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1 **Forms, patterns and extractability of phosphorus retained in**
2 **alum sludge used as substrate in laboratory-scale constructed**
3 **wetland systems**

4

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25 **Abstract**

26 This study examined the form, pattern and extractability of phosphorus (P) retained
27 in alum sludge (an aluminium-containing drinking water treatment residual in
28 dewatered cake form), which was used as substrate in laboratory scale constructed
29 wetland systems for P-rich wastewater treatment. Used alum sludge samples from
30 successive depth ranges in the laboratory scale vertical flow constructed wetland
31 systems were examined for KCl extractable P. The samples were also sequentially
32 fractionated into two main categories consisting of readily available P and P forms that
33 are not easily decomposed. The extracted fractions included labile P, microbial P, (Fe +
34 Al) P, humic P, (Ca + Mg) P and residual P. A major proportion of P retained in the
35 used alum sludge was in forms that are not easily decomposed. Of the P forms that are
36 not easily decomposed, the P associated with Ca and Mg accounted for a higher
37 proportion of the inorganic fraction as compared to the P associated with the Fe and Al.
38 The results also show that in most cases, the concentration of the P forms decreased
39 with increasing depth from the topmost surface of the alum sludge in the systems.

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43 **Key words:** alum sludge, constructed wetlands, labile phosphorus, phosphorus,
44 sequential fractionation, phosphorus extraction

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48 **1 Introduction**

49 The primary phosphorus (P) removal mechanisms in constructed wetland (CW)
50 systems are microbial anabolism, wetland plant uptake and substrate adsorption.
51 However, substrate P adsorption represents the main P removal pathway. Hence,
52 emphasis is being placed on the use of specialized P removal substrates with high P
53 removal capacity. Upon the saturation of such substrates with P, the investigation of the
54 form, pattern and extractability of P retained in them becomes very crucial. This will
55 help in determining the reuse potential of the saturated substrates, plant availability and
56 possible recovery of the adsorbed P. Several studies have examined the form of P-
57 association with elements contained in used CW substrates, but few have identified the
58 form and fate of the retained P. In two separate studies, Sakadevan and Bavor [1] and
59 Johansson [2] showed that large amount of P sorbed onto the surface of blast furnace
60 slag is predominantly associated with light elements, such as calcium, aluminium,
61 magnesium and silicon, with calcium hydroxide found to adsorb the largest amount of
62 P. In other studies, it was reported that P sorbed to blast furnace slag, which had been
63 previously used for wastewater treatment, was readily available for barley growth in a
64 pot experiment [3].

65 In other studies, the variation of P forms with depth was examined. Qualls and
66 Richardson [4] reported that the concentration of most forms of P decreased with
67 distance along the gradient. It was further reported that for most P forms, the trends are
68 weaker or absent at lower soil depths. The insoluble organic P, the humic P, and the
69 Ca-bound P which are all P forms that are not easily decomposed were reported to
70 show the highest decrease in concentration with distance. This observation is also
71 corroborated by the results of Reddy et al. [5] in which it was reported that the
72 concentration of most P forms were higher in the surface layers and decreased with

73 depth and with distance from the inflow structure. Such examination of P adsorbed in a
 74 CW substrate can also be used in evaluating the availability and potential recovery of P
 75 accumulated onto the CW substrate. The P examination is usually determined by
 76 sequential extraction with acid and alkaline reagents as modified for wetland soils by
 77 Qualls and Richardson [4] and Reddy et al. [5]. The P forms in various groups are
 78 usually identified and the form of P will influence its availability and recycling capacity.
 79 It should be noted that based on the sequential extraction, the term availability refers to
 80 both extractable and non-extractable P forms. The non-extractable P implies a form of
 81 P that is also available but in a form that cannot be extracted. Table 1 shows the
 82 different P forms including those that are grouped based on their availability for plant
 83 uptake. The classification is adopted from Qualls and Richardson [4]
 84

Table 1 Forms of extracted phosphorus

1. Forms readily cycled or available to plants (labile)	
Inorganic	Organic
➤ Exchangeable inorganic P (bicarbonate extractable)	➤ Microbial tissue P (chloroform-released P) ➤ Bicarbonate extractable organic P
2. Forms not easily decomposed (refractory)	
Inorganic	Organic
➤ Calcium bound P (soluble in 1M HCl)	➤ Humic organic P (organic P bound to humic and fulvic acids extractable in 0.1M NaOH)
➤ Iron and aluminium bound inorganic P (0.1M NaOH-extractable inorganic P)	➤ Residual (insoluble P)
➤ Iron and aluminium bound inorganic P occluded in particle matrix (0.1M NaOH-extractable inorganic P after sonication)	

