Phosphorus extractability in surface soil samples as affected by mixing with subsoil

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Samples taken from the plow layer (Ap horizon) and subsoil (B horizon) of six cultivated soil profiles were analyzed as original samples and as mixtures containing 25% or 50% material from the B horizon. Acid ammonium acetate extractable phosphorus, degree of phosphorus saturation (DPS), and a phosphorus Q/I-plot were determined for each sample and mixture to evaluate the effect of bulking of dissimilar materials on results and to assess the possibilities of reducing P solubility in P-enriched surface soils. The results obtained for the mixtures were compared with mass-weighed average results of the original samples. Measured values of DPS corresponded well and those of acetate-extractable P reasonably well to the estimated values, and the results were linearly correlated with the mass fraction of horizon B material in the mixed samples ($r^2 > 0.85$). Water-extractable P behaved dissimilarly; the equilibrium P concentration (EPC) estimated from the Q/I-plots decreased dramatically when the fraction of highly sorptive horizon B material increased in the mixture. The marked effect of subsoil material on EPC values may provide a technique to reduce potential losses of soluble P by deep tillage.

Key words: phosphorus, sampling, soil horizons, sorption isotherms

Introduction

In cultivated soils, phosphorus (P) reserves may exhibit large vertical variability created by soil formation and P fertilization (Peltovuori et al. 2002). Steep gradients between adjacent soil horizons are relevant both in environmental and agronomic contexts, and their recognition is essential for representative soil sampling. Increasing minimum tillage cultivation may also add to the vertical stratification of soil and lead to enrichment of a shallow surface layer with fertilizer or manure P.

Even though the relationship between soil test P and P concentration of runoff water is far from predictable, a high P content of surface soil due to intensive agriculture poses the highest risk for receiving waterbodies (Yli-Halla et al. 1995, Sibbesen and Sharpley 1997). High P status of soil increases the risk of losses of particulate P with erosion material as well as losses of...
soluble P released from soil to runoff water, irrespective of the runoff route. Algal-available P in runoff waters from agricultural soils primarily consists of dissolved reactive P (Ekholm 1998); therefore, soils able to maintain a high concentration of soluble $\text{PO}_4^-$-P in the water phase pose the highest immediate risk for eutrophication.

In noncalcareous soils, the $\text{PO}_4^-$-P concentration of soil solution is determined by sorption-desorption reactions on short-range-ordered oxide surfaces and can accurately be measured with a Q/I-plot technique. The technique gives a comprehensive picture of the solubility of P in a particular soil (Hartikainen 1991) but is not suitable for routine analyses. A less laborious approach used to estimate the $\text{PO}_4^-$-P concentration in runoff waters from divergent soils is the concept of P saturation, which relates the molar concentrations of sorbed P to those of active Fe and Al in soil (e.g. Beauchemin and Simard 1999). The degree of P saturation (DPS) in surface soil has been shown to correspond to the $\text{PO}_4^-$-P concentrations in surface runoff water in rain simulator experiments (Pote et al. 1999). The relationship between DPS and equilibrium P concentration is well established in the laboratory. Under field conditions, the complexity of related hydrological factors diminishes the correlation between runoff $\text{PO}_4^-$-P and soil DPS. One of the key factors in field conditions is the depth of interaction between surface soil and water. Rainfall intensity, slope, plant cover, crop residues, and soil aggregation, among others, affect the depth of interaction (Sharpley 1985); generally, the effective interaction has been considered to occur in a layer ranging from millimeters to a few centimeters.

Due to the shallow layer of interaction between surface runoff and soil, sampling of only the top few centimeters is often recommended for environmental soil P analyses (Sharpley et al. 1978, Gartley and Sims 1994, Turtola and Yli-Halla 1999). This is reasonable if the P conditions in the immediate surface differ from those in the rest of the surface soil: a pattern not unexpected in grassland or no-till soils. Differences between the surface soil and subsoil horizons are practically always distinct. In both cases, too deep sampling leads to bulking of dissimilar materials in unknown ratios, and hence, to ambiguous soil test results. Despite the importance of sampling as part of soil P management, few studies have dealt with possible errors induced by improper sampling or bulking of original samples to composite samples. This study was conducted to explore the effects of bulking of dissimilar soil samples on results of various P analyses. Samples were taken from the Ap horizons and the respective B horizons to obtain clear contrasts. A secondary objective was to estimate possibilities of reducing environmental risks for $\text{PO}_4^-$-P losses by mixing low P material from the subsoil with a P-enriched surface soil layer.

