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A novel rhizosphere trait-based approach to evaluating soil phosphorus availability across complex landscapes

Thomas H. DeLuca

Highlights

- Existing methods for assessing phosphorus (P) availability do not adequately reflect plant P acquisition strategies
- We evaluated a novel P extraction procedure to explore the concept of biologically based P protocol
- Soil P was extracted in parallel with CaCl_2 , citric acid, phytase and phosphatase solution and 1 M HCl
- We tested this method on 204 soil samples collected in the United Kingdom and compared it with the standard Olsen P method
- This method helped explain an observed downward trend in Olsen P from 1998 to 2007 as a shift from inorganic to organic P.
- This method can be used as a means of assessing P availability across complex landscapes

1 **A novel rhizosphere trait-based approach to evaluating soil phosphorus availability**
2 **across complex landscapes**

3

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16

17 **ABSTRACT**

18 Plants employ a range of strategies to increase phosphorus (P) availability in soil. Current soil
19 P extraction methods (e.g. Olsen P), however, often fail to capture the potential importance of
20 rhizosphere processes in supplying P to the plant. This has led to criticism of these standard
21 approaches, especially in non-agricultural soils of low P status and when comparing soil
22 types across diverse landscapes. Similarly, more complex soil P extraction protocols (e.g.
23 Hedley sequential fractionation) lack functional significance from a plant ecology
24 perspective. In response to this, we developed a novel procedure using a suite of established
25 extraction protocols to explore the concept of a P protocol based on biologically significant P
26 pools, fluxes and transformations. Soil P was extracted in parallel by using 10 mM CaCl₂
27 (soluble P), 10 mM citric acid (chelate-labile P), phytase and phosphatase solution (enzyme
28 labile organic P) and 1 M HCl (mineral occluded P). To test the integrated protocol, we
29 conducted the analyses on 204 soil samples collected as part of a UK national ecosystem
30 survey (Countryside Survey) in 1998 and repeated again in 2007. Overall, Olsen P showed a
31 net decline in national soil P levels during this 10 year period. In accordance with these
32 results, soluble P, chelate-labile P and occluded P were all found to decrease over the 10 year
33 study period. In contrast, enzyme labile organic P increased over the same period likely due
34 to the accumulation of P in litter and O horizon organic matter. This new method is simple
35 and inexpensive and therefore has the potential to greatly improve our ability to characterise
36 and understand changes in soil P status across complex landscapes.

37

38 *Keywords:* Bioavailability, Ecosystem assessment, Nutrient index, Phosphate, Soil quality
39 indicator

40 **1. Introduction**

41 Increasing food security concerns and decreasing mineable phosphorus (P) supplies
42 necessitate efficient use of soil P resources; however, current methods used to assess plant
43 available P are often ineffective when used on landscapes with a great degree of plant and soil
44 heterogeneity. Soil P exists in a variety of forms including soluble inorganic, insoluble
45 inorganic (P_i), organic, and surface adsorbed with the amounts present in each fraction
46 varying greatly between soil types (Bielecki, 1973).

47 The ability to effectively assess soil P status and phytoavailability is extremely
48 important in terms of environmental protection and agricultural productivity; however,
49 phytoavailable P is not a distinct value for any given soil (Withers et al., 2014). Importantly,
50 plants express unique mechanisms for releasing P from different pools of differing
51 recalcitrance, each contributing to varying extents depending upon several plant and soil
52 parameters (Neumann and Römheld, 1999; Lambers et al., 2006). Current efforts to monitor
53 soil P status are based on methods specifically developed for agricultural purposes with the
54 specific objective of estimating the phytoavailability of soil P and enabling fertiliser rate
55 recommendations (e.g. Mehlich, 1978; Menon et al., 1989; Saggar et al., 1992; Sims et al.,
56 2000). Commonly, these are single solution extractions (e.g. NaHCO_3 or acid NH_4F) which
57 correlate with plant P_i uptake in a controlled environment (e.g. Bray and Kurtz, 1945; Olsen
58 et al., 1954; Mehlich, 1984). These extractions have proved very useful for agriculture as they
59 offer a straightforward index of P fertility. Across complex landscapes; however, single
60 extraction methods do not adequately characterise the bioavailability of P which is directly
61 influenced by plant community and shifts in soil biophysical conditions. Phosphorus
62 fractionation schemes were developed in an attempt to better characterize the P status of soils
63 (e.g. Hedley et al. 1982). Such fractionation approaches expose a single soil sample to a
64 sequence of extractants to quantify pools of progressively occluded P. These approaches

65 offer a more detailed picture of soil P status, are more suited to use over complex landscapes,
66 offer some sense of how P might become available over time and they can provide an
67 indication of the mechanisms controlling P solubility in a given soil (Cross and Schlesinger,
68 1995; Levy and Schlesinger, 1999; Negassa and Leinwieber, 2009). Examples of
69 fractionation methods include the widely adopted Hedley procedure (Hedley et al., 1982) or
70 the Chang and Jackson method (Chang and Jackson, 1957). Unfortunately, fractionation
71 methods are time consuming and require careful preparation making them inappropriate for
72 routine use, especially in agriculture. Furthermore, these fractionations do not adequately
73 reflect rhizosphere processes (Johnson et al., 2003; Yang and Post, 2011). Phosphorus
74 solubilised by rhizosphere processes (in particular organic acid, proton and ectoenzyme
75 excretion) are not individually characterised in these schemes. Instead, chemical analogues
76 are used which, while they may correlate well with plant availability or P accumulation with
77 soil development, they do not offer insight into the potential P uptake mechanisms or
78 rhizosphere P transformations that drive ecosystem P dynamics.

79 In this paper we introduce an alternative functional plant trait-based approach to
80 evaluate soil P status. Here we combine together four established approaches to assessing
81 different pools of bioavailable P thereby simultaneously assessing soil P as influenced by
82 plant rhizosphere mediated processes across a diverse array of soils. The extractants were
83 chosen to emulate four common and significant plant rhizosphere mediated P acquisition
84 mechanisms: (1) root interception, (2) organic acid complexation, (3) enzyme hydrolysis and
85 (4) proton excretion induced acidification. Rather than sequentially extracting these P pools
86 as in the Hedley fractionation, we run the extractions in parallel to measure the total amount
87 of P mobilised by each individual test. The purpose of this effort was to create a simple P
88 assessment regime that reflects rhizosphere mediated P availability, is sensitive to landscape
89 variation in soil P status, and facilitates evaluation of short, medium and long term fluxes

90 between P pools. The combined analyses are collectively referred to as the Rhizosphere
91 Based P (RBP) extraction regime. The RBP method is compared with the standard Olsen P
92 method across a variety of soils and is compared on field moist and air dried soils.

