

2.1.3. Sell Schelen P

2.1.3.1. P form in sell selection

Plant roots absorb P from soil solution. Therefore, an understanding of the forms of P in soil solution and mechanisms that maintain adequate concentrations are important.

P is a component in sell organic metter. When soluble organic completes are present they will contribute to the total P in sell colution.

Polyphosphetes form several ionic species in solution. The $H_2P_2O_7$ and the HP_2O_7 ions are the predominent polyphosphete species in solution in the pH range of most soils. Polyphosphetes are metastable in soils and hydrolyse to orthophosphete.

Phosphete ions, ions pairs, and complexes are present in soil solution, but the orthophosphetes are the most stable forms. The pH dependency of orthophosphete species allows for the calculation of the activity of any one of the orthophosphoric acid species when the P concentration in solution is known (Figure 1.1). Phosphete combines with many of the di- and trivalent cations to form ions pairs and complexes. Whenever ionic activities are calculated from total P in solution, these ion pairs and complexes must be considered. P is absorbed by plants largely as primary and ascendary phosphete ions (H2PO4 and HPO4), which are present in solution.

2.1.3.2. Phosphote concentration required by plants

Maintainance of adequate concentrations of orthophosphate ions in soil solution is essential to plant growth. Classon and Barber (1974) use kinetics to depict a plants dependence upon the P concentration in soil solution. The required phosphate concentration in soil solution depends primerily on the crop species and the level of production desired. Tiedate at al (1965) list a range of phosphate concentrations required for the growth of a variety of crops, ranging from 0.003 to 0.3 ppm, with wheat requiring 0.01 to 0.03 ppm. Also, the optimum phosphate concentration varies with the stage of growth and the occurrence of stress.

Maintainence of adequate phosphate concentration in soil solution depends on the solid phase P entering into adultion to replace the amounts withdrawn by plant uptate. There are two different theories explaining the relationship between solid and solution phase P. The first describes the quantity (Q) of soil P as the amount of P in the solid phase that sate as a reserve and the intensity (I) refers to the phosphate concentration of soil solution. The second theory depicts an equilibrium resolien between soil solution P and two levels of solid P, the labile and non-labile freedoms (Barrow, 1980). The table soil P is the readily

evallable P that permits rapid dissociation. The depletion of labile P initiates the slow movement of non-labile P to labile P.

2.2. FATE OF FERTILIZER P WITHIN THE P CYCLE

2.2.1. Soil-Fertilizer P Reaction

2.2.1.1. Introduction

When P fertilizers are added to the soil nearly saturated solution of the phosphorus fertilizer material forms in and around fertilizer granules. Demotic notantial differences bring more water to the P saturated solution causing fertilizer solution to move to the autrounding soil (Chasewneth et al., 1974). As the saturated solution of phosphorus salts move into the first increments of soil, the chemical environment is dominated by the solution preparties rather than by the soil properties. The soil components after their solubilities as a result of the ion charge and pH changes caused by the fertilizer solution. This increased solubility of soil components will then allow for a flush of cations to enter the solution thus altering the solution composition.

2.2.1.2. Sources for Procedes lone

The sell acts as a cink for the ions in the fertilizer solution and also as a source of reactive cations. The major source for basic cations is the exchange complex. The sum of exchangeable cations varies from 1 cmal(+) Arg sell on coarse textured sells to 50 to 60 cmal(+) Arg sell on time sexterned sells. The exchangeable ions in productive agricultural sells are almost charge $\text{Ce}^{2+} > \text{Mg}^{2+} > \text{K}^+ = \text{Ne}^+$ (Bohn et al, 1967). Specific et al (1963e,h) determined that monomorphism clays did not express differences in exchange leatherms of basic cations. However, when erganic master is present, the exchange leatherms indicate a professore of the sell for calcium and magnesium over sedium, and for calcium over magnesium.

While the exchange complex is the source for the basic cations, metal ions like allicon, iron, aluminum and mangances are derived from the soil mineral constituents. The low pH of saturated MAP fertilizer solution will initiate the dissolution of AI, Fe and Mn axides releasing these cations into solution. Also some soil minerals near the fertilizer site may undergo dissolution, placing relatively large quantities of reactive cations (AI, Fe and Ca) into solution (Lahr, 1972). Lindsay and Stevenson (1959a,b) found large concentrations of AI and Fe in the fertilizer solution. As all these ions enter solution, their reaction with the added shoothate would seem iminent.

2.2.1.3. Thermodynamics of Phosphate Reactions

With the dissolution of phosphete fertilizer granules, ion activities in sall solution become supersaturated with respect to many phosphete minerals. The soil matrix acts as a source of water for granule dissolution and as a sink for the fertilizer solution (Bouldin et al, 1960). As successive increments of sell are contacted by the moving front of fertilizer solution, sell derived cetion concentrations build up both from the replacement of exchangeable cetions and from the dissolution of sell minerals (Kittrick and Jackson, 1965 & 1966). These ions in solution can react with fertilizer phosphete to precipitate as phosphete minerals.

When the ion activity product of a mineral is greater than its solubility product, the solution is superesturated with respect to that mineral. When a solution is superesturated thermodynamics indicate that the mineral in question precipitates; likewise when a solution is underesturated thermodynamics indicate that the mineral in question should be undergoes dissolution. Therefore, when a mineral is in thermodynamic equilibrium with a solution, the ion application will correspond to the solutions product of the mineral in question.

Phosphoto mineral achieffly leafines are a way of deploting how mineral superesturation values change. The construction of these curves requires the calculation of the free lanks consentration in solution by taking into account ion-pair and complex

formation. To calculate these ion activities in the presence of multicharged ions, an extended form of the Debye-Huckel (Device, 1962) may be used.

Solutions containing calcium and phosphate are very complex as many mineral phases become supersaturated. The possible solid phases resulting from these solutions, in order of increasing solubility, are: Hydroxyspesite (HAP), Ca₅(PO₄)₃OH; whitiochite (W), Ca₄H(PO₄)₂; octacalcium phosphate (OCP), Ca₄H(PO₄)₃·2.5H₂O; monetite (M), CaHPO₄; and Brushite (M), CaHPO₄·2H₂O. In addition to these crystalline forms there is an amorphous phase (ACP) of probable composition Ca₃(PO₄)₂·xH₂O. Since kinetics is a determining property in addition to thermodynamics, often the solid phase that precipitates first undergoes changes in solution towards phases of higher stability.

2.2.1.4. Nucleation of maction archites

The precipitation process, therefore, involves more than thermodynamics, as can be determined by figure 2. The first stage, nucleation, corresponds to the production of new centers from which mineral growth can occur. Phosphate minerals then form either through crystal growth or by the aggregation and coagulation of crystallites. Therefore, without nucleation phosphate mineral precipitates cannot occur. It is thought that every mineral has a critical limit of supercenturation Plancelles, 1984) or formation product (Abbana et al., 1988) that allows for the initiation of the process.

Knowledge of nucleation rate in addition to superseturation, seem to be required to understand the dynamics of phosphate mineral formation. Minerals such as Brushles with higher solubilities but lower critical superseturation values than minerals the HAP, often form in solutions that are measurable with respect to HAP. Therefore, these measurable solutions support a higher $H_2PQ_4^-$ autivity than what would be expected by thermodynamics (b. the most stable minerals controlling solution P activity).

Abbana et al (1900) found that in calcium and magazium phosphate salutions, Brushite and manadia occur as the final provipitate phosos, even though only manadia is thermodynamically stable. Brushite owes its wider ecourence to its higher nucleation rate, and its persistence in solution to the high value of the critical supersaturation of monetite. The precipitation of the ACP is usually accompanied by the nucleation of Brushite. White the Brushite crystals precipitate, the amorphous phase dissolves at a rate which is related to the growth of Brushite crystals.

Therefore, the calcium phosphetes which precipitate first in solutions of high to medium concentration do not always have the highest supersaturation values (Abbone, 1988). For example, soluble minerals the brushite and ACP usually form before stable minerals the HAP because their lower fermation product values offset the high supersaturation values of HAP. Abbone at al (1988) also found that no major changes occurred in these metastable solutions with respect to brushite.

2.2.1.5. Nature of mection products.

Brushite (CaHPO4'2H2O) is the most important product of precipitation in moderately sold solutions of 4.0<pH<6.0. Lundager Medicen (1984) found that superseturated solutions produced precipitates of (1)brushite, (2)monotite, (3)OCP, (4)amorphous TCP. Brushite was found to be a precursor of OCP, serving as a substrate for the latter. At pH>6, brushite is unstable with respect to OCP; the transformation takes place in a few hours, but is usually preceded by an induction period which may amount to several days (Lundager Medicen, 1984). The final products seem to be metastable with respect to more stable phases.

One of the possiliarities of the aqueous calcium pheaphate system is the occurrence of an amorphous phase. It is generally recognized that it is formed in the higher pH range and that it is a hydrous tricalcium pheaphate (Cag(PO₄)₂'::H₂O). ACP is also thought of as the critical model of whitestite. ACP has also been reported to contain Mg when precipitated from solutions containing Mg. ACP is measurable over the whole pH range and may be formed simultaneously with crystalline precipitates.

Knowledge of the conditions for precipitation of ACP is accorded for understanding the behavior of superconstructed coloium phosphate solutions, since it strongly. Influences the

occurrence of domeins of other phases (Lundager et al, 1986). For example, as ACP precipitates Ca and P are consumed and pH decreases; this reduces the supersaturation of other calcium phosphates often below the critical value of supersaturation. However, brushite, struvite and calcium exalete were often observed to crystallize more or less simultaneously with the formation of ACP (Lundager et al, 1986).

When Mg is in high concentration in phosphate solutions struvite (MgHH4PO4*8H2Otend newberyles (MgHPO4*3H2O) are often the main reaction products (Abbane et al,1982). However, displie the fact that solutions of Mg, NH4 and H2PO4 easily form superesturated systems, newberyles and struvite in general do not precipitate under low superesturation values (Abbane et al, 1982). Struvite precipitates alone or with the other phosphates depending upon the pH and solution composition. Struvite salubility and nucleation are affected by the phosphatic species which is dominant at the pH of the solution. Therefore the nucleation of minerals in such solutions is mainly controlled by thermodynamic parameters (Abbane and Belessie, 1986).

Treces of Mg added to calcium phosphete solutions supersaturated with respect to HAP, TCP, QCP, and brushite reduce the rate of calcium phosphete crystallization and delay the transformation of ACP to more stable phases. The Mg ions kinetically hinder the nucleation and subsequent growth of calcium phosphete minerals by competing for lattice alter with the chemically similar but larger Ca ions. Salimi et al (1985) noted that although the presence of Mg ions markedly retards the rate of seeded QCP crystallization, there was no detectable affect on the rate of brushite fermation. Where Mg ion concentrations are appreciable the marked reduction in the rate of QCP crystallization may stabilize the more acidic hydrated calcium phosphetes such as brushite during precipitation receitens. Therefore, the numerous observations that brushite is a viable phase may be traced to the

The presipitation products found in pure existent plicityhete colutions are also found when phosphate colutions are added to cell or self-deletion. Morene, Brown and Ochomo

(1960) found that, at 25 C, brushite hydrolysed to OCP in soil solutions with a pH value of 6.36 or higher and that this hydrolysis reaction increased with increasing pH. They found that OCP was metastable with respect to HAP. Bell and Black (1970a) detected OCP 4 weeks after brushite was added to the soil, however they could not detect the more stable HAP in the soil even after 44 weeks. This hydyloysis of brushite to OCP was noted to be inhibited by the presence of a soil solution concentration of Mg of 10⁻³M. In the presence of soil solution Mg, struvite and newberylte precipitate in addition to brushite (Bell and Black, 1970b; Recz and Soper, 1967; Lindsey et al. 1962).

2.2.1.6. Mobility of Added Phosphate

Khasawach et al (1974) found that when P fertilizers were bended at a rate of 112kg P/ha, P moved up to 3cm from the pellot with most of that P approximately within 1.5 to 2cm from the band. When Bell and Black (1970b) placed 2.28 g MAP on top of a column of soil, the P fertilizer was detected 2.8cm down in a SiCL soil. They noted that P movement was lower in soils with monomorfilante than in soils with kaolinite or vermiculties the deminant clay mineral.

However very few researchers have been able to establish the presence of precipitated reaction products at verying distances away from a fertilizer using direct methods. Bell and Black (1970s) identified traces of Brushite and Struvite 3 to 5cm away from a MAP fertilizer alts. Sample (1972)determined that adsorption could account for 20% of P retention adjacent to the fertilizer pollet and 80% at a distance of 15mm.

2.2.1.7. Role to coll P couls

When a P fertilleer is added to sell, a complex sequence of reactions fellows. Three zeroes are generally recognized mear a fertilleer granule (Sample et al, 1980). The control zeroe contains the residue of the P fertilleer. Around this zeroe the concentrated solution results in the precipitation of P minerals. This zero grades into an outer zero in which concentrations are lower and P resistence are mainly with elements on the surface of sell particles. Therefore, the rate at which the sell P syste returns to steady state conditions

after fertilizer addition increases as one moves away from the fertilizer zone.

2.2.2. Effect of Management upon the Fate of Fertilizer P.

2.2.2.1. Introduction

Many factors are involved in P fertilizer management. However, granule placement and size are the management factors of significance to this study.

2.2.2.2. Uotaka Efficiency of P

Most experiments show that the uptake efficiency of added P fertilizer in the first sesson of application is less than 25% (Barber, 1978). At an application rate of 8.4kg P/hs, Egball and Sander (1988) showed how a particle mass of 0.39 mg resulted in an increase in uptake efficiency from 26 to 47% over a particle size of 20 mg.

2.2.2.3. Placement

P placement has recieved considerable research attention because with time soluble phosphates revert to less available forms and remain immobile in the soil. Four broad objectives are involved in studying fertilizer placement. These are: (ii) to enhance the efficient use of fertilizer by the plant, (iii) to prevent or reduce environmental contamination, (iii) to prevent fertilizer selt injury to plants, and (iv) to provide an economical and convenient operation (Plandell et al, 1985). These factors are quite important as they relate to the crop and its rooting pattern, the concentration and closeness of fertilizer to the seed, and the soil test level. (Plandell and Hooft, 1986).