### Material and methods

Six soil profiles were sampled according to genetic soil horizons. From each soil, the topmost Ap horizon (thickness 20–30 cm) and a B horizon starting at an approximate depth of 30 cm were used. Depths and selected characteristics of the horizons are given in Table 1. Prior to the analyses, the samples were sieved, homogenized, and stored at sampling moisture at 5°C in the dark. The analyses were carried out on original Ap and B horizon samples and on mixtures containing 25% or 50% (mass) of horizon B material. Mixtures were made in the extraction vessels by weighing appropriate amounts of the two materials for extraction. Field-moist samples were used in all analyses, but reported mixing ratios and extraction results are calculated on an oven-dry basis (gravimetric moisture at 105°C).

Assuming a hypothetical 100% selectivity and specificity of the analyses, the results for soil sample mixtures ($Y$) were estimated as a mass-weighed average of the original Ap and B horizon sample results according to Equation 1:

$$ Y = M_{Ap} - (M_{Ap} - M_B) \times X, $$
where $M_A$ and $M_B$ are measured results for the original Ap and B horizon samples and $X$ is the mass fraction of horizon B material in the mixture. The estimated results were compared with the values measured for actual soil sample mixtures.

Acid ammonium acetate extractable P ($P_{\text{AAA}}$) was determined in triplicate as in Vuorinen and Mäkitie (1955), with the exception that soil was weighed for the extractions and the results were calculated on a mass basis ($0.5 \text{ M CH}_3\text{ COONH}_4$, $0.5 \text{ M CH}_3\text{ COOH}$, pH 4.65, 1:20 w:v, 1 h shaking). Chang and Jackson P fractions were determined in triplicate as in Hartikainen (1979): 1) easily soluble P was extracted with 1 M NH$_4$Cl (1:50 w:v, 30 min shaking); 2) secondary P bound to Al oxides was extracted with 0.5 M NH$_4$F, pH 8.5 (1:50, 1 h); 3) P bound to Fe oxides with 0.1 M NaOH (1:50, 16 h); and 4) Ca-bound P with 0.25 M H$_2$SO$_4$ (1:50, 1 h). The fractions are referred to as NH$_4$Cl-P, NH$_4$F-P, NaOH-P, and H$_2$SO$_4$-P. Short-range-ordered iron (Fe$_{\text{ox}}$) and aluminum (Al$_{\text{ox}}$) oxides were extracted, also in triplicate, with acid ammonium oxalate ($0.05 \text{ M (NH}_4\text{)}_2\text{C}_2\text{O}_4 \cdot \text{H}_2\text{O}$, pH 3.3; 1:20 w:v; 2 h shaking in the dark) (Niskanen 1989). Phosphorus fractions and oxalate-extractable Al and Fe were used to calculate the degree of P saturation (Peltovuori et al. 2002):

$$\text{DPS} = \frac{(\text{NH}_4\text{Cl-P} + \text{NH}_4\text{F-P} + \text{NaOH-P} + \text{H}_2\text{SO}_4\text{-P})}{(0.5 \times (\text{Al}_{\text{ox}} + \text{Fe}_{\text{ox}}))}$$

Phosphorus fractions were used in Equation 2 instead of oxalate-extractable P because single oxalate extraction grossly overestimates the P saturation in Finnish soils (Uusitalo and Tuhkanen 2000, Peltovuori et al. 2002).

Phosphorus sorption characteristics of the samples were determined with a Q/I-plot technique at a soil-to-solution ratio of 1:50 using P
additions of 0, 0.5, 1.0, 1.5, and 2.0 mg l\(^{-1}\) as KH\(_2\)PO\(_4\) in a H\(_2\)O matrix. After shaking the samples with solutions for 21 h, the suspensions were filtered (Nuclepore\textsuperscript{®} polycarbonate, 0.2 mm) and the PO\(_4\)-P concentrations of the filtrates determined with a molybdenum blue method. The amount of sorbed or desorbed P was calculated from the difference in the P concentration in the solution before and after the equilibration. A modification of the Freundlich adsorption equation \((Q = Q_0 + k \times \ln(F)), Fitter and Sutton 1975\) was fitted to the Q/I points obtained in duplicate determinations, and the equations were used to calculate equilibrium phosphorus concentrations (EPC) (no net sorption or desorption of P). Coefficients of determination for the equations obtained ranged from 0.98 to 1.00. Water-extractable P (P\(_{\text{H}_2\text{O}}\)) was measured from the Q/I-plot filtrates of zero P addition.