93

94 **2. Materials and methods**

95 *2.1. Soils*

96 For the main study, soil samples were collected throughout the UK as part of the
97 Centre for Ecology and Hydrology Countryside Survey (CS) in 1998 (CS98) and 2007
98 (CS07) with sites representing all the dominant landscape types and soil groups in the UK
99 (Emmett et al., 2010; Reynolds et al., 2013). To encompass all the major soil and land use
100 types, a total of 2614 soil samples were collected throughout the UK, based on a stratified
101 random sample of 1 km squares at gridpoints on a 15 km grid using the Institute of Terrestrial
102 Ecology (ITE) Land Classification as the basis of the stratification (Wood et al., 2012). At
103 each grid intersection, a 1 km² sample area was selected. Within the 1 km² sample area, 3
104 plots (5 × 5 m²) were randomly located and a single 15 cm long × 4 cm diameter soil sample
105 was collected from each of the plots. Additional information about vegetation and soils were
106 also collected from the same plots. To facilitate comparison of P pool concentrations during
107 the two sample dates, we used the vegetation and soil categories provided in the CS (Emmett
108 et al., 2010). For plant communities we used the ‘Aggregate Vegetation’ grouping which
109 includes eight categories: 1) lowland wooded; 2) upland wooded; 3) crops and weeds; 4) tall
110 grass and herbs; 5) fertile grassland; 6) infertile grassland; 7) moorland; 8) heath and bog. For
111 soil types, we use the loss-on-ignition categories of: 1) mineral; 2) humus-mineral; 3) organo-
112 mineral; 4) organic. The 1 km² areas were stratified within the 45 major Land Classes of the
113 UK. All the sites were characterised by a temperate climate with a North-South mean annual

114 temperature range of 7.5 to 10.6°C and East-West mean annual rainfall range from 650 to
115 1700 mm.

116 Samples were stored at 4°C prior to analysis for key characteristics including pH,
117 total C and N, mineralisable C and N, Olsen-P (0.5 M NaHCO₃, pH 8.5), bulk density and
118 soil biota as described in Emmett et al. (2008), Emmett et al. (2010), Simfukwe et al. (2011)
119 and Reynolds et al. (2013). All remaining sample was then air-dried and sieved prior to long
120 term storage and use in this study.

121 To assess the changes in soil P seen between the 1998 and 2007 Countryside Survey, a
122 subset of 102 spatially paired soils (204 in total) from the CS98 and CS07 archived soils was
123 selected randomly. In order to represent the archive's spatial diversity, the samples were
124 stratified according to their "Environmental Zone" – nine classifications derived from
125 Institute of Terrestrial Ecology Land Classes which reflect an array of geographically distinct
126 regions of Britain (Bunce et al., 1996). Across all land use and vegetation classes the
127 dominant soil types (% of total) were brown soils (33%), surface water gley soils (19%),
128 podzolic soils (14%), peat soils (12%), groundwater gley soils (11%), lithomorphic soils (8%)
129 and pelosol soils (3%) (Avery, 1990; Simfukwe et al., 2011). These soils were assessed using
130 the novel Rhizosphere Based P (RBP) extraction regime described below and for total C
131 based on loss-on-ignition (Nelson and Sommers, 1982; Reynolds et al., 2012).

132

133 *2.2. Principles behind the proposed RBP method*

134 We employed four existing soil P analysis methods to provide a clear picture of soil P
135 status as influenced by plant rhizosphere mediated processes. Phosphorus in soil can be
136 grouped into three primary pools: (1) readily available, dissolved orthophosphate, (2) more
137 recalcitrant "active P" forms which, over time, are solubilised to replenish this readily
138 available pool, and (3) fixed P which may remain unchanged in soil for many years. The

139 method below herein uses a combination of established extraction procedures to represent the
140 P solubilised by the four primary plant P acquisition mechanisms: (1) root interception, (2)
141 organic acid complexation/dissolution, (3) enzyme hydrolysis and (4) proton excretion
142 induced acidification. The procedures were adapted in order to correspond to the maximum
143 level of each extractant reported in the literature.

144 Each fraction was measured in parallel by shaking 0.5 g of soil with each extractant
145 (10 ml; described below) in separate 15 ml centrifuge tubes for 3 h on a reciprocal shaker at
146 200 rev min⁻¹. Preliminary work showed 3 h to be the point at which equilibrium was reached
147 between soil- and solution-P. Extracts were then centrifuged (3,220 g, 30 min) to negate the
148 need to filter the supernatant (Poile et al., 1990). An aliquot of the supernatant was then
149 decanted and stored for no more than 3 d at 4°C prior to analysis.

150 Soluble P was assessed using a 10 mM calcium chloride (CaCl₂) solution which
151 corresponds to labile P that is easily available to plants (Bielecki, 1973; van Raij, 1998).
152 Typically, this is a relatively small pool of P which root hairs and arbuscular mycorrhizas
153 might remove directly from the soil solution.

154 Organic acid extractable P was assessed using a 10 mM solution of citric acid to
155 quantify the chelate-extractable, active pool of P sorbed to clay particles or as compounds of
156 Ca, Fe or Al which have been shown to be accessible to plants following the release of
157 organic acids into soil (Jones and Darrah, 1994; Hinsinger, 2001; Johnson and Loeppert,
158 2006; Li et al., 2007). Citrate extractable P was chosen over acetic acid or oxalic acid,
159 because it does not interfere with the P analysis reagents described below and is frequently
160 implicated in root and microbial P mobilization in soil.

161 Phosphatase (acid phosphatase from wheat germ; Sigma P3627; Enzyme Commission
162 Number 232-630-9) and phytase (from wheat, Sigma P1259; Enzyme Commission Number
163 3.1.3.26) enzymes were used to evaluate the quantity of available organic P. The final

164 concentration of the enzymes in the extraction solution was 0.02 enzyme units ml⁻¹. This
165 concentration was sufficient to ensure that they would be present in excess. The solution is
166 prepared by the addition of phosphatase and phytase to a sodium acetate buffer (50 mM, pH
167 6.5) with MgCl₂ (0.08 mM) added as a pre-enzyme activator (Ahlers, 1974). We should note
168 here that in more recent enzyme assays we have found commercially available phytase
169 (purchased from Sigma) to be contaminated with P so we have since switched to only using
170 phosphatase.

171 The more recalcitrant P was extracted using 1.0 M HCl. This recalcitrant P fraction is
172 thought to be solubilised by proton excretion in the rhizosphere and by microbial processes
173 (Petersen and Böttger, 1991; Gahoonia et al., 1992).