85 However, it is also important to note that different CW substrates may have
86 specific properties that may be related to the P dynamics. Hence, it is necessary to carry
87 out individual examination of P forms for different substrates. In this study, alum
88 sludge was used as substrate in laboratory scale constructed wetland systems. Alum
89 sludge, an aluminium-containing drinking water treatment residual in dewatered cake
90 form, has been revealed as a new potential CW substrate due to its high P adsorption
91 capacity [6], and physico-chemical characteristics which conform to the recommended
92 standards for CW substrates [7]. The alum sludge (in dewatered cake form) was used as
93 main substrate in laboratory scale CW systems and it was shown to significantly
94 enhance P removal [8]. The study of beneficially reusing alum sludge as raw material is
95 part of a larger project aimed at the eventual development of a novel alum sludge based
96 CW system for wastewater treatment. The project has been carried out for the past five
97 years at the University College Dublin (UCD), Ireland and it has entailed series of sub-
98 projects of different scales. The sub-projects have included: (a) extensive studies on
99 identifying the P adsorption ability of the dewatered alum sludge and the removal
100 efficiencies of other pollutants (COD, BOD₅) using alum sludge as main substrate in
101 the novel CW system [6,8,9]; (b) investigating the lifespan of the system [9]; (c)
102 examining the mechanism and characteristics of P adsorption onto the alum sludge [10]
103 and (d) examining the release of other materials/elements from the alum sludge. As
104 such, the current study is aimed at further gaining a holistic view of the development of
105 the novel alum sludge based constructed wetland system by examining the fate of the P
106 adsorbed by the alum sludge, alongside other sub-projects as highlighted above.

107 Examination of the fate of P adsorbed in the alum sludge is crucial in determining
108 the recycling potential of the alum sludge when it is saturated and also in evaluating the
109 potential for recovery of the adsorbed P. Therefore, the objectives of this paper are to

110 address the following concerns: (i) in what forms is the adsorbed P stored in the used
111 alum sludge substrate (ii) is the P adsorbed stored in rapidly recycled forms or in forms
112 that are resistant to decomposition and plant uptake; (iii) does the concentration of P
113 forms increase or decrease with increasing depth from the topmost surface of the alum
114 sludge in the lab-scale CW systems.

115

116 **2 Materials and methods**

117 **2.1 Constructed wetland systems and used alum sludge samples**

118 The simulated CW systems from which the used alum sludge samples were
119 collected are shown in Fig. 1. The systems were made from Pyrex columns (1.0m long,
120 $\text{\O} = 0.095\text{m}$) and they were unplanted. The dewatered alum sludge was taken from the
121 Ballymore Water Treatment Plant in Southeast Dublin, Ireland and its characteristics
122 have been detailed in Yang et al., [10]. The sludge depth in the two systems was 0.6m.
123 Similar configuration utilizing columns to depict laboratory scale CW systems are used
124 in such laboratory scale studies [11]. The two systems were fed with wastewater from
125 the same feed tank. The wastewater used was collected periodically from an animal
126 farm, located in Lyons Estate, Newcastle, Dublin. Influent P into the systems ranged
127 from 9.8 ± 2.1 mg-P/l to 309 ± 27.9 mg-P/l. It should be noted that the wastewater
128 collected in the feed tank was sometimes diluted with tap water or spiked with artificial
129 P-rich solution (prepared using potassium dihydrogen phosphate (KH_2PO_4) salt) to
130 obtain desired operational P concentration before it is pumped into the systems. The
131 wastewater pumping operation into the two systems was carried out intermittently, i.e.
132 in batch strategy, and this was realised using peristaltic pumps connected to each
133 individual systems. The pre-set digital electronic timers ensured that the pumping
134 operation was done in batches according to the intermittent feeding strategy design.