Sell placement methods involve varying the valume of sell that the fartilleer is to react with. As mixing of the fartilleer is restricted to a smaller sell valume, the P levels in the fartilleed sell increase. Higher P levels in the fixed sell valume mean the nutrients remain available for a longer period of time. Band placement decreases the sell surface area in contact with added fartilleer with a consequent reduction in the amount of fixeden. A possible drawback of smaller sell valumes per mass of fartilleer is that a smaller rest mass is responsible for the P upsales. Thus, for maximum fartilleer efficiency, a compremise must be

reached between the soil volume that is fertilized and the root distribution so the probability of root-fertilizer contact is optimized and fixation minimized.

It is generally found, with low fertilizer rates, the efficiency of nutrient uptake is greater for bend then broadcast application. For soils with low soil test available P, placement in the effective rooting zone generally results in greater efficiency of P use (Young et al., 1985; Engelsted and Terman, 1980). Slight et al (1984) also concluded that the early beneficial effects of bending are obtained mainly from placing all of the fertilizer P where contact by active roots is likely, so localized placement of fertilizer should be more efficient then broadcast application.

However, bending alone on low P soils is shown to be inadequate and supplementary broadcast P is needed to reach top yields (Barber 1958; Welch et al, 1966b). Wager et al (1965) also found the combination of broadcast and seed placed treatments enhance P uptake.

Plants take up P when it is in solution; in a dry climete seed placed P may not enter solution and thus would not be available to the plant. Herapiak and Beaton (1986) found that under dry conditions, wheat responses are greatest when all of the fertilizer P is deep-banded and smallest when the P is seed placed. The subsoil retains a higher moisture content allowing P to remain in solution longer. McConnel et al (1986) also found deep bends 15cm deep to be the most efficient depth for P fertilizer uptake.

Bending an immobile nutrient such as P generally becomes less important as soil P reserves rise over time. Wager et al (1985) found that a single broadcast application of 40 kg P he⁻¹ increased yields for 5 years and increased average yield and P uptake similar to that of annual seed placed applications of 10 and 20 kg P he⁻¹. This indicates a single breadcast P application can produce yields comparable to annual seed placed P applications without requiring additional fertilizer inputs.

2.2.2.4. Fertilizer Granule Size

Particle size of P fertilizer granules influences the utilization efficiency of applied P by affecting the distribution of P within the band and by determining the effective surface area of the fertilizer granule. The distribution of fertilizer P within the band, whether continuous or discontinuous, is shown to be an important factor in the efficiency of applied P (Anghinoni and Berber, 1980; Sleight et al, 1984). The effective surface area of a fertilizer granule is related to the reactivity of P with the soil (Kilmer and Webb, 1988).

It is widely recognized that the optimum distribution of P granules has to be weighed against the rate of P fixation. Distribution of pellets within a bend influences contact of roots with fertilizer P (Berber, 1974). Nonuniform distribution of fertilizer P within a bend affects root contact and thus yield (Stryker, 1974). While it has been shown that the total P needs of a plant can be met by a single root (Kiasal and Ragland, 1967), it is desireable to have more roots exposed to available P (Stryer et al., 1974). When fertilizer perticles are not placed in a continuous band the number of root fertilizer contacts have to increase to attain the same yield (Eghball and Sander, 1988).

Numerous investigators have reported increases in early crop response with an increase in granule size of water soluble P fartilizers v.co. applied at a low rate to acid solls. For water soluble P fartilizer granules up to 6mm die. .ar, effectiveness is related to the amount of water soluble P per granule which in turn determines the volume of soil affected by P (Taylor and Terman, 1964; Sample and Taylor, 1964). Crop response to larger granules depends on the probability of roots finding the very few diffusion zones at a given rate of P application (Moreno, 1966; van Burg, 1963).

Granulation or banding of existum phosphotos has little or no agranomic adventage for alkaline soils. The local acidification near MAP fortilleer as NH4-N is nitrified may increase availability of P. Solutions, suspensions and solid P fortilleers having similar solubilities and when similarly placed in an alkaline soil at a given P rate, show similar

availability to crops. P solubility effects related to granule size in general tend to be obscured at higher available P levels in soils.

Particle size of P fertilizer significantly affects measured crop parameters only on soils which have the greatest yield increases due to applied F. Sender and Eggbell (1988) observed the optimum fertilizer particle size within a bend at 8.4kg P/ha to be 0.39mg/particle. Grain yields were lower for smaller size pritis because of the larger surface area for reaction with soil constituents which reduces fertilizer P availability. Grain yield was lower for larger prill sizes because root - P fertilizer contact was probably lessened by a discontinuous band. This mid range particle size had an average specing within the band of only 0.003cm. Sender and Egbell (1988) noted that as the rate of P application increased to 16.8 and 23.2 kg/ha the effects of particle size and surface area disappear. It is apparent that yield differences due to P fertilizer particle size were most important on soils low to very low in available P where yield responses to applied P were high.

2.3. OBJECTIVES

The objectives of this thesis were to:

- 1) Identify the reactions that occur when MAP fertilizer is placed in soil. This objective will be analysed by menitering the ion concentrations in soil solution before and after the addition of MAP fertilizer. Direct observation of P mineral precipitates formed from added P will be attempted with the use of soil thin sections.
- 2) Identify the movement of added P through the sell P cycle in the season of application. This objective will be accomplished by mentering the sell P fractions throughout a 3 menth period.
- 3) Identify how the efficiency of the uptake of added P can be effected the granule size and bend placement of MAP fertilizer. This objective will be accomplished by menturing the uptake of ³²P by barley. Organic amendments will also be used in an attempt to reduce the context of added P with the cell mental.

3. MATERIALS AND METHODS

3.1 Soil Properties and Semale Properation

Soils of the Breton (Orthic Gray Luvisol) series and Paddockwood (Dark Gray Chernozem) association were selected as representative of soils which demonstrate low apparent recoveries of added P. The Breton samples were taken from research plots in NE25-47-4-W5 near Breton, Alberta. The Paddockwood samples were taken from a field in NE31-50-10-W3 northeast of Prince Albert. Seekstchewen. Solls of the Breton series and soils of the Paddockwood association are developed on glacial till parent material. The experimental soil sample for the Breton series was taken from a field which had not recieved P fertilizer for 20 years, whereas the experimental soil sample for the Peddeckwood association was taken from a field which has received approximately 15 kg P hs⁻¹ for 35 years. Serveral routine analyses were conducted on these experimental soils. Surface samples (Ap horizon) of each of these soils were collected from 0-15cm depth, air dried and passed through a 5mm slave. Particle size analysis was determined using a pipotte method (Day, 1966), pH was measured using both 2:1 CaCl₂:soil and 2:1 water:soil ratios (M^CLean, 1982). Organic carbon was determined with a Loco combustion furnace (Nelson and Semmers, 1982). Cation exchange (CEC) analyses were conducted using 1M ammenium acetate (pH 7) for the displacement of exchangeable cations followed by alcohol washing and replacement of exchangeable ammonium with sedium ions (Chapman, 1965). Exchangeable cations (Ca. Mg. No. K) from the ammonium acetate extraction were determined by stemic obserption (Instrumentation Laboratory) (Baker and Sulv., 1982) and CEC was determined by measuring displaced ammenium with an autoensiyeer (Technican) (Pheedes, 1982). Extresiable eluminum, Iron and mangeness were determined using both ammentum evalute (AC) entrestant and distante-altrate-bleerbenate (DCB) redustantentrestant (McKeegue, 1978). The entrested AO and DCB estudens were analysed for AI, Fe and Mn using an ICP (Department of Chemistry-Spectral Services- University of Alberta). The clay and allt fractions separated during particle size analysis were subjected to X-ray diffraction analysis for minerological determination. The Breton and Paddockwood clay samples were prepared using the paste technique and were subjected to seven pre-treatments to permit a clear identification of the minerals present (De Kimpe, 1978). The x-ray diffractograms were prepared using Cok radiation at 50 kv and 25mA, and at a scan appeal of 1°20 min⁻¹.

3.2 Greenhouse Experiment

During the first part of the experiment plastic pane were set up in the greenhouse to determine the effect of different fertiliser granule sizes and methods of application on P fortilizer uptake by plants of barley (Hardeum variane L. cv Express) and transformations within the soil. Sloved soil meterials were placed in each of the 30cm (length) by 25cm(width) by 20cm (height) pane. Each pan was an experimental unit. MA³²P fertilizer bands were placed 2cm below and 2cm to the side of the seed row. The locations of fertilizer granules were marked for identification with plastic toothpicks. In other treatments fertilizer bands were also placed 2cm to theside by 10cm below the seed row. After garmination the sell meterial was wetted regularly to keep the moisture content near field casecity.

There were seven treatments applied, each replicated three times for each of the solls plus three additional treatments replicated three times for the Breton sell only (Table 3.1). The variables in the experiment were furtilizer granule size, furtilizer application location and addition of an organic amendment. $MA^{32}P$ furtilizer was added as a tracer of P upushe by barby. The MAP furtilizer was added at a rate of 20 kg ha^{-1} (or 0.50g MAP pat^{-1}), K_280_4 and MI_4M0_3 were added to the sell surface of all treatments at a rate of 20 and 80 kg ha^{-1} , respectively. Thirty barby seeds were added per pat.

30

TABLE 3.1. Form of fertilizer application, depth of application and nature of organic amendment used in the greenhouse experiment.

Code	Mass of MAP fertilizer granule (g)	Number of gramules per pot	Depth of band below seed row(cm)	Carbon source for amendment/ encasement
Control	•	•	•	•
SolutionP	•	•	2	•
0.01g ²	0.01	50	2	-
0.10	0.10	5	2	-
0.10g ² 0.50g ²	0.50	1	2	•
Straw	0.10	5	2	Straw
Peal	0.10	5	2	Post
Straw ² Peat ³ 0.01gds	0.01	50	10	•
0.10gd#	0.10	5	10	•
0.50gd#	0.50	Ī	10	•

[#] Indicate treatments applied to Breton soil only.
s,d Indicates the application of MAP fertilizer in shallow and deep bands, respectively.

Plants were harvested at 30, 60 and 90 days by cutting the shoots at ground level. The hervested plant material was dried and weighed. The dried plant material was then acid digested (Olsen and Sommers, 1982) and measurement of ³²P uptake was conducted with a scintillation counter.

After the third hervest, soil cores 6cm in diameter were removed for construction of soil thin sections. The sell cores were centered around previously merked fertilizer placements. The soil cores were sir-dried in the laboratory, impregnated with apoxy resin and cut into 7 by 5 cm thin sections using procedures described by Brewer and Pavoluk (1975).

Soil cores of 2cm diameter were also collected at three leteral distances from the fertilizer placement shelf0-1cm, 2-4cm and 5-7cm). Each 2cm diameter core was then divided at the depths 0-2cm, 2-4cm and 4-5cm. The sell material was analyzed for organic P content by the ignition method (McKeegue, 1978). The sell material was also analyzed to determine the distribution of P among sell P fractions using a sequential extraction procedure (O'Helleren et al. 1967).

Soil thin sections were analyzed patrographically for soil fabric and for P crystal identification. The thin sections were also mapped for P, Ca, Mg, Mn, Fe, Al and Si on the SEM/EDX to determine the distribution pattern and elemental association of added P. Mean contrast analyses were conducted on 32P uptake and yield data by the General Linear Models program of SAS (1986).

3.3 Smillbootles Superinees

The Broton and Paddechareed sell metarials were also used in this experiment. MAP forefleer solution was evenly added to 250g of sleved sell to bring the molecure content to field expectly. Forefleer rates corresponded to the P levels found within 1cm radius of the 0.10g granulo in the greenhouse experiment. The sell metarials equilibrated with the MAP in plants corresponds to they are the particles. Delenhood

water was added regularly to maintain field capacity. Thus there were four treatments and three time periods each replicated three times (Table 3.2); two of the treatments were not tested with the Peddockwood soil.

The soil pore water was extracted at each time period by immiscible displacement (Barrow, 1962). The following chemical analyses were performed on the pore water extected by immiscible displacement from the equilibrating soil samples: the cations Ca, Mg, Na. K. Fe. Al. Si and Mn were aniyeed by atomic absorption (Beker and Suhr, 1982; Rhoades, 1982); the anions PO4, SO4 and NO3 were analysed by ion chromotography (Dionex); ammonium ion was anlysed by an autoenelyser (Technicon) (Keeney and Nelson, 1982); and electrical conductivity (EC) and pH were analysed with a conductivity meter and an electrode, respectively (Rhoedes, 1982). These data were then used to calculate len activities with the use of the thermodynamic models 'Soll Solution' and 'MINTEQ'. Analytical concentrations of cations and anions together with pH and EC values, all derived from extracted pare water, were used as input parameters for the models. The calculated ion activities were then used to calculate the IAP of the solution species which compass the clay and P minerals of interest. The Kap of these minerals were calculated separately using thermodynamic data obtained from Rai and Lindsay (1975) and Lindsay (1979). Tables 3.3 and 3.4 show the Ken values calculated at 25C for selected P and clay minerals considered in this thesis their and Linear, 1975; Lindayr, 1979). The IAP calculated from modelled ion activities was then compared to the Kap values for minerals listed in tables 3.3 and 3.4 to obtain the acturation index (SI) value (log IAPKap). SI<0, SI=0, and SI>0 in legarithinis form rater to underseturated, acturated and superacturated conditions, respectively, with reference to the minerals in question. The ion activities were also compared to day mineral and P mineral leeline diagrams. Samples from the same experimental unit also underwant sequential entraction to determine the distribution of sell P among fractions (O'Helleren et **d.** 1987).

The freedometen precedure was conducted as a split-plot experiment

33
TABLE 3.2. Concentration of P and equilibration time for P equilibrium studies.

Soil	Equilibrating P concentration (mg P/g soil)	Time	Amount of alfalfa amondment (g)
Breton	0 5.2	1 hr	0 0 50
	5.2	1 he	0
	5.2	1 hr	50
	16	1 hr	0
	0	30 days	0
	5.2	30 days	0
	5.2	30 days	50
	16	30 days	0
	0	60 days	0
	5.2	60 days	0
	5.2	60 days	50
	16	60 days	0
Tiller	0	1 hr	
	0 5.2	1 br	
	0	30 days	
	0 5.2	30 days	
		-	
	0	60 days	
	5.2	60 days	

TABLE 3.3. Equilibrium constants of relovant clay mineral equilibria.