174 All extracts were diluted appropriately and analysed colorimetrically (630 nm) using
175 the malachite-green method as described in Ohno and Zibilske (1991) using a PowerWave-
176 XS microplate spectrophotometer (BioTek Instruments Inc., Winooski, VT). Malachite-green
177 was chosen over the standard molybdate blue method (Murphy and Riley, 1962), as it is
178 highly sensitive and not susceptible to interference from organic acids. The method was
179 slightly modified to incorporate a ten-fold in-plate dilution where necessary.

180 The standard method used for assessing P availability in the CS is the Olsen-P method
181 (Allen, 1989). Briefly, 5 g of air-dried soil was mixed with 100 ml of 0.5 M sodium
182 bicarbonate at pH 8.5. Phosphate in the extract was then determined colorimetrically by
183 molybdate blue at 880 nm using a Skalar continuous flow analyser with the addition of a
184 dialysis step to overcome the effect of the Olsen's reagent.

185

186 *2.3. Comparison of Olsen P and the RBP method in field-moist soils*

187 The soils evaluated in Section 2.2 were all air-dried prior to extraction (following the
188 UK national soil inventory protocol). To compare the proposed RBP method with the

189 standard Olsen P method in field-moist and air dried samples, we collected 27 independent
190 soil samples (0-10 cm) from different farms within the Hiraethlyn catchment in North Wales
191 ($53^{\circ}10'N$, $3^{\circ}45'W$; area = 27 km²). The samples were characterised as described above with
192 exception of ammonium (NH₄⁺) and nitrate (NO₃⁻) which were measured in 0.5 M K₂SO₄
193 extracts as described in Jones and Willett (2006). The samples ranged in soil organic matter
194 content from 4.61 to 18.19 % (mean \pm SEM, $10.54 \pm 0.62\%$), pH from 4.76 to 6.36 (mean \pm
195 SEM, 5.57 ± 0.08), moisture content from 7.8 to 80.8% (mean \pm SEM, 49.5 ± 4.0), available
196 NO₃⁻ from 2.4 to 49.1 mg kg⁻¹ (mean \pm SEM, 15.4 ± 1.9 mg kg⁻¹), available NH₄⁺ from 0.8 to
197 42.9 mg kg⁻¹ (mean \pm SEM, 5.6 ± 1.7) and available K from 61 to 364 mg kg⁻¹ (mean \pm SEM,
198 157 ± 15). The soils were sieved to pass 5 mm and stored at 5°C until weighed out for
199 extraction as either fresh (field moist, corrected to dry weight based on moisture content) and
200 air dried (dried for 48 hours at room temperature) were extraction using the RBP procedure as
201 described above.

202

203 *2.4. Statistical analysis*

204 A one-way ANOVA was used to detect changes in P concentration between the two
205 survey years for the different fractions. Data were then split according to one of three
206 grouping variables, namely (1) vegetation community type, (2) broad ecosystem type, and (3)
207 soil organic matter content (measured via loss-on-ignition) and ANOVA undertaken to
208 identify differences in P concentration. Pearson correlations were used to assess the
209 relationship between our individual extraction techniques and that of the standard Olsen P
210 method employed on the Countryside Survey. Principle components analysis (PCA) was used
211 to explore variability, patterns, and relationships between P concentrations (mg kg⁻¹) of the
212 four P pools and Olsen P. Significant ($p < 0.05$) environmental and soil characteristic vectors
213 were fit onto the PCA ordination. In a PCA, maximum variances are accounted for but a

214 normal distribution of the population is not a requirement (Reimann et al., 2011). Incomplete
215 observations were excluded from PCA except for AgClass where two blank values for 1998
216 data were substituted with 2007 values. Outliers were included in the analysis. Data was
217 scaled to ensure homogeneity of variances. Correlations and ANOVA were analysed using
218 SPSS 16 for windows (SPSS Inc., Chicago, IL) and PCA was run using the vegan package
219 (Oksanen et al., 2013) in the R Statistical Environment (R Version 3.0.3, [http://www.r-](http://www.r-project.org/)
220 [project.org/](http://www.r-project.org/)). For comparison of P fractions in the field-moist soils, linear regression and t-
221 tests were undertaken using Minitab v16 (Minitab Inc, State College, PA).

222

223 **3. Results**

224 *3.1. Relationship between the soil P extractants*

225 Three of the methods used in our rhizosphere-based P fractionation protocol were
226 highly correlated with the Olsen P method with the exception of the enzyme extraction
227 method which was weakly correlated with Olsen P ($P < 0.05$; Table 1). Citrate-extractable P
228 was most highly correlated ($r^2 = 0.563$, $P < 0.001$) with the enzyme extraction closely
229 followed by the 1.0 M HCl extraction ($r^2 = 0.432$, $P < 0.001$). All three of these methods are
230 effective at accessing moderately soluble mineral adsorbed and precipitated mineral forms of
231 P. The HCl extractable P was also highly correlated ($r^2 = 0.732$, $P < 0.001$) with citrate
232 extractable P.

233 The relationship between the four P extraction methods of RBP and that of Olsen P
234 are further demonstrated in Figure 1. Using principal components (PC) analyses, we found
235 that PC1 explains 48.66% of the total variation in the P concentration across methods and
236 PC2 accounts for 20.71% of the total variation. Figure 1 provides a visualization of PCA
237 scores, calculated by observations and displayed by grey dots, in relation to the loadings, or P
238 methods (in blue). The lengths of the arrows are proportional to the variability explained by

239 PC1 and PC2 and angles between loadings represent the correlation between the variables.
240 The arrows labeled with environmental or soils characteristics (in red) point to the direction
241 of the most rapid change across that variable and lengths indicate the correlation of that
242 variable and the P method ordination. Factor loadings for this PCA reveal close associations
243 between citrate and HCl-extractable P. Enzyme-extractable P explains the least variability in
244 the data is markedly distinct from all other methods.

245

246 3.2. Country scale changes in soil P status

247 Assessing the change in P pools in the UK Countryside Survey soils over the 10 year
248 period, we observed a significant decrease in P in the inorganic P fractions (HCl, CaCl₂ and
249 citrate extractable). The largest percentage change was observed in the CaCl₂, or soluble,
250 fraction with a 41% decrease ($P < 0.05$) from 1998 to 2007 (Table 1). Citrate extractable P
251 decreased significantly ($P < 0.01$) from 284 mg P kg⁻¹ to 188 mg P kg⁻¹ between 1998 and
252 2007. The less labile inorganic (P_i), as extracted by HCl, decreased from 573 to 399 mg kg⁻¹
253 ($P < 0.05$) during this same period. Interestingly, enzyme extractable P increased ($P < 0.001$)
254 by more than a factor of two from 130 mg kg⁻¹ in 1998 to 291 mg kg⁻¹ in 2007. The increase
255 in organic extractable P may partially explain the decrease in inorganic P fractions as there
256 was no significant difference between the sum of the averages of the four extractants for the
257 two sampling dates.