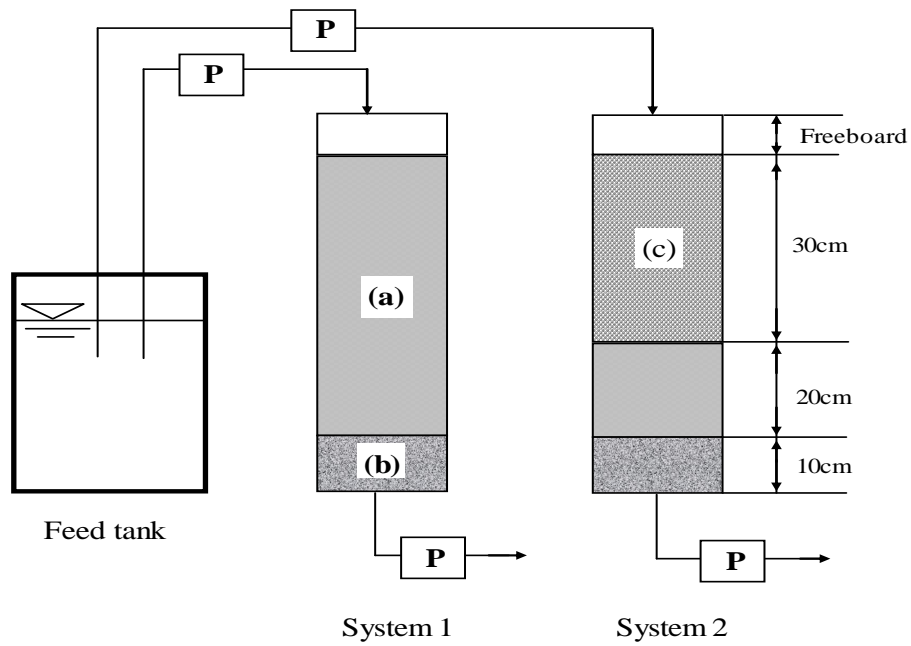


Fig. 1 Schematic diagram of the unplanted laboratory-scale alum sludge based CW systems. P = timer controlled peristaltic pumps, a = dewatered alum sludge cakes, b = gravel layer for support, c = pea gravel

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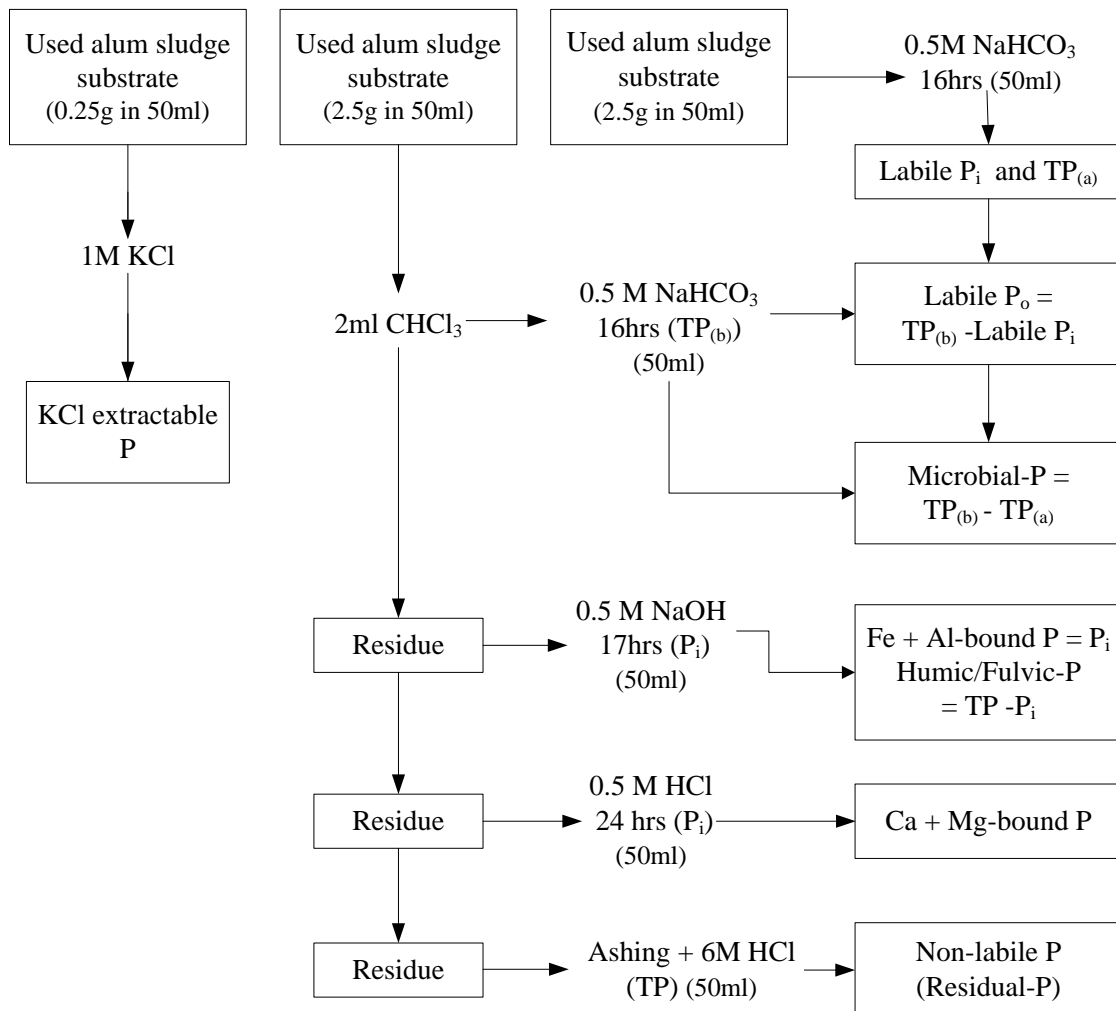
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137 A typical intermittent feeding consists of a ‘fill’ and ‘drain’ period. The fill period
 138 refers to the time of contact of the wastewater with the alum sludge in the system, while
 139 the drain period refers to the pumping out of the wastewater from the system. System 1
 140 was filled entirely with the dewatered alum sludge whereas system 2 had a different
 141 substrate configuration. System 2 was firstly filled with the dewatered alum sludge up
 142 to a depth of 20cm and then topped with 30cm of pea gravel. The experiment was
 143 conducted over a total period of 25 weeks. At the end of the experimental period, the
 144 systems were gently dismantled and the used alum sludge samples were collected at
 145 every 10cm depth increment from the top of the alum sludge in the systems and
 146 sequential P extraction was carried out. Note that in system 2, the used alum sludge
 147 samples were collected from depth range of 30-40cm and 40-50cm.