Dancting		Log K
KALSO 16(OR) Simmoovies) + 1081*	= K ⁺ + 3A ³⁺ + 3H ₄ SiO ₄	17.08
MayAlyBigOlg(Olf)g(chlorin) + 16H+	= 5Mg ²⁺ + 2Al ³⁺ +3H ₄ SiQ ₄ + 6H ₂ O	72.97
Algeria (Off) planteles + 6H+	= 2A1 ³⁺ + 2H4SiQ ₄ + H ₂ O	6.67
(47), gy/le, 12/44, gy/leg, gy/leg, 45/0/gy/OH), j	= 3.16Al ⁺³ + 0.58Mg ²⁺ +0.45Fe ³⁺ +7.57Hg SlO _g	0.48

my, 1975. A Thermodynamic Model for Predicting the Formation, Stability, and Weathering of L. Sell Science Society of America Proceedings, Vol. 39, p.991-996.

TABLE 3.4. Equilibrium constants of relevant phosphete mineral equilibria.

Laction		Los K
APO42H2O(varianths) + 2H+	= AI ⁺³ +H ₂ PQ ₄ ⁻ +2H ₂ O	-2.50
P#O ₄ + 281*	= Fe ³⁺ + H ₂ PO ₄ .	-5.37
PdO4/285_O(strengths) + 251*	= F43+ + Hy PQ4- + 21h O	6
CallFO ₄ ZH ₂ O@makke) + ZH ₂ O + H ⁺	= Cz ²⁺ + H ₂ PO ₄ + 2H20	0.63
Call(PO ₄) ₃ ·2.5H ₂ O(ectecalcies phosphes) + 5H ⁺	= 4Cz²+ + HyPQ4" + 2.5HyO	11.76
ChyffO ₄) ₂ (trinsteinm phosphere) + 4H ⁺	= 3Cs ²⁺ + 21hPO ₄ .	10.18
CayPO4)ORPydemyspeths) + 7H+	= 5C2 ²⁺ + 3H ₂ PQ ₄ ⁻ + H ₂ O	7.
Marie O4:382 Openhenryke) + H+	= Mg ²⁺ + H ₂ PO ₄ · + 3H ₂ O	1.38
Martin Pol, 4815 Operates) + 281 +	= Mg ²⁺ + NH ⁴⁺ + H ₂ PQ ₄ + 6H ₂ O	6.40

cente: Linday, 1979. Chemical Bankhein.

(Dr. Hardin, Dept. of Animal Science, University of Alberta, 1990, personal communication), and the pore water data was analysed as a two way anova. The analyses were conducted by the General Linear Models program of SAS (1985).

4. Discussion

4.1 Soil Characteristics

4.1.1 Introduction

Both the Breton and Paddockwood soils were developed on glacial till parent materials. However, the geographical distance between sites suggests the glacial till parent materials probably originated from different bedrock formations. The geographical location of the two soils, on the fringes of the boreal forest, would generate similar environmental conditions responsible for soil formation. The soil properties reflecting from the parent material and environmental conditions can greatly affect the way added P reacts with the soil.

4.1.2 Soil Mineralogy

The mineralogy of the clay and silk fractions of the two soils is very similar (Table 4.1, Appendix I). Smooths is the dominant clay mineral of both soils. The mineralogy of the clay and silk fractions is similar display the distance between the two sampling sites.

The most important characteristics of the mineralogy of the soil clay fraction that relate to added P are the ion exchange perperties. The ion exchange properties of soil clays very with the mineral crystal structure. Smectle minerals have a large CEC of approximately 110 email (+) kg-1 (Berchardt, 1977). This high CEC is a result of the dominant pH-independent charge associated with amost to minerals. The exchange complex of amost to minerals allows must monovalent and divolent actions to be completely exchangeable at all pH values. A list of the mechanisms involved in the exchange reactions of amost to minerals is exclined in Table 4.2.

Miles has a law CEC relative to amostle slay minerals because the interlayer estions

TABLE 4.1. Minerology of the clay (<2 μ m) and silt (2-20 μ m) fractions of the Paddockwood and Breton soils.

Soil	Fraction	Minerology in order of decreasing abundance
Paddockwood	Clay	Smectite, mica, kaolinite, quartz, chlorite
	silt	Quarts, feldspar
Breton	Clay	Smectite, mica, kaolinte, quartz, chlorite
	silt	Quartz feldspar

TABLE 4.2. Reactions of various elements with smectites in soil environments.

Element	Type of Reaction	
Ca ²⁺ , K ⁺ , NH ₄ ⁺	Cation exchange	
H ₃ O ⁺	Cation exchange, replaces octahedral Al	
Na ⁺	Cation exchange, replaced from exchange by octahedral Mg and Al	
Fe ³⁺	Cation exchange, hydrolysis, and dissolution of smectite	
Al ³⁺ , Mg ²⁺ , Fe ³⁺	Cation exchange, hydroxy interlayer formation	
H ₂ PO ₄	Anion exchange	

Source: Borchardt, 1977. Montmorillonite and Smectite Minerals IN Minerals in Soil Environments. Dixon Ed. ASA. Madison Wisconsin.

are not exchangeable. Potassium dominates the interlayer region and is released slowly through selective exchange with other monovalent ions such as ammonium (Fanning and Karamides, 1977).

Kaolinite also has a low CEC relative to amerite clay minerals. The low CEC of kaolinite is a result of low pH independent - permanent negative charge and substantial pH dependent charge. Therefore, kaolinite minerals have ion exchange properties similar to amorphous and oxide minerals. Despite the low CEC, exchangeable sodium may be retained in kaolinite mineral structure under acid conditions (Dison, 1977).

Soil extraction with A.O. and D.C.B. (Table 4.3) indicate the presence of oxides of AI, Fe and Mn within the soil matrix. Iron and manganese nodules within the soil matrix were also detected on soil thin sections (Plate 4.1).

Plete 4.1. EDX maps of seequioxidic nodules in the Paddockwood sample: a) the nodules are composed of iron and mangenese oxides; b) P is not seed at the surface of the nodule.

4.1.3 General Analytical Properties of Experimental Soils

In addition to having similar mineralogy in the allt and clay fractions, the two experimental soils have similar particle also distributions (Table 4.4). The Paddockwood soil has a loam texture and the Breton soil has a ality learn texture. Therefore, based on clay mineralogy and particle also distribution the two experimental soils should have similar CEC values.

Actual measurements of CEC and exchangeable cations for the two experimental soils are also presented in Table 4.4. The Paddeskwood sample has a higher CEC (20 $cmol(+) log^{-1}$) then the Broton sample (14.5 $cmol(+) log^{-1}$), and it also has higher exchangeable calcium and magnesium values than the Broton sample.

The Broton samples have much higher values of DCB and AO extractable AI, Fe and Min than the Paddeckwood samples. The Paddeckwood sample has higher organic earbon content and pit values than the Broton sample (Table 4.4). Therefore, some sell properties

TABLE 4.3. Extractable Fe, Al and Mn of Experimental Soils.

Treatment	Al	Pe	Mn
		1	
		PADDOCKI	000
DCB	0.067	0.42	0.029
M O	0.086	0.24	0.029
		202700	
DCB	0.10	0.62	0.066
MO	0.13	0.44	0.000

Ptoto 4.1. EdX maps of sesquioxidic nodules in the Paddockwood sample: (a) the nodules are composed of iron and manganese oxides; (b) P is not associated with the surface of the nodule

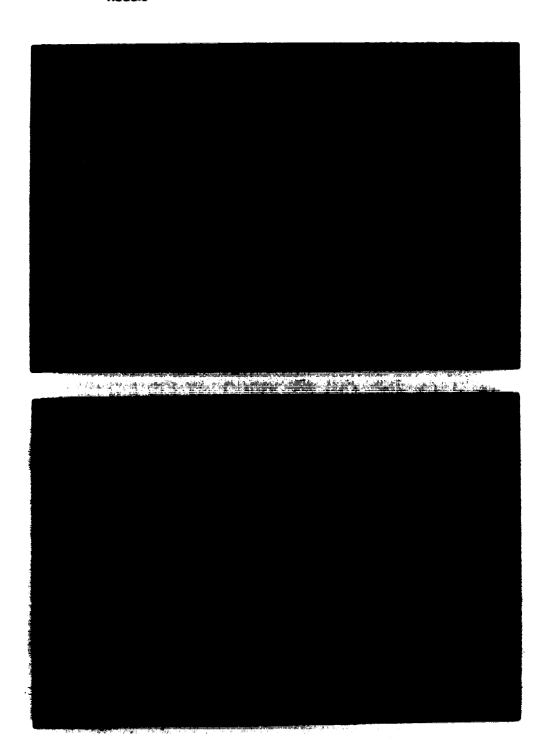


TABLE 4.4. Chemical properties of experimental soils.

1	6	CaC, HOR	8	400	OC Brownik Crim	iga ×	2	8	Participant	Particle size distribution Sand Silt Clay	ribation Clay
			*		ca of (+	1 -21(+),0				4	
Publicativood	3	6.8	3.4	20.9	2.8	0.37 0.41	0.41	90.0	38	\$	11
	73	5.8	3	12.3	17	0.28	0.45	14.5	35	5	±

are very contrasting between the two experimental soils.

The fertilizer history of the two soils previous to the time of sampling is quite different. The application of 15 kg P he⁻¹ annually to the Peddockwood sample site for the 35 years prior to sampling is in concrest to the Breton sample site which had no fertilizer addition during the same time period.

4.1.4. Soil Solution Characteristics

Breton and Paddockwood samples were equilibrated at field capacity. The pore water was extracted after equilibration periods of one hour, one month and two months. The soil solution was then analysed for a suite of ion concentrations.

The cations ${\rm Ce}^{2+}$, ${\rm Mg}^{2+}$ and ${\rm Ne}^+$ steadily increase and then stabilize with time while the remaining suite of cations maintain their original soil solution concentration through a two month period (Table 4.5 and 4.6). The concentrations of ${\rm Ce}^{2+}$, ${\rm Mg}^{2+}$ and ${\rm Ne}^+$ increased at the same relative rate and then stabilized near the same relative concentrations.

The increase and subsequent strbillastion of Ca⁺², Mg⁺² and Ne⁺ ion concentrations may be the result of their dominance on the exchange complex of the two experimental soils (Tables 4.5 and 4.6). Therefore, their ion concentration in soil solution is a reflection of the equilibrium relationship between the soil solution and the exchange complex. We can then use these equilibrium values to determine the effect of MAP fertilizer addition to soil.

4.2. Characteristics of the Reaction Between Sell and Facilitar in Solution

Broson and Peddockwood complex were also equilibrated at field expectly after the addition of 5.2 flow rate) and 16 mg P/g sell (high rate) of MAP fartillizer. The pure water was extracted after equilibration periods of one hour,

one menth and two menths. The sell solution was then analysed for a suite of ion concentrations.

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o addition levels for u		5.2 + AMIG		8	2°128188°9818
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leld especity with three different				8	22262282828222
4	100	4		Я	2228125813824 F
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		4		8	32×22×2=2-2*
				4	33×3512×2=5.3×
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ns is significant at: ne, not eignificant; *, affected p < 0.05; **, adjusted p < 0.01; *** adjusted p < 0.001.

That 4.6. Ion concentration means of Peddockwood and Breton soil pore veter equilibrated at field capacity with added MAP fortiliser for up to two months.

[401 L402	80'0 80'0	0.04 0.04	2.2	0.07 0.05	30.0	0.02 0.17 71 39.2 32 41	0.4	ty level	ity level
1.2	TOM SO	4.1 5.1 5.5	30 6.4 4.7	6.2 3.3 3.2 54 20 14		280 100 93 570 110 61	• •	means at the 0.05 probability level	at the 0.05 probability level
P. in sentitibrating soil	•	2.0 6.3	10 24 27 3.5 5.1 6.1 0.23 0.24 0.33	9.0	1.9 2.6 3.3 1.1 1.1 2.1	0.27 0.24 0.28 0.46 0.49 0.44	34 34 33	the treatment means at	be treatment means at ments.
Concentration of 1/2 (mg P/s	900	1.8 8.2 8.3		3.6 2.7 3.0 45 23 26	10.00	150 120 130 200 200 130 2.7 3.0	2	itterence between the treatments.	lifferense between ti Broton sample treets
Description of the last of the	o casa coura	6.6 6.9 1.6 1.9 2.2 10 14 16	0.43 0.36 0.46	1.0 1.0 1.0 0.0 0.0 0.0	1.1 1.0 2.3 0.46 0.64 0.88	1.2 0.97 1.2	1000 01000	Posterior and	to Peddodinos and
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		8	8	•	1	ŧ	:	ŧ	ŧ	ŧ	į	2	ŧ	2	:
		•													

4.2.1 Sobsion Semi-Separated with MAP Fertilizer

The characteristics of a solution saturated with MAP fertilizer are presented in Table 4.7. The pH of a solution saturated with MAP fertilizer is 3.5 when equilibrium with the dissolution constant of $H_2PO_4^-$ is attained with an initial concentration of 2.9 M. The pure water extracted from the experimental soils one hour after the application of the low rate of P fertilizer (Tables 4.5 and 4.6) has characteristics similar to the solution saturated with MAP fertilizer. The pH of the extracted pure water (Tables 4.5 and 4.6) was slightly higher and the PO_4 and NH₄ concentrations were lower than the solution saturated with MAP fertilizer.

4.2.2. Flush of lone into Soil Schulen

The addition of MAP to sell caused a unique and consistent change in the ionic concentrations in the sell pere water. The EC of sell pere water dramatically increased with the addition of MAP fortilizer (Tables 4.5 and 4.6). The increase in EC of the pere water reflected increases in the ion concentration of all elements in addition to the components of the fertilizer solution, ammenium and phosphate. One hour after the addition of the low rate of MAP fertilizer the ion concentrations were generally two to three times higher than in the control sell solution (Tables 4.5 and 4.6). The ion concentrations of the basic cations and allics were slightly higher after the high rate of addition of MAP fertilizer. However, the ion concentrations of Iren, mangenese and aluminum were substantially higher after the addition of MAP fertilizer at the high rate as compared to the low rate (Table 4.5).