258 Taking the UK as a whole, the pattern of decreasing available inorganic P (based on
259 an Olsen-P bicarbonate extraction) described in 2007 CS is corroborated by the shift in
260 inorganic P pools as demonstrated by the RBP.

261

262 3.3. *Changes in soil P with vegetation community and soil organic matter types*

263 The general trend of decreasing inorganic P and increasing organic P is apparent when
264 soils are grouped by plant community. Ecosystem type or aggregate vegetation class (AVC)
265 describes the predominant habitat of the parcel of land on which the sampling plot is located.
266 The HCl-extractable P consistently made up the largest P fraction as it likely accounts for
267 most of the P in the more labile inorganic P pools. Enzyme extractable organic P (P_o)
268 increased (Fig. 2) from 1998 to 2007 and inorganic P as extracted by citrate and HCl
269 decreased during this same period (Fig. 2). However, no significant changes were observed
270 for the labile $CaCl_2$ fraction (Fig. 2). No significant changes were seen for either of the AVC
271 woodland classifications (Upland woodland and Lowland woodland) or under the crop and
272 weed category. The HCl extractable P decreased by 569 mg P kg^{-1} ($P < 0.05$) under tall grass
273 and herb. Enzyme-extractable organic P increased ($P < 0.05$) in fertile and infertile
274 grasslands, heath and bog, and moorland, while citrate-extractable inorganic P increased;
275 however, the changes in both fractions in heath and bogs are much larger than in the
276 grasslands.

277 Within the four soil organic matter (SOM) status groupings, larger changes in P were
278 observed in the soils with the highest C contents. In particular, we observed a decrease in the
279 inorganic P fractions extracted with $CaCl_2$, citrate and HCl. Enzyme-extractable organic P did
280 not follow a specific pattern with soil SOM status (Fig. 3). However, in all but the organo-
281 mineral classifications there was a significant ($P < 0.05$) increase in enzyme-extractable
282 organic P and significant ($P < 0.05$) decreases in HCl-extractable P in the highest and lowest
283 SOM categories as well as large significant decreases in citrate-extractable inorganic P (Fig.
284 4).

285

286 3.4. Comparison of Olsen P and the RBP method in field-moist and dried soils

287 Using the Olsen extraction method, the field-moist samples from the Hiraethlyn
288 agricultural catchment in North Wales showed a wide range of P levels ranging from 6 to 63
289 mg P kg⁻¹ (mean ± SEM, 27 ± 3). Overall, P concentrations in the Olsen extracts were
290 significantly correlated with P recovered in all four proposed RBP extraction regime (Fig. 4).
291 Of these, the best correlation was seen with the citrate extraction ($r^2 = 0.87$), while the
292 weakest correlation was found between the enzyme-based and Olsen bicarbonate extraction
293 ($r^2 = 0.16$). Soil P pools in moist versus dry soils were found to be closely aligned for all P
294 pools (Fig 5); however, air drying nearly doubled P extraction by citrate ($P < 0.01$) and
295 enzymes ($P < 0.001$) and slightly increased CaCl₂ soluble P ($P < 0.05$). Air drying of soils
296 slightly decreased P extraction by using 1 M HCl ($P < 0.05$).

297

298 4. Discussion

299 4.1. Basing an assessment of available P on known rhizosphere processes

300 Bicarbonate extraction of soil, or Olsen P, is one of the most widely adopted test used
301 for assessing soil P availability. Further, it is often used in broad regional or national scale
302 assessments of soil P status (e.g. Sparling and Schipper, 2004; Emmett et al., 2010; Zhang et
303 al., 2012). While highly suited to near-neutral or alkaline pH agricultural soils, Olsen P has
304 been shown to be of less use in predicting plant available P in semi-natural acidic and peat
305 soils (Kuo, 1996; Emmett et al., 2008). For example, across a diverse range of agricultural
306 soils ($n = 164$), Speirs et al. (2013) demonstrated that Olsen-P only provides an approximate
307 guide to plant P availability (correlation between Olsen P and wheat yield, $r^2 = 0.064$).
308 Further, Jordan-Meille et al. (2012) have openly criticised current soil P availability testing
309 procedures calling for “a more mechanistic approach in which the processes involved in plant

310 P nutrition are truly reproduced by a single standard method". This has led to the emergence
311 of alternative approaches such as diffusive gradient thin films (DGT) which have proven to
312 provide better predictors of plant P availability than Olsen P (Six et al., 2014). The DGT
313 technique is highly suited to soils receiving high levels of fertiliser where plant capture is
314 largely related to sorption-desorption reactions and where rhizosphere P acquisition
315 mechanisms are down-regulated. However, we do not feel that a single chemical extraction or
316 technique like DGT adequately represents P availability in more P limited non-agricultural
317 environments where plants may be expressing a diverse array of mechanisms to exploit soil P
318 reserves. In our view this complexity needs to be captured by parallel extractions.

319 In both the national and regional scale examples used here, we clearly demonstrate
320 that the three inorganic P accessing extractants of the RBP method (CaCl_2 , citrate and HCl)
321 all correlate to some extent with Olsen P, but each provides insight into the source of the P;
322 soluble (directly available to roots and arbuscular mycorrhizas; Bolan, 1991), chelate labile
323 (available by the release of organic acids from roots and ectomycorrhizas; Jones and Darrah,
324 1994), or proton labile (release of H^+ by root tips and ectomycorrhizas; Römheld et al.,
325 1984). Enzyme extractable P, however, represents labile organic P (Tabatabai, 1994), a
326 component of soil P not effectively accessed by bicarbonate (Kuo, 1996) thereby explaining
327 the relatively weak factor loadings for enzyme extractable P compared to inorganic P
328 methods. The orthogonal correlation between soluble P by CaCl_2 extraction and HCl-
329 extractable P, and the proximity of other methods, supports the conclusion that CaCl_2 and
330 HCl access labile and recalcitrant forms of P, respectively. Inclusion of the environmental and
331 soil characteristics reveals that vegetation class is most strongly correlated with the PCA
332 ordination and it has a negative directional gradient.