**Results**

Degrees of phosphorus saturation measured in the mixtures containing both Ap and B horizon materials agreed well with the estimates calculated with Equation 1 (Fig. 1a). The 95% confidence interval for the regression coefficient in Fig. 1a (0.94–1.14) indicates no difference between measured and estimated values of DPS. Acetate-extractable P results for the mixtures, in contrast, deviated slightly from those estimated with Equation 1 (Fig. 1b). The regression coefficient excluded the value 1 (0.88–0.93, \(P = 0.95\)), indicating that at high P concentrations the measured values were lower than expected. In water extraction (P\(_{\text{H}_2\text{O}}\)), the measured values for the mixed samples were distinctively lower than those estimated with Equation 1 (Fig. 1c, regression coefficient 0.49–0.68, \(P = 0.95\)). Coefficients of correlation both for DPS and P\(_{\text{AAA}}\) were also high compared with that of P\(_{\text{H}_2\text{O}}\).

The behavior of DPS in mixed soil samples was close to ideal and that of acetate-extractable P was predictable (Fig. 1); in these cases,
Table 2. Linear regression equations between acid ammonium acetate extractable P (P_{AAA}, \text{mg kg}^{-1}) and the mass fraction of horizon B material (x) in the composite sample (Ap+B), and between the degree of P saturation (DPS) and the mass fraction of horizon B material in the composite sample.

<table>
<thead>
<tr>
<th>Soil</th>
<th>y = P_{AAA}</th>
<th>n</th>
<th>R^2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kotkanoja</td>
<td>y = 8.88 – 8.69x</td>
<td>12</td>
<td>0.98</td>
</tr>
<tr>
<td>Lintupaju</td>
<td>y = 11.33 – 10.01x</td>
<td>10</td>
<td>0.95</td>
</tr>
<tr>
<td>Sjökulla</td>
<td>y = 4.38 – 3.85x</td>
<td>12</td>
<td>0.99</td>
</tr>
<tr>
<td>Toholampi</td>
<td>y = 3.39 – 2.60x</td>
<td>12</td>
<td>0.95</td>
</tr>
<tr>
<td>Loppi</td>
<td>y = 62.12 – 62.20x</td>
<td>12</td>
<td>0.98</td>
</tr>
<tr>
<td>Loppi2</td>
<td>y = 13.49 – 8.40x</td>
<td>12</td>
<td>0.92</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Soil</th>
<th>y = DPS</th>
<th>n</th>
<th>R^2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kotkanoja</td>
<td>y = 0.193 – 0.096x</td>
<td>12</td>
<td>0.86</td>
</tr>
<tr>
<td>Lintupaju</td>
<td>y = 0.360 – 0.253x</td>
<td>12</td>
<td>0.86</td>
</tr>
<tr>
<td>Sjökulla</td>
<td>y = 0.160 – 0.106x</td>
<td>12</td>
<td>0.94</td>
</tr>
<tr>
<td>Toholampi</td>
<td>y = 0.099 – 0.069x</td>
<td>12</td>
<td>0.97</td>
</tr>
<tr>
<td>Loppi</td>
<td>y = 0.460 – 0.372x</td>
<td>12</td>
<td>0.94</td>
</tr>
<tr>
<td>Loppi2</td>
<td>y = 0.167 – 0.057x</td>
<td>12</td>
<td>0.85</td>
</tr>
</tbody>
</table>

The use of Equation 1 is meaningful. The linear regression equations between measured acetate-extractable P or DPS and the fraction of horizon B material in the composite samples are presented in Table 2. The fit of all equations was good (r^2 > 0.85), and the regression coefficients as well as the constant terms of the equations were in accordance with the analytical results for the original Ap and B horizons (P_{AAA} in Table 1, DPS not shown) and Equation 1: the differences between the coefficients in Table 2 and those calculated on the basis of the results of original Ap and B horizons [M_{Ap} and (M_{Ap} – M_{B})] were all less than 5%.