Both components of MAP fertilleer solution, ammonium and phosphoto, decrease to similar levels at different relative rates after their addition to the sell. One hour after addition, ammonium ion concentration was approximately one helf of the phosphoto ion concentration. This rapid decrease in ammonium ion concentration coincides with the ion flush into sell solution after the addition of MAP fertilleer. The ammonium ion coturates the auchange complex, decreasing its sell solution generaturation, and then releases the

TABLE 4.7. Characteristics of Saturated MAP Fertilizer Solution

Property	MAP Solution Value	
pH	3.5	
PO ₄ (mol/L)	2.9	
MH ₄ (mol/L)	2.9	

Source: Sample et al., IM Khasawneh et al., Eds., The Role of Phosphorus in Agriculture, p. 275. Medison, Wis.; ASA, 1980.

exchangeable cations into solution.

4.2.3. Change in Ion Concentration with Time After the Addition of MAP Fertilizer

One month after the addition of MAP fertilizer at the low rate the EC of the soil solutions decreased to and levelled off around 6 dS m⁻¹. The decrease in EC reflected decreases in concentrations of all ions except sulfate.

Calcium and magnesium show the greatest relative decrease in cation concentration with time after the initial flush caused by the addition of MAP fertilizer at the low rate (Tables 4.5 and 4.6). These two ions reach an equilibrium concentration which is below the control equilibrium concentration (Tables 4.5 and 4.6). The remainder of the cations in soil solution decrease to the level of concentration attained for the control treatment at equilibrium.

After the high rate of MAP fertilizer was added to the soil, the soil solution was dominated by ammonium and phosphate to such and extent that only minor changes in ion concentration occur after their initial flush into soil solution. However, calcium and manganese, in contrast to the remainder of the suite of ions analysed, had quite large decreases in ion concentration from the time of MAP fertilizer addition to a time of one month.

4.2.4 Change in Ion Concentration with Time After the Addition of MAP Fertilizer to Soil with Alfalfa Amendment

When MAP fertilizer was added to sell containing an alfalfa amendment many characteristics of the sell solution were different then when MAP fertilizer was added to the sell without the alfalfa amendment. With most sell solution parameters, the difference between the alfalfa and non-alfalfa treatments occur with the 30 day treatment.

The pH values of the alfalls treatment immediately after the addition of MAP fertilizer are similar to the non-alfalls treatment. The pH values between the two treatments are also quite similar after two mentils of equilibration. However, after one mentil of

equilibration the pH values of the alfalfa treatment after the addition of MAP fertilizer are two whole units higher than at 2 months (Table 4.4).

Soil solution values of calcium and magnesium are three to four times higher in the alfalfa treatment than in the non-alfalfa treatment one month after the addition of MAP fertilizer. The soil solution values of calcium and magnesium are similar for the two treatments two months after the addition of MAP fertilizer (Table 4.4).

Soil solution values of potassium are higher in the alfalfa treatment than in the nonalfalfa treatment one and two months after the addition of MAP fertilizer. Whereas sodium values are similar between treatments one and two months after the addition of MAP fertilizer (Table 4.4).

Soil solution values of iron and manganese are very high both one hour and two months after the addition of MAP fertilizer to the alfalfa treatment. The soil solution values for iron and manganese one month after MAP addition are similar between alfalfa and non-alfalfa treatments (Table 4.4).

Phosphete values in soil solution are reduced much faster in the alfalfa treatment than in the non-alfalfa treatment. One hour after the addition of MAP fertilizer the phosphete level in soil solution of the alfalfa treatment was half that of the non-alfalfa treatment (Tables 4.5 and 4.6). Also, the phosphete level in soil solution was lower in the alfalfa treatment than in the non-alfalfa treatment two months after MAP addition.

The nitrate levels detected in soil solution were very high one month after the addition of MAP fertilizer to the alfalfa amendment. The nitrate levels in soil solution were similar between alfalfa and non-alfalfa treatments one hour and 60 days after the addition of MAP fertilizer.

The color of the pore water after passing through 0.2µm filters were quite different. The pare water from the control, with no MAP added was clear. The pare water from the soil after MAP was added was clear but yellow (10Y 6/6). The pare water from the soil after MAP was added with an alfalfa amendment was blackleb brown (10YR 2/1) and

yellow, clear (10Y 6/6) after one and two months of equilibration, respectively.

4.3. Discussion of the Reaction Between Soil and MAP Fertilizer in Solution

Many researchers have detected a flush of soil cations into solution after the addition of MAP fertilizer (Lindsay et al., 1962; Racz and Soper, 1967; and Bell and Black, 1970a). Similar results have been detected here (Tables 4.5 and 4.6). The possible sources for this flush of ions are from the exchange complex, from oxides, oxyhydroxides and amorphous minerals, and from secondary clay minerals.

4.3.1. Characteristics of Solution Containing MAP Fartilizer Responsible for the Flush of Ions into Soil Solution

The low pH of solutions containing MAP fertilizer, when reacting with soil constituents, affects many properties. Increased ion solutility resulting from fertilizer induced soldification leads to the destabilization of soil minerals and soil organic metter. The soil as a complete unit may not be greatly affected by the low pH due to the buffering capacity of clays and organic metter (Barrow and Show, 1976). However, within the vicinity of solutions containing MAP fertilizer, the low pH has a great influence on soil properties which may greatly affect the fate of added P.

In addition to low pH, the high concentration of ammonium ion in the solutions containing MAP fertilleer reacts with the soil constituents. The ammonium ion has a relatively low hydration energy resulting in easy dehydration and strong resention by 2:1 clay srinerals (Bohn et al, 1987). Therefore, the high concentration of ammonium ion in solution will tend to replace enchangeable cations on the exchange complex. This may partially explain the increase in SC and ion concentrations after the addition of MAP fertilleer (Tables 4.5 and 4.6). Analytical methods for cation exchange capacity determination such as ammonium assesses acturation (pH>7.6) use high ammonium ion concentrations, similar to those found in the fartilleer treatments, for replacing other ions on the exchange complex.

The ammonium ions in solutions containing MAP fertilizer will affect the exchangeable cations on the exchange complex in a similar manner.

The most probable mechanism of cation exchange with high ammonium levels present in soil solutions containing MAP fertilizer is mass action. The exchange reactions move to completion as the ammonium ions are adsorbed onto the exchange complex thus releasing the exchangeable ions into solution.

4.3.2. Effect of Fertilizer Ammonium on the Soil Exchangeable Cations

Earlier discussion noted the cley mineralogy of both experimental soils is dominated by amerite. Smooths cley minerals have high CEC properties with pH-independent charge. The high exchange properties of ammonium would cause the ions on the exchange complex to be replaced with added fertilizer ammonium. Therefore, the exchange complex of amerite clay mineral may be expected to be the major source for cations entering soil solution.

An example may indicate the amount of exchangeable cation that could be released into soil solution through exchange with furtilizer ammenium. If the clay fraction contains 80% amounts and has a 20% moisture containt, ion exchange can account for an increase of 400 mmel Ca L⁻¹ or 800 meq L⁻¹. Therefore, ion exchange from sell day amounts can fully account for the ion concentrations found in extracted pure water before and after the addition of MAP furtilizer (Tables 4.5 and 4.8).

The exchange complex of the other clay minerals can also release cations into soil actual through exchange with ammonium. The lower CEC and pH-dependent charge of mice and itselfnite minerals, in addition to their lower content in the sell stay fraction, reduces the effect ammonium exchange has an releasing their advantad actions into sell solution. The change in pH-dependent charge on the exchange complex of exide and hydracide sell minerals, brought about by the decrease in sell pH through the addition of MAP fertilizer, may size result in the release of extient into sell calution.

4.3.3 Effect of the all of Solution Containing MAP Fertilizer on the Solubility of Soil Minerals

The low sH of solution containing MAP fertilizer increases the solutility of soil minerals. The increased solutility of soil minerals with the addition of MAP fertilizer may release ions into soil solution, by affecting both the exchange complex and the structure of the mineral surface. The solutilities of the soil minerals are different from each other and are important in affecting the release of ions into solution.

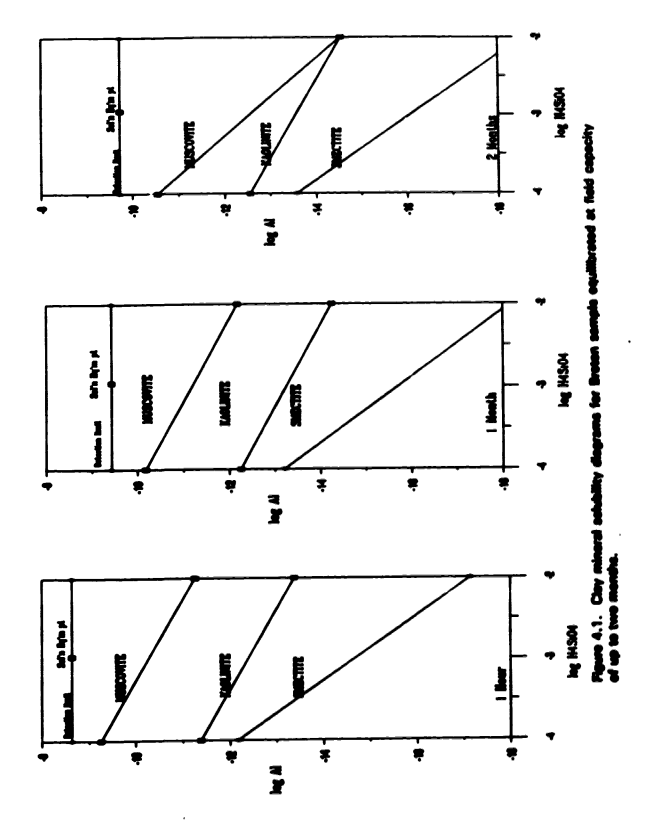
4.3.3.1. Oxide and Hydroxide Minerals

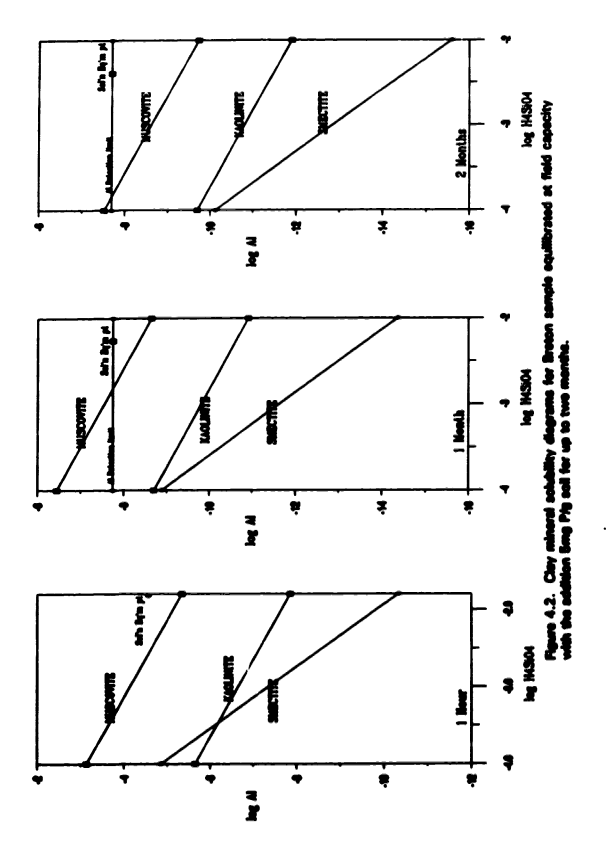
Onide and hydroxide minerals, in addition to their exchange complex, may release ions into sell solution through their dissolution. After the addition of MAP fertilizer, the pH of the sell solution decreases, causing the solutility of exide and hydroxide minerals to increase (Acquade and Tinelay, 1985; Schwertmann and Taylor, 1977). Therefore, the low pH conditions of the sell solution containing MAP fertilizer, may cause exide and hydroxide minerals to be sources for eleminum, iron and manageness ions.

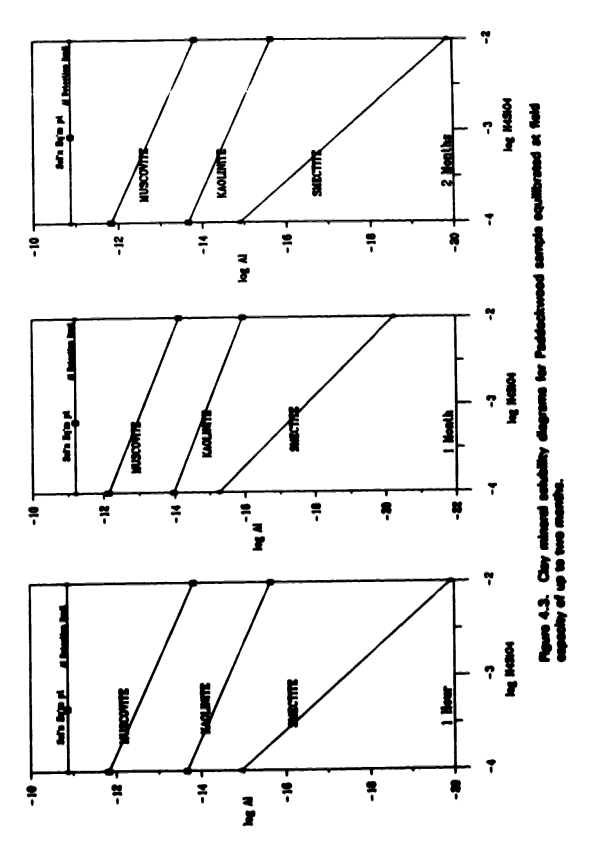
Oxide and hydrenide minerals amount to less than one per cent (mass) of the experimental sells (Table 4.1). However, the amount of AI, Fe and Mn in soil solution after the addition of MAP fertilizer is extremely low compared to the amount extracted from the sell by AO and DCS.