333

334 *4.2. National scale changes in soil P status*

335 The final report from CS07 (Emmett et al., 2010) described a surprisingly large
336 decrease in mean Olsen-P concentration in all broad habitat types across the UK from 43 mg
337 P kg⁻¹ in 1998 to 32 mg P kg⁻¹ in 2007 (Table 2). The greatest change was seen in soil beneath
338 dwarf shrub heath, whilst the highest Olsen-P concentration and smallest significant change
339 was seen in arable soils. The RBP procedure described here effectively confirmed the
340 declining trend in inorganic available P described in the UK national survey, CS07 (Emmett
341 et al., 2010) and provided the clear pattern of increasing labile organic P. Therefore, the
342 observed decrease in inorganic P over a 10 year period does not specifically reflect a net loss
343 of P from the system; but rather demonstrates a noted change between pools of P from
344 inorganic to organic with the significant increase in enzyme extractable organic P. This is
345 seen across soils in all SOM categories and under all vegetation types to varying extents. As
346 there is overlap in the P pools quantified by each extractant this cannot be taken as the
347 average total available P value (in mg P kg⁻¹) across the UK. However, it does indicate there
348 is no net loss of P from UK soils. Further, our results suggest that the inorganic P is not
349 simply precipitating out into increasingly insoluble forms otherwise we would have observed
350 a smaller net decrease in the more stringent HCl extraction method where in reality, the
351 largest decrease in extractable inorganic came with the HCl extraction (e.g. Fig. 2).

352 The declining chelate and proton labile P could reflect consumption of residual P
353 without replenishment in the form of fertilisation (Withers et al., 2014). Chelate labile P
354 reflects P that is available to P-efficient plants whereas HCl labile is a gross proxy for proton
355 release at plant root tips (Jones, 1998; Hinsinger, 2001; Dakora and Phillips, 2002). Given
356 that the largest decreases are associated with grasslands (which have progressively been
357 receiving less P fertilization; 29.5 kg P ha⁻¹ in 1983 to <10 kg P ha⁻¹ in 2013; Defra, 2014)
358 suggests that plants harvested for fodder may be mining soil P reserves. The increasing
359 organic P across many categories of vegetation suggests that P is being taken out of the

360 mineral soil by plants and soil biota and is accumulating P in an organic form in litter and O
361 horizon organic matter. The organic P fraction can make up between 20 and 80% of total P
362 (P_i) in some soils (Dalal, 1977). A proportion of this will be easily hydrolysed (George et al.,
363 2002; Tang et al., 2006; Tarafdar and Jungk, 1987) and made available for plant uptake, but
364 the remainder is relatively stable and will remain occluded (Stewart and Tiessen, 1987). The
365 C density in the four SOM categories corroborates this theory; the patterns of increasing
366 organic P (Fig. 3c) and C density (Emmett et al., 2008) are very similar.

367 Increases in organic P in soil O horizon and litter may be attributed to increased
368 primary productivity due to several confounding environmental changes happening across the
369 UK over the study period. Increasing atmospheric nitrogen (N) deposition in the UK as
370 reported in a number of studies (e.g. Galloway et al., 2004; Stevens et al., 2006) has been
371 shown to increase primary productivity (Cannell et al., 1998) and consequently induce P
372 limitation through depletion of phytoavailable P. The increased uptake of inorganic P would
373 then be returned to the soil as organic P. Longer term increases in atmospheric CO₂
374 concentrations (IPCC, 2007) and temperature (Jones and Hulme, 1997) along with increasing
375 yields due to increasing N fertilization and use of improved hybrids (Jones et al., 2013) may
376 exacerbate the removal of labile and semi-labile inorganic P. Further, a decrease in external P
377 inputs may also be partly responsible for this shift in P status of UK soils. P fertiliser use on
378 grass and crops over the study period decreased by 40% and 35% respectively primarily due
379 to the increasing cost of P fertilizer (Defra, 2011).

380 The observed increase in soil pH reported in CS07 from 1998 to 2007 may also
381 contribute to the observed decrease in P associated with labile fractions. This soil pH
382 increased was particularly strong in soils with lower organic matter contents and soils with
383 neutral to alkaline pH (Emmett et al., 2010). With increasing pH in acidic soils one would
384 expect an increase in P solubility; however, an increase in the pH of alkaline/calcareous

385 would likely enhance precipitation of P as insoluble Ca-P (Samadi and Gilkes, 1999)
386 rendering the P unavailable to plants. However, the small degree of the change in pH makes it
387 unlikely that this represents the main driver of the change in P status with the exception of
388 microsite effects.

389 The lack of significant changes in any P pools in woodland habitats suggests that
390 more complex and successional advanced habitats were less susceptible to changes in soil P
391 status. Woodlands often express limited presence of soluble or labile P as nutrient
392 mineralization and solubilisation is balanced by nutrient uptake and immobilization
393 associated with litter fall and decomposition (Glenn-Lewin et al., 1992). It could also be that
394 the slower life histories associated with tree dominated habitats yield slower to responses to
395 shifts in nutrient inputs. For example, Cannell et al. (1998) modelled the response of conifer
396 forests to increasing N deposition, atmospheric CO₂ and temperature and predicts changes in
397 soil and plant response over decadal or century timescales. However, Shaw et al. (2002) and
398 Stevens et al. (2006) saw responses to similar parameters in grassland habitats in a matter of
399 months and years in both laboratory and field studies.

400 Given that British soils are relatively immature (ca. 10,000 years old; Avery, 1990), it
401 is likely that they are still undergoing the changes in form and amounts of P described by
402 Walker and Syers (1976). They describe soils reaching a terminal steady state at
403 approximately 22,000 years, before which occluded P and organic P increase at the expense
404 of more labile fractions. This can be seen to some extent in these results with the increase in
405 organic P fractions and decrease in labile fractions. However, the changes seen over the short
406 period studied here likely cannot be attributed wholly to pedogenic processes. Similar to the
407 CS results for Olsen P, there were no clear relationships between change in any P fraction and
408 2007 values for soil pH, SOM, moisture content, or with change in soil pH and SOM between
409 1998 and 2007.

410 Drying of soils prior to extraction has been shown to increase P solubility (Turner and
411 Haygarth, 2001; Styles and Coxson, 2006). The evaluation of moist and dry soil samples
412 from Hiraethlyn catchment in North Wales further demonstrates differences between the
413 Olsen method and the RBP method (Fig. 4) and indicates that use of fresh soils would be a
414 preferable approach for the RBP method. This is consistent with the findings of Styles and
415 Coxson (2006) which demonstrated an increase in extractable P with drying as a result of
416 destabilization of soil organic matter. Turner and Haygarth (2001) suggested that rewetting of
417 dried soils released P from the lysing of microbial cells and questioned the use of soil P
418 analyses that did not take soil moisture into account. In this study, we used air dried soils that
419 had been previously collected and archived as part of the CS; however, in future efforts, we
420 would recommend using this method with field moist soils and correcting to dry weight based
421 on soil moisture content.

422 Finally, it is important to note that we observed a great deal of variation in the P
423 content of batches of phytase enzyme reagent and found the some batches to be highly
424 contaminated with P. This required extensive dilution which compromised the overall assay
425 or pre-analysis treatment of the enzymes with dialysis membranes, a time consuming step.
426 We recommend only using acid phosphatase for the enzyme component of the assay.