148 Sequential P extractions were then carried out. The sequential P extraction
 149 procedures used in this study were modified from those developed for subsurface flow
 150 CW substrates by Forbes et al. [12]. In each step of the fractionation scheme, both
 151 organic and inorganic P forms were extracted. The extraction scheme used are
 152 discussed below and also depicted sequentially in the extraction scheme shown in Fig.
 153 2.

154

155



156

Fig 2. Scheme of sequential phosphorus extraction procedure for used alum sludge substrate

157 **2.2 Sequential P extraction**

158 **2.2.1 KCl extractable P**

159 KCl extractable P was determined by modifying the method proposed by Makris and
160 O'Connor [13] and used for the alum sludge. 0.25g of the alum sludge was equilibrated
161 with 50ml of 1M KCl for 2 hours using a rotary shaker. Thereafter, the mixture was
162 centrifuged and then filtered through a 0.45µm membrane filter. The filtrate was
163 acidified to pH < 2 and then analysed colorimetrically for soluble reactive P using a
164 HACH DR-2400 spectrophotometer according to standard methods [14].

165 **2.2.2 Labile inorganic (P_i) and organic (P_o) phosphorus forms**

166 In order to determine the different fractions of the labile P_i and P_o, duplicate
167 samples (2.5 g each) were used. The first sample was fumigated with about 2-5ml of
168 chloroform to lyse the microbial cells, and then extracted with 0.5M NaHCO₃. The
169 mixture was then mechanically shaken using a mechanical shaker for 16 hours. After
170 this, the samples were analysed for Total P (TP) by ashing samples at 550°C for 4 h
171 followed by extraction with 6M HCl [12]. The extract was then analysed for TP
172 following sulfuric acid-potassium persulfate digestion, using an autoclave. All P forms
173 were determined analytically using a HACH DR-2400 spectrophotometer. The second
174 set of the samples was extracted with 0.5M NaHCO₃, but these samples were not
175 fumigated with chloroform. The samples were shaken for 16 hours, filtered using the
176 whatman no. 41 filter and analysed for both SRP (soluble reactive phosphorus) and TP.
177 The SRP determined is referred to as labile P_i. It should be noted that this step is the
178 first stage in the entire sequential extraction scheme. The residue obtained from this
179 particular set of extraction was used in subsequent extraction as explained in sections
180 2.2.3 to 2.2.5.