As demonstrated in Fig. 1c, the behavior of water-extractable P differs from that of DPS and acetate-extractable P because of the very high sorption affinities of the B horizons and consequent resorption of P in mixed samples during the extractions. Two-way desorption-resorption reactions during extraction result in a nonlinear behavior of water-soluble P in soil sample mixtures. This nonlinearity is evident in Fig. 2, which depicts the EPC values of composite soil samples as a function of the mixing ratio. Especially the high EPC values decreased dramati-
cally when subsoil material was added to the mixture. The values of the original B horizons were lower than 0.01 mg l\(^{-1}\), with the exception of the Loppi2 soil (0.06 mg l\(^{-1}\)).

**Discussion**

The degree of P saturation (DPS) and acetate-extractable P determined for a composite sample proved to be a mass-weighed average of the original samples. Thus, it can be estimated with Equation 1 if the results of the original samples and the mixing ratio are known. The predictable performance of these analyses is a consequence of net desorption of P from all mixed materials during the extraction. In the P fractionation used in the DPS determination, resorption is prevented by the strong extractants used. In the acid ammonium acetate extraction, the organic ligands of the extractant have a similar effect. At high P concentrations, however, the ability of the extractant to prevent resorption is insufficient and some resorption of P occurs, as can be seen from the regression coefficient in Fig. 1b (< 1). Resorption of P has been demonstrated to occur even with strong extractants when long extraction times are used (Rodrigues and Mendoza 1993). During water extraction or equilibration in Q/I-plot analyses, resorption from solution is probable to materials of high sorption affinity, leading to a nonlinear relationship between equilibrium P concentration and the mixing ratio. According to del Campillo et al. (1996), the possible deviations in P test results between a measured result of a composite sample and the calculated average of the original samples is highest when the samples differ in their intrinsic properties, e.g. sorption affinities.

The ranges of DPS (0.03–0.47) and acetate-extractable P (0.5–71 mg kg\(^{-1}\)) measured in this study were large and cover the values typically encountered in the field – although values at the lower end of the scales were dominant. For both analyses, correlations between measured and estimated values for mixed samples (Fig. 1) were high, considering that the results of the original Ap and B horizon samples were excluded. These values were not included because Equation 1 returns the average result of Ap or B horizon when \(X\) is 0 or 1, thus raising the correlation inappropriately. The values in Table 2 were calculated using all observations, including the results of the original samples.

The extensive use of moldboard plowing has created marked differences in P characteristics between Ap and B horizons in cultivated soils in Finland (Kaila 1963, Jokinen 1984, Puustinen et al. 1994). The boundary between the horizons is visually distinct and unintentional mixing of the horizons is usually easy to avoid in soil sampling. However, an apparently homogeneous Ap horizon may contain materials differing widely in P reserves; Humphreys et al. (1998) found up to 200% higher soil P test results in samples taken from the top 7.5 cm than in samples taken from the top 10 cm in the same grassland soils due to biological P translocation and surface fertilization or manuring. In Finland, a three-year surface application of slurry and P fertilizer on grass ley elevated the P contents and acetate-extractable P in the top 5 cm of soil, while soil at 5–25 cm depth was unaffected (Tuttoo and Yli-Halla 1999). Placement of P fertilizers with planting creates distinct differences between the in-row and between-row soil test results (Urvas and Jussila 1979), but the differences disappear during the growing season. Minimum tillage practices may also create vertical stratification of nutrient concentrations, especially when combined with surface application of phosphorus. Howard et al. (1999) found distinctive P enrichment in the top 8 cm of soil after six years of no-till cotton farming. Further research on the most appropriate depth of sampling, as well as other aspects of representative sampling for environmental P analysis, is obviously required.