427

428 *4.3. Conclusions*

429 Soil P transformations occur over both a dynamic, rapid biological cycle and a much
430 more gradual pedogenic cycle. Further, plant community directly influences P availability
431 making a single extraction approach inappropriate for natural or seminatural settings with
432 diverse plant assemblages. Given the limited solubility of P and its propensity to adsorb to
433 organic and mineral surfaces, almost all plants have evolved to develop specialized
434 mechanisms for enhancing P acquisition from soil. Therefore, measurement of P across

435 landscapes using a single extraction technique is likely to generate artefacts and will not
436 adequately reflect P bioavailability. The exhaustive, repeated sampling of CS offers an
437 invaluable opportunity to assess shifts in soil conditions at the national scale. The use of the
438 single solution bicarbonate method (Olsen P) for assessing soil P status does not adequately
439 evaluate the P status of soils in the UK. The RBP method has great promise for this type of
440 survey by providing a simultaneous assessment of biologically available P through the use of
441 four accepted P methods: 1) Soluble or solution P; 2) Enzyme extractable organic P; 3)
442 Chelate extractable P; 4) Proton extractable inorganic P. This suite of P extraction methods
443 offers a great deal of insight into changes occurring across diverse landscapes. The RBP
444 method proposed here has the potential to greatly improve our ability to characterise the soil
445 P status across complex landscapes. The RBP method is relatively quick (full assessment of
446 four P pools on ~56 soils in a day), inexpensive, and requires no specialist equipment making
447 P fractionation more accessible and feasible for large scale studies. It has proved accurate and
448 reliable on soils with a range of characteristics.

449 Future national surveys such as the UK Countryside Survey will help shed light on
450 whether this is a temporary change in P status in UK soils or a continuing trend. Whichever is
451 found to be the case, it is not necessarily a worrying phenomenon. Soils in the UK are
452 typically enriched in P which can cause eutrophication of water bodies (Withers et al., 2000).
453 If this is removed from the soluble and labile inorganic phase and stabilised in the organic
454 fraction it might have positive implications for water quality without greatly altering long-
455 term P fertility. Simultaneously, agricultural P fertilizer costs are climbing with increasing
456 limitation of minable P resources which makes plant P acquisition strategies that much more
457 important when assessing P availability. The long-term change in P pools observed herein
458 may also have implications for vegetation community structure and ecosystem dynamics

459 especially in a changing climate where community composition is likely to change in semi-
460 natural ecosystems.

461

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471

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660 southern China. *Environmental Earth Sciences* 67, 1725-1734.

Figure legends

661 **Fig. 1.** Principle component analyses (PCA) of the four P analysis methods of the rhizosphere
662 based P (RBP) extraction regime and the conventional Olsen P method as determined for 102
663 soil samples collected in 1998 in the UK Countryside Survey (Emmett et al., 2010).
664 Observations are displayed by grey dots, in relation to the loadings, P methods are displayed
665 as blue arrows and environmental or soils characteristics by red arrows.

666

667 **Fig. 2.** Mean change between 1998 and 2007 in P content (mg kg^{-1}) in (a) CaCl_2 , (b) citrate,
668 (c) enzyme, and (d) HCl extract fractions of soils collected from different ecosystem types
669 within the UK. Values indicate means \pm SEM. Asterisks indicate significant differences
670 between years (* $P < 0.05$, ** $P < 0.01$).

671

672 **Fig. 3.** Mean change between 1998 and 2007 in P content (mg kg^{-1}) in (a) CaCl_2 , (b) citrate,
673 (c) enzyme, and (d) HCl extract fractions within soils of differing soil organic matter status
674 within the UK. Values indicate means \pm SEM. Asterisks indicate significant differences
675 between years (* $P < 0.05$, ** $P < 0.01$, *** $P < 0.001$).

676

677 **Fig. 4.** Relationship between Olsen P content and the four fractions of the proposed
678 rhizosphere trait-based method for field-moist soils collected from within the Hiraethlyn
679 catchment in North Wales. (a) Olsen P vs. CaCl_2 ; (b) Olsen P vs. citrate; (c) Olsen P vs.
680 enzyme; (d) Olsen P vs. HCl extract. Lines and associated r^2 values are linear regression fits
681 to the experimental data.

682

683 **Fig. 5.** Relationship between field-moist and air dried soils for the four soil extractions, (a)
684 CaCl_2 ; (b) citrate; (c) enzyme; (d) HCl of the proposed rhizosphere trait-based method for

685 collected within the Hiraethlyn catchment in North Wales extract. Lines and associated r^2
686 values are linear regression fits to the experimental data.

687

Table 1

Mean concentration of P (mg kg^{-1}) solubilised by 10 mM CaCl_2 , 10 mM citric acid, 0.02 enzyme units of phosphatase and phytase enzymes, and 1.0 M HCl across 102 soil samples collected both in 1998 and 2007 in the UK Countryside Survey.

Extract	1990	2009	Progression
CaCl_2	33 ± 6^a	19 ± 3^b	Decrease
Citrate	285 ± 26^a	188 ± 26^b	Decrease
Enzyme	130 ± 28^b	291 ± 31^a	Increase
HCl	572 ± 40^a	399 ± 34^b	Decrease
Total, sum of averages	903 ± 16	897 ± 15	No change

Data represent means \pm SEM, $n = 102$. Different letters following numeric means indicates significant ($P < 0.05$) change in P between 1990 and 2009.

Table 2

Pearson correlation matrix for P solubilized using the Olsen bicarbonate method and the 4 extractants used in the rhizosphere-based P fractionation procedure (10 mM CaCl₂, 10 mM citric acid solution, 0.02 enzyme units of phosphatase and phytase enzymes, and 1.0 M HCl).

	Olsen	CaCl ₂	Citrate	Enzyme	HCl
Olsen	1.000				
CaCl ₂	0.372**	1.000			
Citrate	0.563**	0.153	1.000		
Enzyme	0.145	0.143	0.169	1.000	
HCl	0.432**	0.013	0.732**	0.18*	1.000

Significance indicated by asterisks, * $P < 0.01$, ** $P < 0.001$ ($n = 204$).