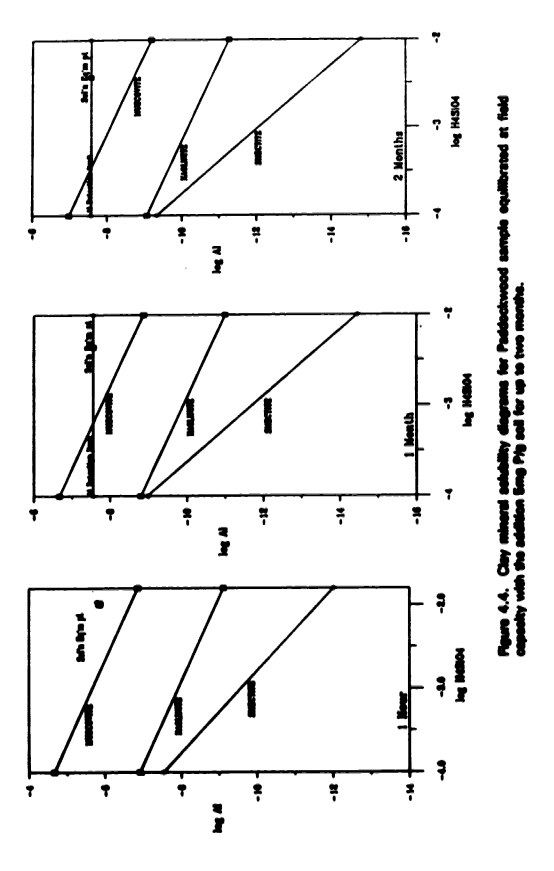
4.3.3.2 Clay Minerals

The solubility of secondary clay minerals is also affected by changes in pH. The equilibrium solubility status of clay minerals can be determined from sell solution data. Figures 4.1 to 4.4 depict on increase in the solubility of clay minerals after the addition of MAP fortilleer. The increase in AI concentration associated with the clay mineral leatines of Figures 4.2 and 4.4 after the addition of MAP fortilleer, one sell solution appears to be decreasing pH. One hour ofter the addition of MAP fortilleer, the sell solution appears to be









in equilibrium with muscovite, the most soluble clay mineral present (Figures 4.2s and 4.4s). The low AI ion concentration, undetected in all other pore water extractions both before and after the addition of MAP fertilizer, prevents the identification of the soil mineral phase in equilibrium with soil solution at that point in time.

Al, Si, Fe, Mg, Ne, O, H and K concentrations are directly affected by changes in the solubility of clay minerals in the Breton and Peddockwood samples. The ions associated with the mineral edges of muscovite are the mest directly affected by increased solubility of clay minerals. The ionic concentrations of Si, Fe, and Mn would also be in equilibrium with the soil clay minerals, however, we are unable to determine which clay mineral is in equilibrium with soil solution.

Exchangeable cotions and ion complexision properties are indirectly affected by increased solubility of clay minerals. The exchangeable cotions are the first to go into solution because they are retained at the surface of clay minerals. The exchange complex was proviously discussed as the major source of exchangeable cotions. As the exchangeable ions move into solution they undergo complexistion reactions with other ions. Therefore, even if the dissolution of clay minerals does not contribute a high concentration of ions to sell solution directly from the lettice structure, it does indirectly affect exchangeable cotion concentration and ion complexistion reactions.

4.3.4. States of Added Phosphones with Time after the Addition of MAP Facilitate

The large decrease in phosphate ion concentration in sell solution from one hour to two mentiles after the addition of the low rate of MAP fertilizer is the result of a complex set of reactions. Sell solution phosphate can be tied up in an array of P mineral precipitation and P adsorption reactions.

4.3.4.1 Relation of Soil Solution Phosphote and Colcium/Magnesium with Time

ions in soil solution decrease in concentration from one hour to one month after the addition of MAP fertilizer. The ion concentrations have appeared to stabilize somewhat by the end of the first month after the low level of MAP addition. Two cations, calcium and magnesium decrease in concentration to levels lower than the control treatment whereas all other ion concentrations decrease to levels alightly higher than the control treatment (Tables 4.5 and 4.6). The exchangeable cations would be expected to have higher concentrations after the addition of MAP fertilizer than prior to the addition of MAP fertilizer; assuming the exchange complex would become dominated by fertilizer ammonium. Therefore, there must be some additional reaction occurring that removes these two cations from solution.

Phosphete has the largest decrease in lank concentration in sell solution from one hour to one month after the addition of MAP fartilleer (Tables 4.5 and 4.6). The decrease in calcium ion concentration in sell solution caincides with the decrease in sell solution phosphete ion concentration. The decrease in calcium in sell solution deeples an exchange complex solurated with ammonium, suggests that calcium and phosphete may be precipitating to form calcium phosphete minerals. The decrease in calcium ion also provides an indication of the amount of P mineral formation taking place. With a decrease of 86 mmsl Ca L⁻¹ from one hour to two months after the addition of the low rate of MAP fertilleer (Tables 4.5 and 4.6), about 43 mmsl PO4 L⁻¹ could be tied up as dicalcium phosphete.

4.3.4.2 Relation of Soil Scholen Phosphote with Iron. Absoluter and Managemen Over Time

After the initial ion flush reculting from the addition of MAP fertilizer, the ione Fe, Mn, AI and SI also decreased in concentration (Tables 4.5 and 4.6) resulting from their decreased solutility at pH values above 5 (Acquiago and Tinaloy, 1986). Therefore, one may

anticipate the formation of sluminosilicate minerals. Unfortunately, the exceedingly low level of Al ion concentration does not allow us to observe if the clay minerals are superseturated.

Fe, Al, Mn and Si ions could also be precipitating as exide costs on surfaces of preexisting sell minerals. Amorphous allics accumulates as an intermediate in supersaturated solutions because it precipitates more readily than crystalline allics (Wilding et al, 1977). As can be seen from the day mineral solubility diagrams (Figures 4.1 to 4.4), the allics activity in sell solution moves towards amorphous allics equilibrium (pH₄SiO₄ = 3.1) supporting the possibility that amorphous allics components are forming.

Although ferritrydrite will form quickly if its reasonably high solubility product is exceeded, geothite is more likely to form at pH levels present in the fertilized sell solutions. Geothite fermation is faster and its proportion increases as the pH of the solution decreases from 7 to 4 (Schwertmann and Taylor, 1977).

The decrease in Fe, Al and Mn concentrations in soil solution after the addition of MAP fertilizer could be explained by the precipitation of Fe, Al and Mn phosphate minerals in addition to the precipitation of alumininealisate minerals. The decrease in ion concentrations of Fe, Al and Mn in the soil solution from one hour to one month after the addition of MAP fertilizer is very low in comperison to the decrease in Ca ion concentration for the same time period. This may lead one to assume that only a small amount of phosphate is reacting with Fe, Al and Mn. However, the extraction of pere water is a static measurement that connex determine soil solution dynamics. Therefore, Fe, Al and Mn may be continually released into soil solution with immediate reaction with added phosphate. Sell solution dynamics such as these would not be detected with pere water extraction. The maintaneous of high concentrations of Fe, Al and Mn in soil solution after the addition of the high rate of MAP fertilizer indicate the presence of a large source of these ions which may allow them to enter into soil solution form.

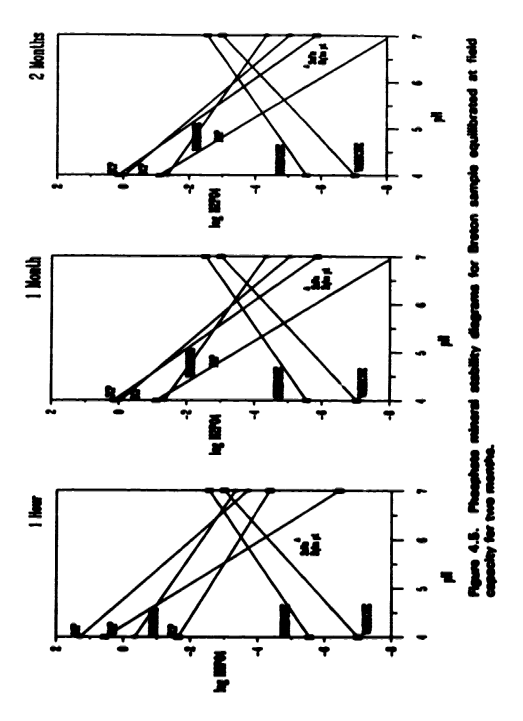
4.3.4.3 Phosphote Mineral Schubilly Disgrams

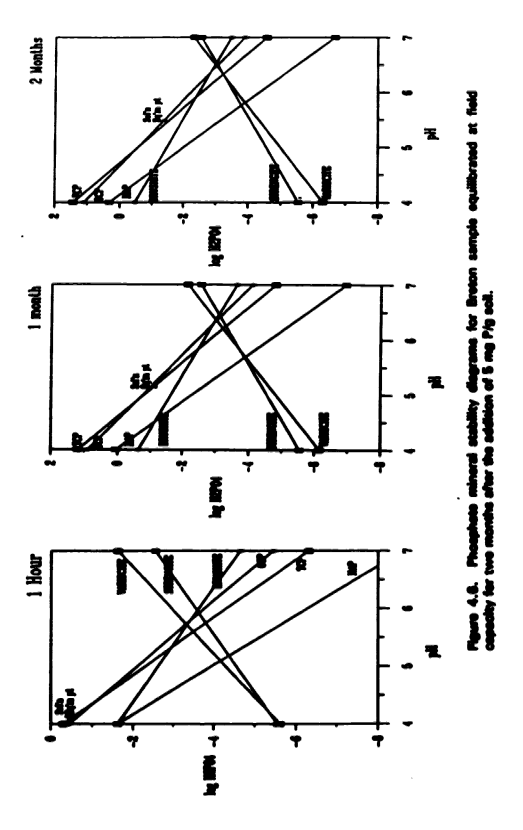
4.3.4.3.1 Calcium and Magnesium Phosphate Minerals

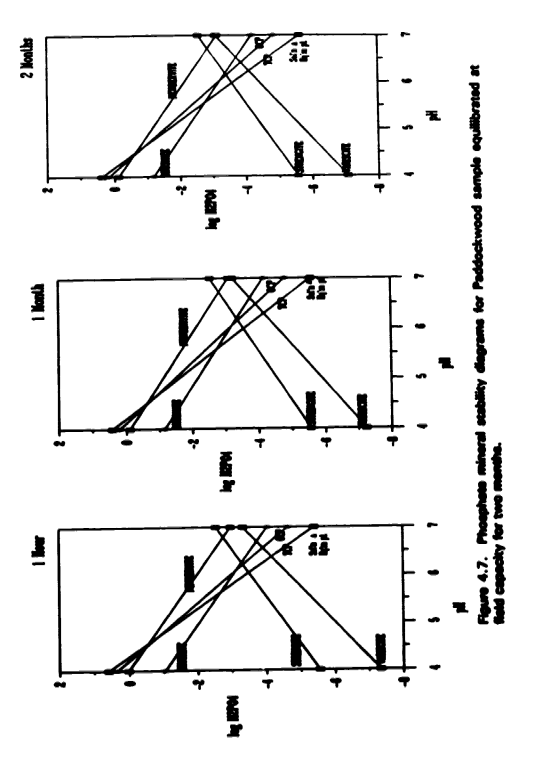
Calcium and magnesium ions precipitate with phosphote in supersaturated solutions (Abbons et al., 1986). The lower concentration of calcium and magnesium two months after the addition of MAP fertilizer compared to the control may possibly be related to the decrease in soil solution phosphote after the addition of MAP fertilizer (Tables 4.5 and 4.6). As discussed in 4.3.4.1, the precipitation of P with Ca or Mg cannot account for the total decrease in phosphote concentration. However, the precipitation of P minerals associated with calcium and magnesium is an important component of chemical reactions between sell and P fertilizer.

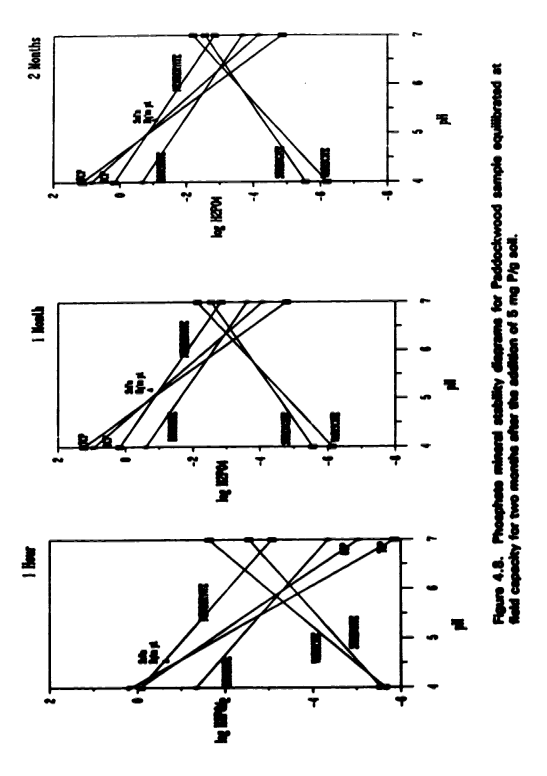
Solubility diagrams of P minerals depict the saturation index (SI) values of different P minerals in graphical form. As can be seen in Figures 4.5 to 4.8, calcium phosphate minerals are in equilibrium with phosphate levels in soil solution at all times after the addition of MAP farillier.

P minerals could not precipitate from solution within one hour of the addition of MAP fortilleer (Abbana et al, 1808). The adultity diagrams of P minerals depict SI values only and do not indicate P mineral precipitation. One hour after the addition of MAP fortilleer TCP and OCP are in equilibrium with sell solution (Figures 4.6 and 4.8), but probably have not precipitated out. However, the maintainance of equilibrium conditions between sell solution phaspitate with P mineral establishes for a two menth period, may suggest that TCP and OCP are precipitating out of sell solution. The relationship of P minerals to the sell solution phaspitate concentration of both samples indicate that phaspitate processes are occurring in each sample.