Water extraction has been suggested as a universal environmental soil P test for predicting P losses in surface runoff. Because of the rapid decrease in high EPC values with a small amount
of unintentional sorptive material in the sample, this analysis seems more vulnerable to errors induced by inappropriate soil sampling than the other two methods, acid ammonium acetate extraction and DPS. On the other hand, the non-linearity between the concentration of PO₄-P and the fraction of sorptive material in a mixture provides a promising technique to reduce the solubility and potential losses of P in P-enriched top-soils; in the Loppi soil with an excessive P-test value, 25% of horizon B material mixed with the Ap horizon increased the EPC value by 70% with a simultaneous increase in buffering capacity. Success of the technique requires a B horizon of high P-fixing capacity. In the Loppi2 soil, the marginal decrease in the EPC values in mixed samples (Fig. 2) is most likely due to competitive sorption by soluble organic matter originating from the Bw horizon containing almost 10% organic carbon.

Of the several mandatory and voluntary measures currently used in reducing P loss from agricultural soils – fertilization limits, buffer zones, reduced tillage, constructed wetlands, and sedimentation ponds – only limiting the use of P focuses on the P release potential of the soil. Decreasing the area of high P content soils would be important in protecting the surface waters because the concentration of PO₄-P in runoff waters is primarily controlled by the P status of surface soils (Yli-Halla et al. 1995). Unfortunately, high or excessive soil P-test values decrease slowly with existing P fertilization recommendations (Yli-Halla et al. 2001) and even with zero P additions (Jaakkola et al. 1997). This laboratory study showed that on selected soils the leaching risk of the potentially most harmful form of P can be reduced quickly by mixing the surface soil with P-fixing subsoil. This remediation method could be applied on areas possessing high P-loss potential (hot spots). The method might even be attractive to some farmers since deep tillage has been shown to increase the accessibility of P by roots, and consequently cereal yields, on drought-prone clay soils (Saarela et al. 2000).

Acknowledgments. I thank the Finnish Drainage Research Foundation and the Finnish Cultural Foundation for financial support and Helena Soinne for the analytical work.

References


Peltovuori, T. Phosphorus in mixed soil samples


Muokkauskerroksen sekoitetun pohjamaan vaikutus maasta uuttuvan fosforin määrään

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Tutkimuksessa selvitettiin muokkauskerroksen mukaan maanötteen mahdollisesti joutuvan pohjamaan vaikutosta tehtävien fosforianalyysien tuloksiin. Samalla tutkittiin mahdollisuksia pienentää runsaasti fosforia sisältävää muokkauskerroksesta vapautuvaa liukosien määrää sekoitamalla siihen fosforia tehokkaasti pidättävää pohjamaata.

Kuudesta viljellystä maaprofiilista otettiin maanötteet muokkauskerroksen ja muokkauskerroksen alapuolisesta pohjamaasta. Puhtaiden maanöttestä ja maanöttestä, jotka sisälsivät 25 tai 50 % pohjamaata, analysoitiin viljauuskoski, fosforin kyllästysaste ja fosforin Q/I-kuvaaja. Seoksista mitattuja analyysituloksia verrattiin muokkauskerroksen ja pohjamaan analyysitulosten massaosuksilla painotettuihin keskiarvoihin.

Viljauuskoskin ja fosforin kyllästysasteen osalta maanöttestä mitatut ja puhtaiden näytteiden tulosten perusteella lasketut arvot olivat hyvin läheillä toisiinsa. Pohjamaan osuuden kasvaessa maanöt seedeksessä viljauuskoskin j ja fosforin kyllästysaste pienenivät lineaarisesti (r² > 0,85). Lineaarisen riippuvuuden perusteella maanötseedeksen analyysitulos voidaan ennustaa, mikäli puhtaiden näytteiden tulokset ja maiden sekoitussuhde tiedetään. Vesiuttoisen fosforin kohdalla seoksista mitatut arvot olivat sen sijaan huomattavasti pienempää, kuin alkuperäisten pinta- ja pohjamaanöttestä perusteella lasketut tulokset, ja vesiuttoisen fosforin pitoisuus sekä fosforin Q/I-kuvaajan avulla lasketut tasapainotason pohjamaan osuuden kasvaessa maanötseedeksessä. Pohjamaan voimakas taipumus pienentää vesiliukoisen fosforin määrää ja fosforin tasapainopitoisuutta seoksessa saattaa tarjota mahdollisuuden pienentää fosforin liukoisuutta runsaasti fosforia sisältävissä maisissa syvään muokkausen avulla, ja siten vähentää välittömästi rehellöitymistä aiheuttavan liukoisen fosforin huhtoutumista vesistöihin pintavalunnan mukana.