Figure 1

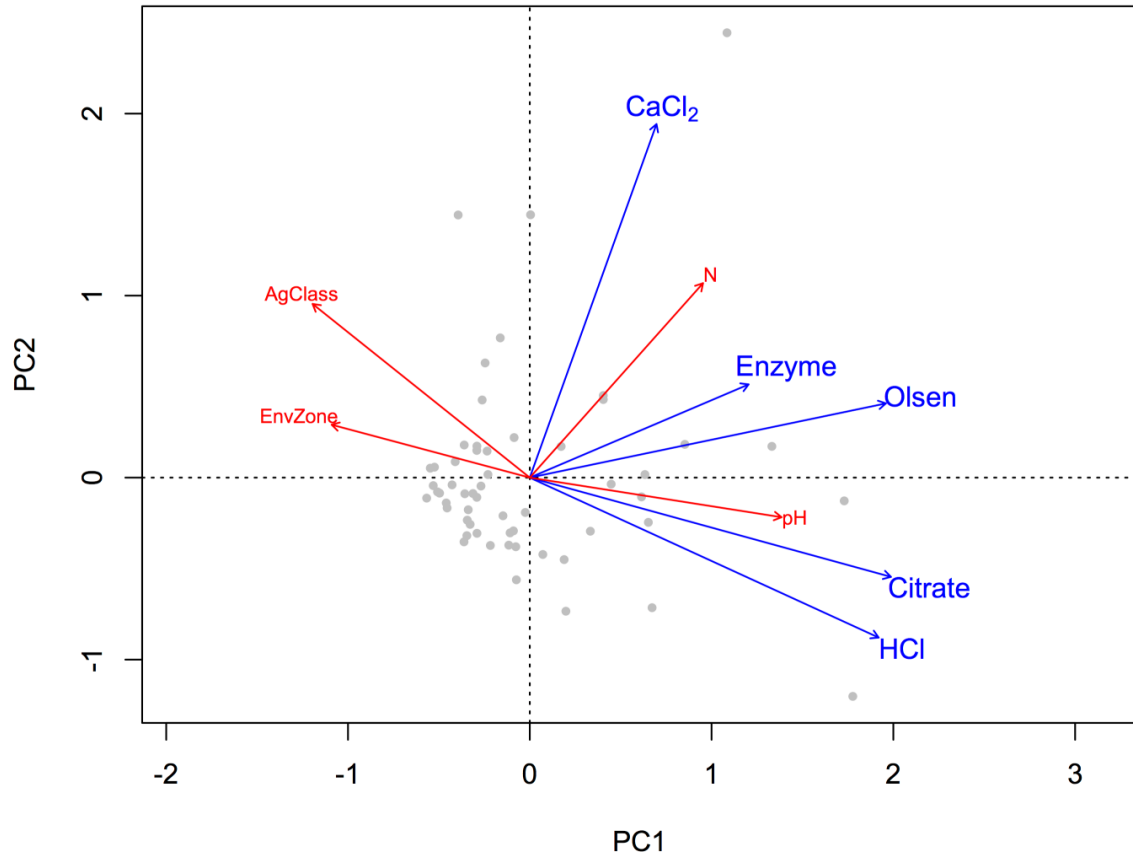


Figure 2

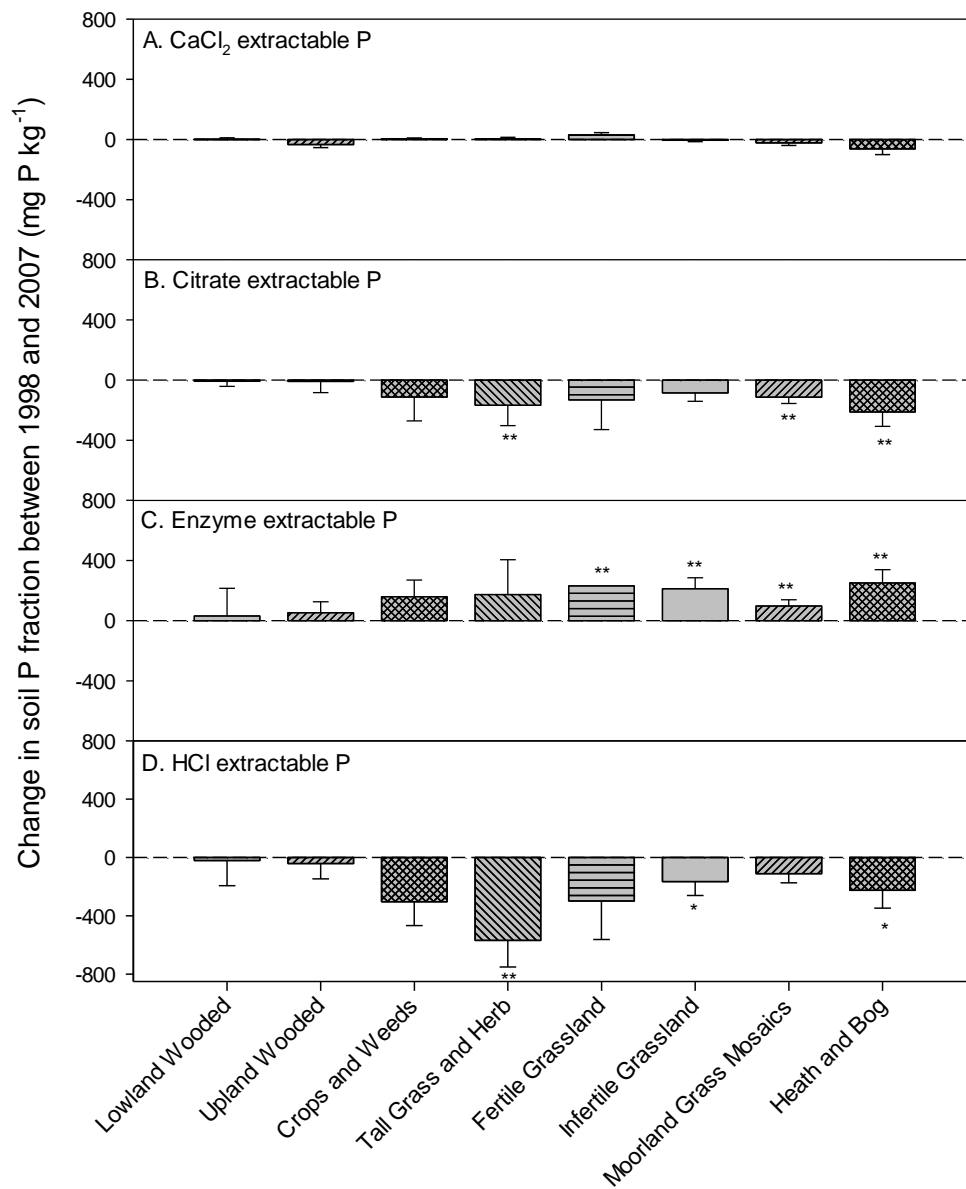