4.3.4.3.2 Iron, Aluminium and Manganese Phosphete Minerals

The presence of iron and aluminium P minerals have been documented within the zone where P fertilizer reacts with soil (Sample et al, 1980). P minerals of iron, aluminum and manganese are undersaturated prior to, and are supersaturated after the addition of MAP fertilizer in both soils (Figures 4.5 to 4.8). The supersaturated condition of the solution with respect to P minerals of iron, aluminum and manganese after the addition of MAP fertilizer indicates high precipitation potential. Therefore, Fe, Al and Mn P minerals are definitely a potential sink for added P.

4.3.4.4 Fertilizer History and P Mineral Solubility Disgrams

The similarity in the dynamics of the ion concentrations in soil solution is remarkable considering the P fertilizer history of the two soil meterisis. The solubility diagrams of P minerals are also similar for the two soils. The annual addition of P fertilizer to the Paddockwood sample site for the 36 years prior to sampling does not seem to have affected the manner in which the soil reacts with added P.

Based on the dynamics of ion concentrationsin soil solution, added P may be reacting with the soil constituents in the same manner for both soils. Added P may simply be entering the soil P cycles of both soils. Residual added P in the Paddockwood sample must have been dispersed through cultivation so as to not allow the build up of zones of high P concentration in the sell.

4.3.4.5 Advocation of Phosphorus

The precipitation of P minerals after the addition of MAP fortilleer could not possibly

account for the decrease in soil solution phosphate. The decrease in the amounts of calcium, magnesium, AI, Fe and Mn from soil solution after the high rate of addition of MAP fertilizer if tied up with phosphate could not account for the total decrease in the phosphate ion concentration over the same period. The only other option is that much of the phosphate has been removed from soil solution through its adsorption to soil minerals and organics.

4.3.5 Discussion of the Effects of Alfalfa Amendments on Soil Solution Values

The effects of the alfalfa amended to the soil samples during equilibration can mainly be attributed to alfalfa decomposition. The effects were mainly noticeable after one month of equilibration after which the soil solution values became similar to the non-alfalfa treatment (Table 4.6). The color of the soil solution may indicate the level of decomposition of alfalfa. After one month the pore water was very dark and contained high organic matter, whereas after two months of equilibration the pore water was the same yellow, clear color characteristic of the non-alfalfa treatment. The yellow color of the non-alfalfa treatment pore water is due to the solubilization of soil organic matter as a result of the low pH values (pH = 5.2). The dark color of the pore water after equilibration with alfalfa for one month is characteristic of organic matter dissolution through alkaline extraction. The pore water after equilibration with soil plus alfalfa for one month may represent the alkaline soluble fraction of organic matter. Therefore, the addition of MAP to soils high in organic matter may affect the alkaline aggregate stability.

Enriching the soil with an easily decomposible organic N compound with a low C:N (alfalfa meal) will always cause a rise in pH which is due to an accumulation of ammonia. The duration of such a pH increase will depend on the rate of alfalfa applied (Barnett and Dionne, 1980). The accumulated ammonia represents the quantity of substrate N in excess of microbial demand. The deamination of amino-acids with N mineralization releases ammonia which upon hydrolysis forms ammonium and increases the pH (Alexander, 1977).

The high pH of the pore water caused by excess ammonia must have undergone nitrification after extraction to account for the high nitrate values detected one month after the addition of MAP to the alfalfa treatment (Table 4.6). At high pH, nitrification occurs rapidly (Alexander, 1977).

Betremieux and Henin (1965) found that with the decomposition of luceme, the pH of the solution became slightly alkaline, and the iron content in solution decreased. They also noted the evolution of soil organic matter.

The lower iron and manganese values in soil solution one month after the addition of MAP to the alfalfa treatment, corresponds to the period of highest decomposition of alfalfa. Betremieux and Henin (1965) noted that the decomposition of lucerne decreases the solubility of iron and manganese.

The removal of phosphate from soil solution after MAP addition to soil with an alfalfa amendment can be attributed to both adsorption and immobilization. Immobilization may occur during the initial decomposition of the carbon source where the C:P is high (300:1), especially when the carbon source is high in N. The CEC of organic materials doubtes within 40 days of decomposition (Alexander, 1977); therefore CsHPO4 complexes may be adsorbed to the organics at the low pH after two months of equilibration.

4.4 Cheracterization of Phoenhorus in Fertilized Soil
4.4.1 Introduction

The mineral and organic components of soil are important factors that partially determine the fate of added P. Understanding the relationship of added P to mineral and organic soil components may compliment our previous discussion of soil solution dynamics after the addition of P familiaer. The importance of understanding the relationship of added P to mineral and organic soil components is identified by the small amount of P in sell solution. For example the phosphate in soil solution amounted to \$1% of Pt in the Breten

sample one hour after the addition of MAP fertilizer and decreased to 9% at two month. The phosphate in soil solution amounted to 47% of Pt

in the Paddockwood sample one hour after the addition of MAP fertilizer and decreased to 17% at two months. Therefore, to understand the fate of added P in the soil it is necessary to study the soil P distribution in addition to soil solution dynamics.

The Pt of each soil was analyzed by a sequential extraction procedure (O'Helloran ex al, 1967). The sequential extraction procedure attempts to duplicate the effect a growing plant would have on the system by repeatedly depleating the soil solution of Pi. This technique isolates the various components of the soil P cycle according to their bioevallability (Roberts and Stewart, 1967).

The extractants involved in this fractionation procedure are: anion exchange resin (pH 8.2), 0.8M NeHCO3 (pH 8.2), 0.1M NeOH (pH 8.0), 1.0M HCI (pH 3.0) and $\rm H_2SO_4$ - $\rm H_2O_2$ digestion. The sequential extraction precedure divides soil Pt into either five or seven fractions; two of the extractable fractions, 0.8M NeHCO3 and 0.1M NeOH can be divided further into inorganic and organic components.

In the fractionation procedure each extractant is responsible for extracting different fractions of soil P (Roberts and Stewert, 1987). Resin-extractable P (Resin-P) is the most plant evallable form of PI; it consists of the P minerals brushles and OCP, and also the PI that is adsorbed to the surface of assignishides and carbonases. NeHCO₃ extractable P (Bloerb-P) is considered to consist of readily plant evallable PI, alightly less evaluable forms of the same components extracted by Resin-P and of easily mineralizable Po traction. NeOH entractable P (NeOH-P) removes both the PI component associated with amorphous and crystalline AI and Po, and the Po component associated with stable organic forms. The HCI entractable P (NCI-P) consists of stable Co-bound P, youthly characterized by NAP, Pinally,

acid digestion extactable P (Residual-P) consists of P associated with stable humus and humic acid and of insoluble PI forms.

4.4.2. Changes in the Fractions of Phosphorous After the Addition of MAP Fartilizer and Subsequent Equilibration

4.4.2.1. Introduction

The sequential extraction technique was conducted on the equilibration samples that soil pure water was extracted from. After complete characterization of soil P fractions, comparison with pure water data may increase the understanding of soil-fertilizer reactions.

The Resin-P fraction generally decreased with time after MAP fertilizer was applied to the soil samples in the equilibration study. The decrease in Resin-P was complimented by increases in other less available P fractions in both Breson and Paddockwood samples. This decrease in Resin-P with time after the addition of MAP fertilizer may be related to the decrease in P concentration in soil solution. As the phosphate concentration in soil solution decreased, there was a corresponding decrease in the amount soil solution phosphate in Resin-P. Also, decreases in soil solution phosphate were generally greater than decreases in Resin-P. Therefore, phosphate that was leaving soil solution with becoming transformed to other soil P fractions in addition to solid forms of Resin-P.

4.4.2.2. Changes in the P Fractions of Brosse Sell After the Addition of MAP Fortilless

The sell P fractions were menitered one hour, one month and two menths after the

addition of MAP fertilizer to the Breton samples. The continual decrease of Resin-P over the two month time period was matched by simutaneous increases of Bicarb-Pi, NeOH-Pi,NeOH-Po and HCI-P (Table 4.8). The highest increases were in the Bicarb-Pi and NeOH-Po fractions. The P associated with the Bicarb-Pi fraction remains readily available, while the P associated with NeOH-Po is more stable (Roberts and Stewart, 1987). A major portion of added P also occurs as relatively insoluble calcium phosphate minerals associated with HCI-P.

The amount of soll solution P in the Resin-P fraction after the addition of MAP fertilizer to the Breton sample decreases with time. For example, one hour after the addition of MAP fertilizer, soil solution phosphete virtually equals Resin-P (Table 4.9). However, one month later soil solution phosphete accounts for 32% of Resin-P; and at two months this value is 19%. Over the two month duration of the equilibration experiment, the amount of soil solution phosphete in the Breton sample decreased by about 3200 μ g P g⁻¹ (calculated from Table 4.5) soil while the amount of Resin-P decreased by about 1300 μ g P g⁻¹ soil (Table 4.9).

4.4.2.3. Changes in the P Fractions of Breson Soil With Alfalfa Amendment After the Addition of MAP Facilities

When alfalfa was added to the Broton sample in addition to MAP fertilizer, the P transformations between soil P fractions during equilibration were similar to treatments without alfalfa addition. However, the amount of P undergoing transformations was much lower with the alfalfa amendment then without it. More P was received in Resin-P fraction, and lower levels of NaCH-P and HCI-P were assessed.

When alfalfs and MAP fartillar were added to the Broton comple, the amount of sell solution phosphote in Resin-P strengly decreased with time (Table 4.5). For example one

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TABLE 4.9 Soil solution phosphate expressed as a percentage of the Resin-P fraction after the addition of the low rate of MAP fertilizer

Sample	lhr	30 days	60 days
Breton	90	32	19
Breton with		_	
errerre	60	6	5
Paddockwood	76	66	28

hour after the addition of MAP fertilizer, soil solution phosphate amounts to 60% of Resin-P; after one month this value is 6% and after two months this value is 5%. Over the two month duration of the equilibration experiment, the amount of soil solution phosphate in the Breton sample with alfalfa amendment decreased by about 1674 μ g P g⁻¹ soil while the amount of Resin-P decreased by about 522 μ g P g⁻¹ soil (Table 4.9).

4.4.2.4 Changes in P Fractions of Paddockwood Soil After the Addition of MAP Fartilizer

During the equilibration experiment, the transformations between P fractions are also similar for the Paddockwood sample and the Breton sample. Although a few differences do exist. For example, there is no measured decrease in Resin-P, however the soil solution phosphate concentration decreases with time.

The amount of soil solution phosphete in Resin-P after the addition of MAP fertilizer to the Paddockwood sample also decreases with time (Table 4.9). For example one hour after the addition of MAP fertilizer, soil solution phosphete amounts to 76% of Resin-P; after one month this value is 86% and after two months this value is 28%. Over the two month duration of the equilibration experiment, the amount of soil solution phosphete in the Paddockwood sample decreased by about 1426 μ g P g⁻¹ soil, about one half of the Breton value (Table 4.9). A decrease of Resin-P over this period is not detected.

4.4.3. Discussion of the Changes in P Fractions After the Addition of MAP Familiar.

4.4.3.1 Introduction

Transformations of P between fractions after the addition of MAP furtillar fellow

similar pathways between soils. P transformations were larger for the Breton sample than for the Paddockwood sample. The P transformations for the Breton sample with the alfalfa amendment were of a similar scale to the Paddockwood sample.

P added to the soil generally moves from plant available forms (Resin-P) to less readily available forms. P moves into most forms directly from soil solution. Soil solution phosphate undergoes both precipitation and adsorption reactions and remains in the Resin-P fraction. However, the majority of the decrease in Resin-P probably occurs as direct movement of soil solution P to other P fractions through precipitation, adsorption, and chemi-corption.

Most of the P transformations within two months after the addition of MAP fertilizer is from Resin-P to Bicarb-Pi. 50% of added P in the Breton sample and 70% of added P in the Paddockwood sample remains in Resin-P and Bicarb-Pi fractions. Wager et al (1986) found that after both five to eight years of addition of P fertilizer to two Seekstchewen soils, 50% of fertilizer P remaining in the soil existed in the Resin-P and Bicarb-Pi fractions.

4.4.3.2 Discussion of Changes in P Fractions After the Addition of MAP Facilities to the Broton Semale

The movement of P out of soil solution after the addition of MAP fertilizer to the Breton comple indicates a decrease in available P forms. The P can move from soil solution to all sell P frections. Some will remain as Resin-P and the rest will move directly from soil solution to the loss available fractions.

The majority of added P in the Breton sample remains in the readily evaluable Resin-P fraction for at least two months. Resin-P contains sell solution P, soluble P mineral

precipitates and adecreed Pi. Therefore, most of the added P remains available for plant uptake for the two months.

Added P in the Breton samples also moves from soil solution to Bicarb-P through precipitation and adsorption reactions. Bicarb-P contains slightly less available forms of P than Resin-P. The Bicarb-P in the Breton soil will then serve as a labile P source to Resin-P.

The increase of NeOH-Po in the Breton samples after the addition of MAP fertilizer is the result of complex reactions. The Breton sample has low carbon content, so the generation of 600 μg P g^{-1} soil is unlikely. Therefore, the increase in NeOH-Po is probably not actual Po. The NeOH-Po increase is probably the result of selective adsorption of P to stable organic compounds associated with iron and manganese (A polt et al, 1975).

The increase of HCI-P in the Breton samples after the addition of MAP fertilizer may result from P mineral precipitation with a subsequent increase in pH. Calcium phosphete minerals are more soluble at the pH levels found in soil solution just after the addition of MAP fertilizer. The increase in soil solution pH with time decreases the solubility of the precipitated calcium phosphete minerals. Therefore with the increase in pH of soil solution, calcium phosphete minerals may move from Resin-P to HCI-P because of the change in solubility.

4.4.3.3 Discussion of Changes in P Fractions After the Addition of MAP Facilities and Alfalia Assessment to the Breton Sessale

The transformations between the P fractions of the Broton sample with the alfalla amendment are occurring for similar reasons as in the Broton sample without the alfalla amendment. Even the increase in NeCH-Pe is similar to that of the non-citalla treatment and is therefore thely the result of adsorption of P to Ai, Mn and Pe organics.