181 The rationale for bicarbonate extraction of the P_i is that soil/samples that contain
182 levels of calcium have been shown to have decreased level of calcium activity in
183 solution by precipitation as $CaCO_3$, and therefore, P extracted using this method is
184 found to be effective in determining plant-available P [5]. The derivation of labile P_o
185 (living and dead microbial biomass) and the labile P_i can be seen diagrammatically in
186 Fig. 2. Specifically, microbial mass is calculated as the difference between TP in the
187 extracts from chloroform treated and untreated samples [15], while total labile organic
188 P is calculated as the difference between TP in solution extracted with $NaHCO_3$ from
189 samples subjected to chloroform fumigation and P_i extracted from soils without
190 chloroform [5].

191 **2.2.3 NaOH extractable P**

192 The residue of the samples extracted as described in section 2.1.2 was treated with
193 0.5M NaOH and allowed to equilibrate for 17 hours on a mechanical shaker. This was
194 followed by centrifugation and filtration and the filtered solution was analysed for SRP
195 and TP. These fractions are referred to as NaOH- P_i and NaOH-TP respectively. The
196 NaOH- P_i is considered to represent iron and aluminium bound P (Fe- and Al-bound P).
197 The difference between NaOH-TP and NaOH- P_i is taken as organic P (NaOH- P_o),
198 which is associated with fulvic and humic acids [5].

199 **2.2.4 HCl extractable P**

200 The residue from the above extraction step was further treated with 0.5M HCl and
201 equilibrated for 24 hours on a mechanical shaker. This was also followed by
202 centrifugation and filtration and the filtered solution was analysed for SRP, while the
203 residue was kept for the next extraction step. The HCl- P_i fraction is taken as calcium
204 and magnesium bound P (Ca- and Mg- bound P).

205 **2.2.5 Residual phosphorus and total phosphorus**

206 The residue from the above extraction step was combusted at 550°C for 4 hours.
207 The ash was then dissolved in 6M HCl and then analysed for TP. The result was
208 referred to as non-labile P (i.e. residual P).

209 **3 Results and discussion**

210 **3.1 KCl extractable P**

211 The result of the KCl extractable P is shown in Fig. 3. It can be seen that the alum
212 sludge had an initial KCl extractable P content ranging from 0.08 to 0.10 mg-P/g and
213 this increased to between 0.08 to 0.17 mg-P/g in the used alum sludge substrate. From
214 Fig 3, it can also be seen that the greatest increase was in the topmost surface of the
215 alum sludge in both systems (i.e. 0-10cm in system 1 and 30-40cm in system 2). Even
216 though the used alum sludge was considerably loaded with P during the operating
217 period, the general increase in the KCl extractable P content of the used alum sludge
218 can be considered marginal in comparison to the P loading. This suggests the strong P
219 binding ability of the alum sludge and also shows that the adsorbed P in the alum
220 sludge may not be easily available. KCl extractable P is the most available (labile) pool
221 of P [13, 16], which can be easily released (most loosely bound pool of P). However, it
222 does not differentiate among the different forms of P.