Figure 3

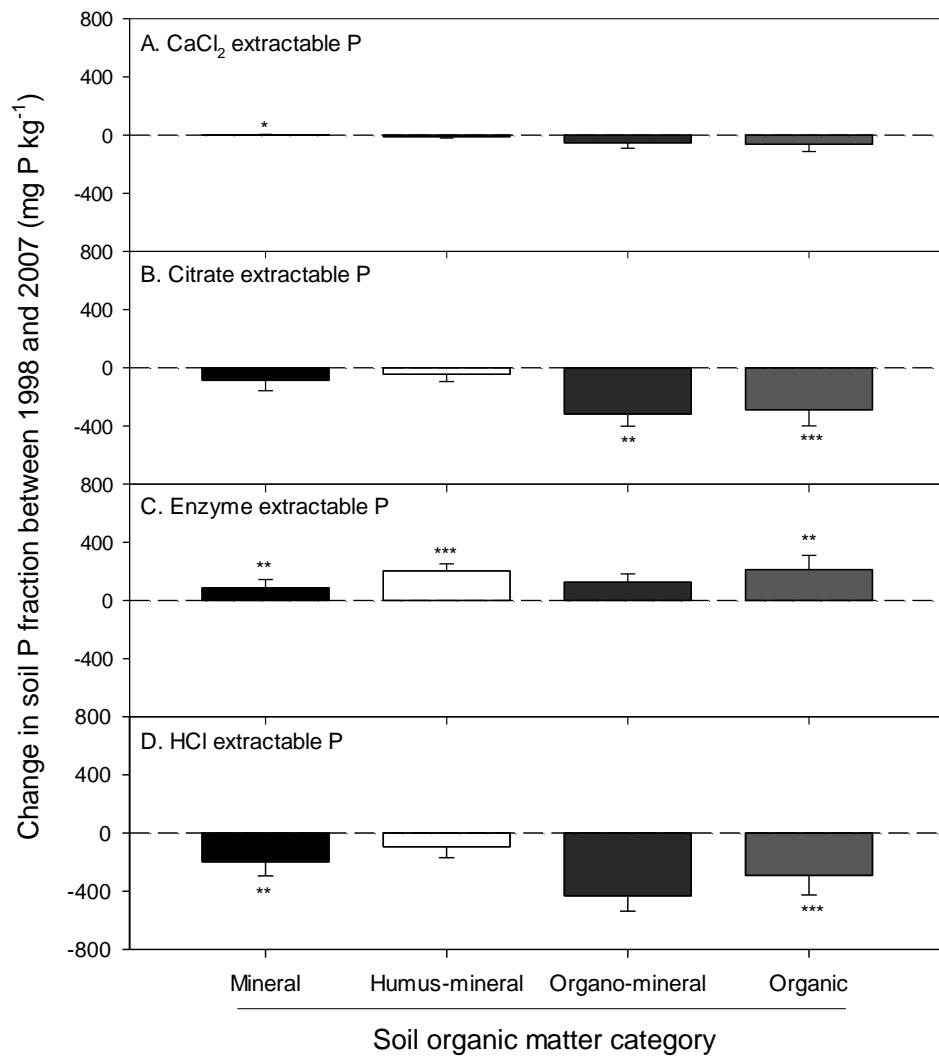


Figure 4

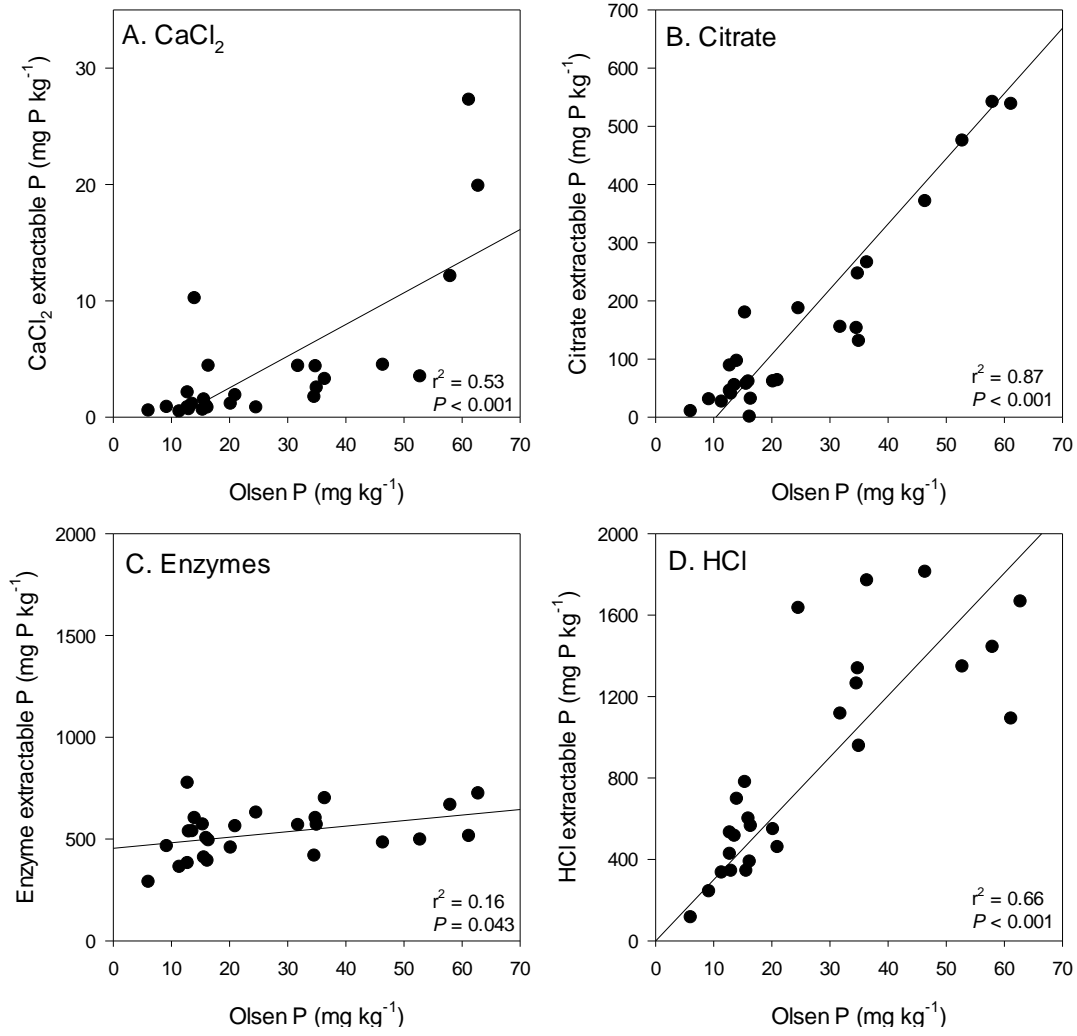


Figure 5