4.4.3.4 Discussion of Changes in P. Fractions After the Addition of MAP Fartilizer to the Paddockwood Semale

After the addition of MAP fertilizer to the Paddockwood samples, the movement of P out of soil solution is not associated with a decrease is Resin-P. The low decrease in Resin-P may result from P mineral precipitation and from phosphete adsorption to associated nodules that are part of this fraction. The P source for increases in Bicart-Pi and NaOH-Pi may be undetected decreases in Resin-P.

The prior addition of P fertilizer to the Paddockwood sample has probably caused an adjustment in the steady state conditions of the soil P cycle. Therefore, higher levels of P would be associated with all components of this soil P cycle (McKenzie, 1989). The high P content of each component of the soil P cycle may reduce the movement of added P away from the most readily plant available form. Therefore, Paddockwood sample Resin-P values may remain nearly constant with time.

4.4.4. Distribution of Added Phosphorus After Barley Growth

4.4.4.1 Introduction

Analysis of P distri-ution within fractions after orap growth may lead to an understanding of the role these different P fractions have in uptake of P. After barely was grown in greenhouse self samples for three menths, added P was observed by three methods. The plant meterial was analyzed for P uptake, self P fractions were analyzed and soil thin sessions were analyzed for direct observation of P distribution.

P abundance decreased with distance from the farillies granule until, at 2 - 3 cm, it

was similar to the control treatment. The P increases near the fertilizer granule were distributed among the P fractions at similar ratios to the equilibrium experiment. The similarity between the greenhouse sample P fractions and the equilibration sample P fractions indicate the same P transformation processes were occurring in the soils of each experiment.

The amount of P involved in the P fractions of the greenhouse samples is lower than the equilibration samples, so comparisons will be dealt with on a relative scale. Distribution within fractions of the greenhouse samples is generally similar to the equilibration samples. Regin-P fractions account for relatively the same amounts of P as the equilibrated treatments in both Breton and Paddockwood samples. The Bicarb-P in greenhouse samples is generally lower, possibly reflecting P uptake by plants.

Large amounts of NeOH-P were found in Breton and Paddockwood samples under both experimental conditions. NeOH-P accounted for approximately 20% of Paddockwood Pt and 33% of Breton Pt in the greenhouse soil samples. Wager et al (1986) measured relatively small amounts of fertilizer P recoverable by NeOH in Waskada soils whereas NeOH-P accounted for 36% of fertilizer P remaining in a Sutherland soil after five years. Therefore, values obtained in this experiment are similar to those values attained after long term fertilization.

The similarities of P distribution in soil after long term P fertilizer addition and our short term experiment may mean P transformation pathways between fractions are similar for both time periods. Therefore, the soil P cycle must be similar for both soils, with different steady state conditions existing for each sample.

4.4.4.2 Distribution of Added P After Berley Growth on Breton Semples

After three months of beriev growth, approximately 50% of the P remaining in Breton samples is retained in readily available Resin-P and Bicarb-P forms (Table 4.10). This is similar to values found in the equilibrated samples and to values found by Wager et al (1986). Therefore, except for the Po returned to the soil in crop residue forms, the P cycle may reach steady state conditions by the end of the growing sesson.

4.4.4.3 Distribution of Added P After Barley Growth on Paddockwood Samples

After three months of barley growth 75% of the P remaining in Paddockwood samples is retained in readily available Resin-P and Bicarb-P forms. This may reflect the low decrease in Resin-P in the Paddockwood samples of the equilibration study.

4.4.5 Discussion of the Changes in P Fractions After Barley Growth

The non-replication of P fractionation analysis of the greenhouse samples does not allow for statistical analysis. Therefore, definite statements about P transformation processes cannot be made. However, general observations depict similar effects happening near the fertilizer polists in the greenhouse samples as were occurring in the equilibration samples.

The only noticeable effect that the growth of barley may have had on the P distribution among soil fractions is a decrease in labile P (Blearb-P). This may indicate that as plants take up phosphate, it may move from Bloarb-P form to Resin-P form.

TABLE 4.10. Greenhouse soil P fractions after P addition and 3 months of barley growth.

			Size of	YAP feetilizer neile	(0)				
		0.01			0.10				
	Distance of soil sample from fertilizer peliet (cm)								
P.Fraction	0-1	1-2	2-3	0-1	1-2	2-3			
				μg P/g soil					
				BRETON					
RESIN-P	58	3 3	1	493	333	48			
NaHCO3-P	52		0	147	138	ŏ			
NaOH-P	114	92	0	391	251	Ŏ			
HCI-P	0_	68	20	146	59	50			
RESIDUAL	26	23	0	5	15	0			
Total-P	250	189	21	1182	796	98			
				PADDOCKWO	00				
RESIN-P	354	345	0	649	450	41			
NaHCO3-P	92	92	0	220	102	15			
NaOH-P	98	120	0	236	85	õ			
HCI-P	84	84	8	34	44	š			
RESIDUAL	9	0	0	37	3	3			
l'otal-P	637	641	8	1176	684	59			

4.4.6. Forms and Distribution of Added Phosphorus as Seen in Soil Thin Sections

Thin sections of the undisturbed soil near the fertilizer granule within the greenhouse experiment soil samples give an excellent depiction of the soil environment including products of reactions between the soil and fertilizer P. Both P adsorption to soil minerals and P precipitation products are clearly shown. The locations and distribution of areas of high P concentration within the soil fabric are also clearly evident. The evidence is perticularly good for the Paddockwood samples. Although thin section evidence has not been found, similar reactions are most certainly occurring in the Breton samples, based on similarities in P mineral solubility diagrams and en P fractionation values (Figures 4.5 and 4.6 and Tables 4.8 and 4.10).

P levels in the soil thin sections were measured semi-quantitatively by EDX. Feirly high quantities of P must be present because of low detection limits of EDX. Much of the distribution of added P that is present in low concentrations in the soil, could not be detected.

The concentration of adecreed P in % per g soil would normally be so low that it would not be detected by the EDX. However, the P on the surface of an iron/mengenese nodule in Plate 4.2 has concentration high enough to be detected.

P mineral cryotals have been found in sell thin sections within a 1cm radius of 0.10g MAP granules and within a 1.5cm radius of 0.50g MAP granules. The P mineral cryotals are enstualvely associated with calcium ions. These calcium phosphota minerals ensur in valds adjacent to sell minerals (Plates 4.3 to 4.5).

Plate 4.2. Depictions of nodules in Paddockwood sample thin sections near the application site of MAP fertilizer: (a) micrograph in plane light (frame length = 3.5mm); (b) SEM scan of the same area





Place 4.2. continued: (c) EDX map showing the nodules to be of iron and manganese oxides; (d) showing P association with the nodules and not with calcium.





Plete 4.3. Depictions of nodules in Paddockwood sample thin sections near the application site of MAP fertilizer: (a) The association of calcium with added phosphorus; (b) calcium phosphate accumulations in voids of the sell matrix;

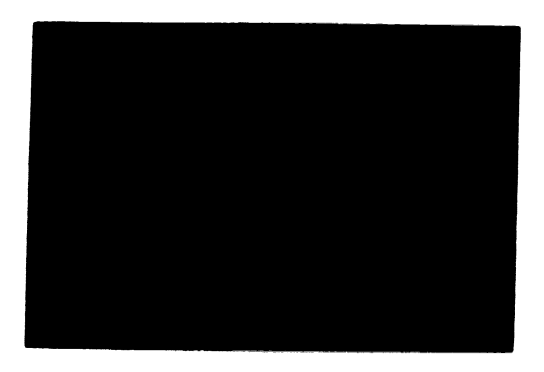




Place 4.3, certificand: (c) calcium. phosphate accumulations in voids left by the fertilizer



Plate 4.4. Micrographs showing calcium pheephate minerals adjacent to soil fabric in valds left by the granule of MAP fertilleer: (a) plane light (frame length - 3.5mm); (b) with filters under cross polarized light (frame length - 3.5mm).





Place 4.5. Micrographs showing calcium phosphate minerals adjacent to soil minerals in valde: (a) in plane light showing distribution in the soil matrix (frame length = 7.0mm); (b) with filters under cross polarised light (frame length = 7.0mm);

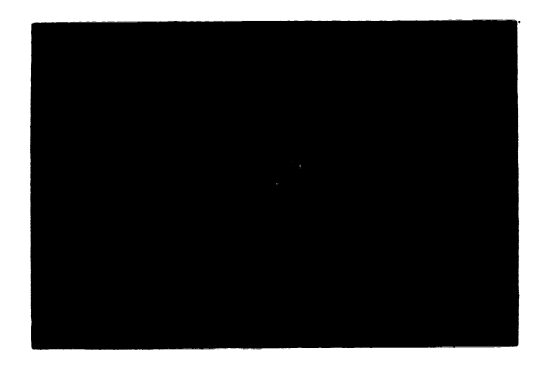




Plate 4.5. continued: (c) in plane light (); (d) with filters under cross polarized light (frame length = 0.5 mm).





The most common calcium phosphete minerals appear to be OCP and Brushite. OCP crystallizes as thin blades arranged in a radial pattern (Plate 4.4) (Lahr et al, 1967). Brushite is present as a larger crystal with a brownish color in plain light with crystal habits that resemble gypoum (Plate 4.5). These OCP and Brushite minerals are located in vaids left by the MAP fertilizer granule and in packing vaids further away from the granule.

4.4.7. Discussion of Thin Section Evidence

The thin section evidence clearly showed detectable P accumulations to be more abundant in the Peddockwood sell samples then in the Breton sell samples. As discussed earlier, the Peddockwood site has had a previous history of P fertilizer addition whereas the Breton site has not. So the more abundant P accumulations of the Peddockwood sample thin sections may be a result of either the previous fertilizer history of the sell or the sell properties themselves.

Thin sections from the Peddockwood central comple, that was not treated with MAP fartilizer in the greenhouse, did not provide any evidence of P accumulation. So the P accumulations in thin sections of Peddockwood complex after barley growth and fartilizer addition could be artifluted to the MAP fartilizer addition. All the P cryotals found were located near the fartilizer granule aloss (Plate 4.1). P accumulation onto accoulantific neglules was also limited to the neglules located near the fartilizer granules (Plate 4.2). The large amount of P fartilizer that has accumulated in the sell over the years has probably generated nuclei that increase the rate of P mineral precipitation, thus enabling factor cryotal farmation in the Peddockwood comple.

The correlation between the distribution of fractions in the greenhouse and equilibration complex is very good. A good relationship also orders between the self thin

sections, taken from the greenhouse samples, and the P mineral solubility diagrams derived from equilibration soil data.

Although OCP, TCP, Brushite and Newberyite all seem to be in equilibrium with Paddockwood soil solution (Figure 4.6), OCP and Brushite are the dominant P minerals present. The presence of Brushite and OCP minerals vertiles the validity of the P mineral solubility diagrams. These Brushite and OCP minerals must be controlling the soil solution phosphate levels in soil. If they control soil solution phosphate levels in soil. If they control soil solution phosphates to the Resin-P fraction.

As the sell pH increases the Brushite and OCP minerals will become less soluble (Figure 4.6 and 4.8). The Resin-P fraction will decrease as the Brushite and OCP minerals become more stable and revert to the HCI-P fraction.

The P accumulated on the surface of the sesquioxidic nodule in Plate 4.2 has moved 1.5 to 2 cm from a 0.50g MAP pellot. The sell that the added P moves through is a pecked greenhouse sell with fabric as described in Table 4.11. Therefore, a large amount of added P is moving through the sell. These quantities and distances are similar to those found by Bell and Black (1970b).

The P accumulated on the surface of the seequioxidic nodules may be either attached through anion adverption or precipitated as surface coats of Fe or Mn phosphote minerals. The potential for the precipitation of Fe and Mn P minerals is evident from the self solution date. Iron phosphote minerals are more stable than iron evides in self. The potential for phosphote adverption to the acceptantide nodule is also high because the low pH will give rise to high anion exchange expectly because of the pH dependent charge of the evides. Regardless, the relationship between P and the acceptantide nodules increase the NaCH-P dependent.

TABLE 4.11. Hieramorphological description of potted greenhouse soil samples at depths from 1 to 6 cm.

<u>Paddockwood</u>

lemence:

Granic

Type:

Matri-granoidic porphyric//porphyric

Description: Deminant fabric type is matri-granoidic perphyric with common matri-perphyric fabric. Size of granoidic units varies from 0.5mm to 2.0mm with an average of 1.0mm diameter. Granoidic units are composed of soil plasma and frequent quarts and foldopar grains. Pew sense of iunctic perphyric fabric are also present. Voids are comprised of crase planes (15%) and fine to medium, irregular, ortho vughs (10%) and comprise of 25%/unit volume of soil. Packing voids are also present in a few areas.

Padalesies)

<u>factures</u> (i) Few, discrete, opherical, cooquiexidic modules with charp boundaries and undifferentiated fabric. They range in size from lmm to Jmm in diameter. (ii) Very few opherulitie, phosphitic crystals within voide, adjacent to coil matrix.

Plassa fabric: Deminant fabric type is silasopic. Hodulos also have silasopic fabric.

Bretes

Secretors:

Granic

Dens.

Matri-granoidic perphyric

<u>Restrictions</u> Deminent fabric type is matri-granoidis perphyris with frequent matri-granoidis and innatis-perphyris fabric. Since of granoidis units very from 0.4mm to 1.5mm with average of 0.8mm diameter. The granoidis units are composed of soil plasma with frequent quarts and foldaper grains. Somes of frequentials perphyris fabric are also present. Voids comprise 30% of soil volume and 30 to 50% above the cood row. Fine to medium irregular ortho waghs are frequent with a few interpodal most planes.

Problemies

<u>Sections</u>

(i) Common discrete spherical esequienties (9%) and manganiferous (9%) nodules with charp boundary and lamaller febrie. They range in size from 0.2mm to 1.0mm with an average of 0.8mm diameter. (ii) Very few discrete, restangular, single ortho-approtubule (2.8mm).

Places fabric: Deminent fabric type is allesspie. Hodulos are accepts. Approtubulos are allesspie.

4.5. Granule Size and Bend Placement of P Fertilizer

The granule size and bend placement of MAP fertilizer were both varied in the greenhouse study. These two factors were studied to develop and understanding of the utilization efficiency of P fertilizer. Also, granule size and bend placement are the two factors that could most easily be adjusted to increase the efficiency of P fertilizer on a practical level.

4.5.1 Effects of the Granule Size of MAP Fertilizer on Phosphorus Untake and Berley Growth

Different treatments were tested to determine how P fixation varies with fertilizer granule size. In the Breton samples the distribution between soil P fraction of each pellet size treatment varied from the P distribution within the equilibration treatment. For example, the 0.10g pellet treatment had relatively lower Bicarb-P and the 0.01g pellet treatment had relatively lower Realn-P (Table 4.10).

In the Paddackwood samples, the distribution between soil P fractions for both - granule size treatments exhibit trands similar to that for the equilibration soil treatments. The majority of the added P remained in the Resin-P form (Table 4.10). Significant increases were determined for the Bicarb-P and NaOH-P fractions. In addition, there was no difference in the HCI-P and Residual-P fractions.

The efficiency of uptake by barley from each granule size of MAP fertilizer was measured. The solution form of MAP had higher P uptake from the Broton soil sample after one menth then other granule size treatments (Table 4.12). This treatment probably had the greatest distribution of P and was therefore the most available for the early growth of

TABLE 4.12. Uptake of added 32 P by barley from Breton and Paddockwood soils for pellet size, pellet placement and organic amendment treatments.

Form of	Harvest these periods (days)						
	BRETON			PADO	<u> </u>		
fertilizer MAP	30		90			90	
Solution P ^{As} 0.01g ⁴ 0.10g ⁴ 0.50g ⁴ Straw ⁴ 0.01g ⁴	2.8	5.9	12.3	2.1	5.4	12.3	
0.0125	2.5	5.8	11.5	2.0	6.1	11.9	
0.10gs	2.5 2.5	6.1	11.9	2.3	6.8	12.6	
0.50g ⁶	1.5	6.8	12.3	1.7	6.6	10.9	
Straw	2.2	6.3	12.1	2.0	7.2	13.3	
Post ⁵ .	2.3	7.1	14.2	2.7	6.8	13.8	
0.01g ⁴	0.7	4.9	8.0				
0.10g	0.9	5.5	9.2				
0.50g4	0.2	3.9	8.0				

- Partition MAP was added at a rate of 20 kg/ hs, and K2904 and NEI4NO3 were added at rates of 20 and 50 kg/hs, respectively.

 Partition was disselved water and added as a band.

 Partition was banded 10 cm below the seed sow.

 Fertilizer was banded 2cm below and 2cm beside seed row.

Significance levels for various treatment contrasts from analysis of variance of uptake of 32 P by barley from Broton and Paddockwood grounbouse self samples.

		Hereset time periods (days)					
Orthogonal Contrasts	MOSTON M			PADOO	90		
soin vs granuler lo,hi vs mid 0.01g vs 0.30g 0.1g vs 0.1g + org Post vs Strw	0.005 0.05 0.001 M	**	0.005 0.005	0.005 10 10 0.005	0.02 M M M	0.0001 0.0001 0.0001	
Nonorthogonal Contrate				·			
6.615 w 6.105 6.505 w 6.105 day w deller 6.615 w 6.105 6.505 w 6.105 6.615 w 6.505	0.0002 0.005 m 0.62 0.65	20 0.01 0.05	0.01 0.01	es 0.005	*	*	

plants.

The large 0.50g pellet had significantly lower uptake of added P after the first month of barley growth on both soil samples then the other granule size treatments (Table 4.15). The narrow distribution of MAP fertilizer in this treatment was probably responsible for the low uptake of added P. By the end of the second month the P uptake from the 0.50g granule size treatment was similar to treatments of other granule sizes. By the end of the third month the uptake of added P from the 0.50g granule size treatment was the same as the other granule size treatments for the Breton sample but was significantly higher than the other granule size treatments for the Paddockwood sample.

4.5.2. Discussion of the Effect of Granula Size of MAP Fertilizer on Phosphorus Untake and Barley Growth

The similarity of the distribution between soil P fractions of different granule size treatments indicate the same reactions are occurring near the fertilizer pallet regardless of the size. One might expect, that with a larger surface area, more P of the smaller size treatment would become fixed. However, since the majority of added P remains in the plant available Resin-P form, no difference in grap P uptake is detected between granule size treatments.

P uptake by barley was generally unaffected by the granule size of MAP fertilizer. Differences that axisted after the first menth were gene after the second menth in most cases. The lewering of the difference in uptake of added P from the sells shows the buffering effect the sell P cycle has. The colution form of MAP had significantly higher uptake of added P from the Broton cample after one menth. This is probably the result of the greater initial P distribution resulting from solution application. The absence of any

significant increase after the second month probably results from the fact that in the other treatments, rooting located P sources and thus sufficient P was taken up.

4.5.3. Effects of Fertilizer Band Placement on P Untake and Barley Growth

The P uptake of deep banded treatments on the Breton soil was also uneffected by MAP granule size (Table 4.12). The larger granule size treatment had lower uptake of added P after one and two months, for the same reasons as this granule size treatment had lower uptake in the shallow band treatments. The poor distribution of P lowers the probability of the plant roots from coming into contact with added P (Sander and Egbell, 1968).

The deep bended MAP pellet treatments had significantly lower uptake of added P then the shellow bended treatments (Table 4.12). In addition to the low P uptake, barley yields were significantly lower (Table 4.13). Assuming that the distribution between soil P fractions in the soil near the deep bended pellets is similar to that near the of the shellow bended treatments, the only explanation for low P uptake and low yields must be the plant root not finding the P source soon enough. Roots definitely reach the P source, as evident in the P uptake values (Table 4.12), but clearly the plant is not able to shearb enough.

4.5.5. Effect of Orașeic Amendment on Untake of P and Barley Growth

The uptake of added P from the treatments with organic amendments was significantly higher than without organic amendment in both Breton and Paddeckwood samples. The past amendment treatment also had significantly higher uptake of added P than the strew amendment. The higher uptake of added P from the treatments with organic amendments may be a result of lower contact between added P and the sell. The barley yields were not significantly greater despite the greater uptake of added P.

TABLE 4.13. Burley harvest weights of dry shoot material from Braton and Paddockwood soils for pellet size, pellet placement and organic amendment treatments.

Form of	Harvest time periods (days)							
fortilizer	BRETON			PADDOCKWOOD				
MAP [®]	30		99	30		90		
	:			ot marcial/po	<u> </u>			
Control [®]	1.83	9.34	26.51	0.61	3.60	16.90		
Solution P ^{Ps}	2.54	10.48	30.16	1.73	9.41	25.16		
0.01g ² 0.10g ² 0.30g ² Name	2.50	11.13	29.94	1.55	8.96	24.71		
). 10 2 3	2.51	11.83	29.25	1.56	9.00	24.90		
1.50 ²³	2.51	11.45	30.86	1.35	8.91	24.39		
	2.43	11.16	20.51	1.49	9.58	24.49		
	2.34	10.96	29.52	1.83	9.63	25.87		
LOLE!	1.06	9.20	25.83		,,,,,	-5.01		
	1.24	8.80	24.85					
1.00 C 1.10 C 1.50 C	1.05	7.20	25.20					

Postliner MAP was added at a rate of 20 kg/ ha, and 123/04 and NEHHO3 were added at a rate of 20 and 30 kg/ha, respectively.
 Postliner was dissolved vester and added as a band.
 Postliner was banded 10 on below the cool row.
 Postliner was banded 20m below and 20m baside cool row.

Significance levels for various treatment contrasts from analysis of variouss of barley barvest dry weights greenhouse experiment.

					Herent time periods (down)			
		M	PADDO	PADDOCKWOOD				
		<u> </u>			99			
0.005	0.02	0.05	0.0001	0.0001	0.0001			
•	-	*	*		-			
•	**		₩.	•	-			
	_	_		-	-			
_	_		8	_	-			
-	-	-	4.60	•	-			
	<u>. </u>							
		-	-					
*	-		0.05	=	=			
0.005	0.01	0.01						
0.01	-	-						
		•						
	-	0.005 0.02 0.005 0.02 0.005 0.01 0.005 0.01 0.005 0.01 0.01 00	0.005 0.02 0.05	0.005 0.02 0.05 0.0001 0.005 0.02 0.05 0.0001 0.005 0.01 0.01 0.01 0.01 0.01 0.01 0.01	0.005 0.02 0.05 0.0001 0.0001			

4.5.6. Discussion of the Effect of Organic Amendments on Uptake of Phosphorus and Barley Growth

Putting the fertilizer pellet in a volume of peet/straw may have lowered the dissolution rate of the MAP granules. The organic material would have a lower matrix potential than the surrounding soil at field capacity causing water to move towards the organic volume. Therefore, the added P would only move out of the organic volume through diffusion against the current. This would lower the contact of added P with the soil and thus reduce the occurrence of reactions between soil and MAP fertilizer. Once a plant root finds the source of P there would be more P available. thus allowing for higher P uptake.

There was no difference in barley yield despite the higher uptake of P from the treatments with post amendment. Therefore, plants may save this excess P for grain fill.

5.0 Conclusion

Based on the results and observations of the studies of this thesis the following - conclusions were drawn:

- 1) The reaction between sell and MAP fertilizer results in the release of cotions into sell solution. The exchangeable cotions are released from sell minerals through exchange with fertilizer ammonium. The law pH of sell solution containing MAP fertilizer initiates dissolution of sell slay minerals and of exides and hydrexides. The dissolution of sell minerals may also release substantial amounts of ions into sell solution.
- 2) Added phosphate reacts with the ions that are released into sell solution. The reactions

of added phosphate with soil ions result in the production of P mineral precipitates. The calcium phosphate minerals Brushles, OCP and TCP form adjacent to soil minerals in voids near the application site of MAP fertilizer granules. Iron and manganese phosphate minerals precipitate on the surface of existing seequiexidic nodules in the soil matrix.

- 3) Added P moves from readily available to less available soil P fractions with time after the addition of MAP fertilizer. However, near the application also of MAP fertilizer, the majority of P will remain in the Resin-P form even though the soil solution phosphate concentration decreases over this time period. The P that leaves the readily available form becomes adeorbed to inorganic and organic forms of iron, aluminium, and manganese; both components of the NeOH-P soil fraction. As the pH in soil solution increases from one hour to two months after the addition of MAP fertilizer, the solubility of calcium phosphate minerals decreases; therefore, forms of calcium phosphate mineral present as readily available P become less available forms associated with the HCi-P fraction.
- 4) The granule also of MAP fertilizer applied at a rate of 20 kg P ha⁻¹ has a minimal effect on the efficiency of P uptake. The larger surface area and wider distribution of the smaller size (0.01g) fertilizer granules and of the solution forms of MAP fertilizer result in higher initial uptake of added P. Conversely the small surface area and poor distribution of the large (0.50g) fertilizer granule results in low initial uptake of added P. These effects only leated for the first month because by the second month the P uptake was similar for all also treatments of MAP fertilizer granules.
- Di Proventing contact of MAP fertilizer with sell minerals may increase the efficiency of P uptake. When granules of MAP fertilizer were encosed in either straw or post the efficiency of P uptake increased. The law density of the organic materials may provent sell solution from readily moving into them. Therefore, water would move through the organic materials

slower and less often causing less MAP to dissolve thus lowering fertilizer contact with soil minerals.

6) The placement of bands of MAP fertilizer greatly affects the efficiency of P uptake by plants. Bands of MAP fertilizer within the vicinity of growing roots, 2 cm below and 2 cm baside the seed row, had a much higher efficiency of P uptake by plants than bands of MAP fertilizer placed 10 cm below the seed row. This effect lested throughout the three months of barley growth.

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APPENDIX

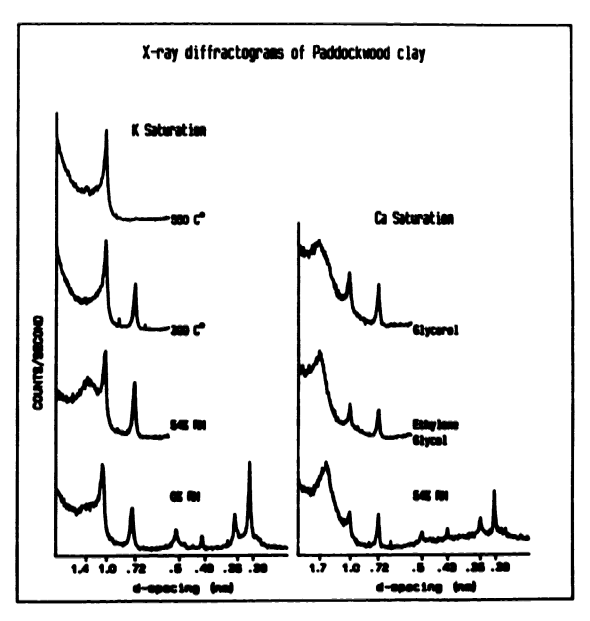


Figure A1. X-ray diffractograms of the clay separate from the Paddeckwood sample.

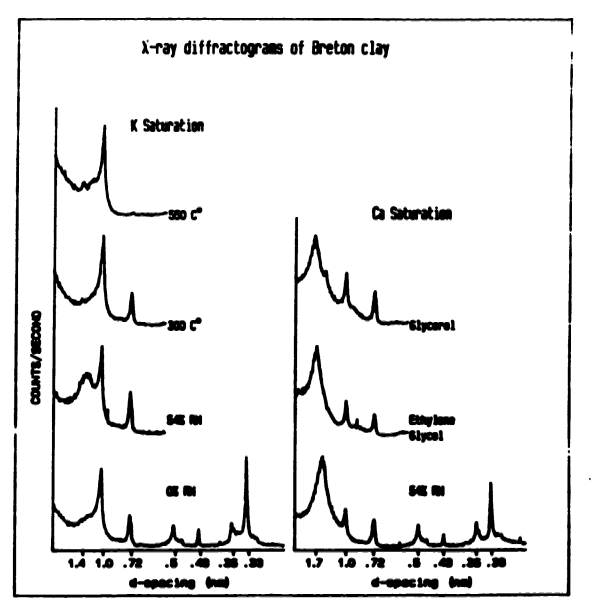


Figure A2. X-ray diffractograms of the clay separate from the Brates sensits.