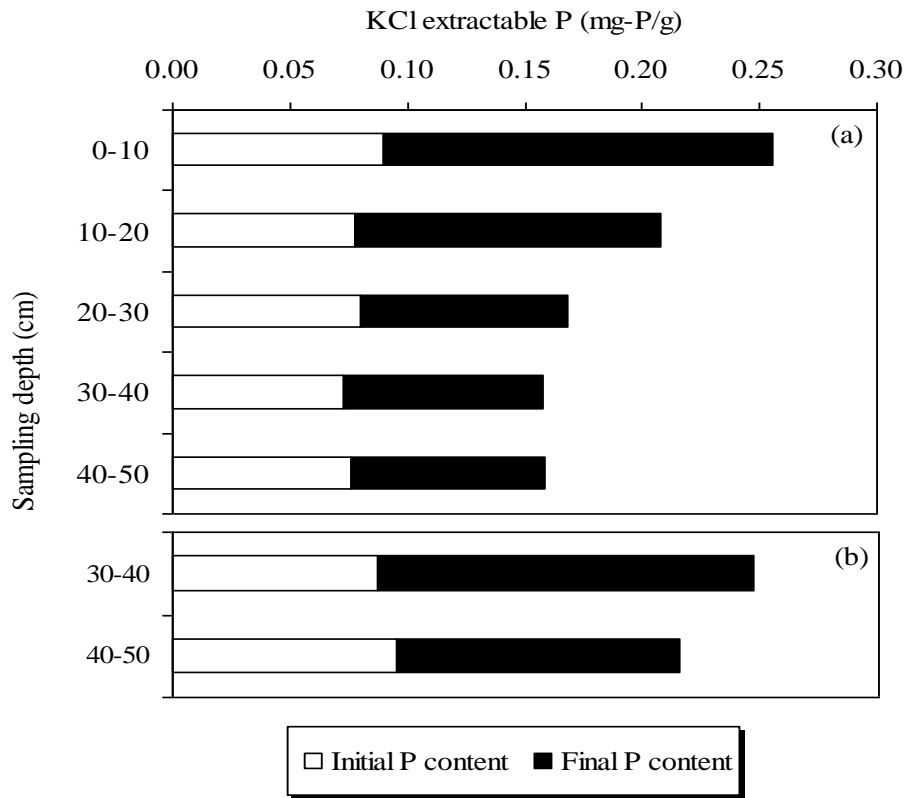


Fig 3 (a & b) Initial and final KCl extractable P in the used alum sludge from systems 1 and 2, respectively.

223

224 3.2 Sequential phosphorus fractionation

225 3.2.1 P forms that are readily cycled or available to plants

226 Fig. 4 shows the concentration of P forms that are readily cycled or available to
 227 plants at various depths in the two systems. This group consists of the exchangeable
 228 inorganic P (EIP) and organic P (microbial and bicarbonate extractable P). From the
 229 figure, it can be seen that the concentration of EIP was mostly highest at the topmost
 230 surface of the alum sludge in the systems. In particular, the result for system 1 shows
 231 that the concentration of EIP decreased with increasing depth from the topmost surface
 232 of the alum sludge. For system 1, the concentration of EIP decreased from 2.2 mg-P/g
 233 to 1.6 mg-P/g for depths ranges of 0-10cm and 40-50cm, respectively.

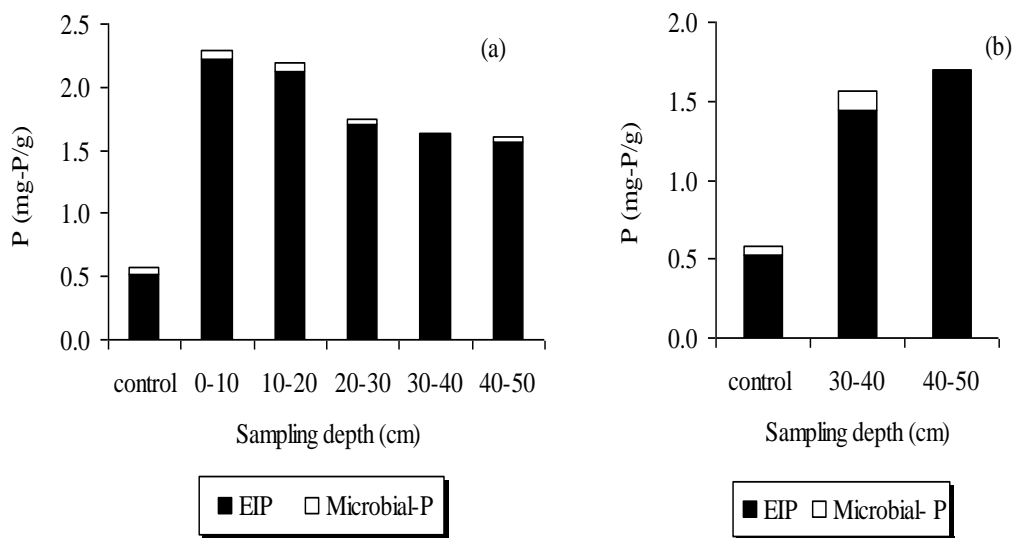


Fig. 4 Profiles of readily cycled/plant- available P in the used alum sludge samples (note: a and b refer to systems 1 and 2, respectively; control refers to the unused alum sludge).

234

235 The high concentration of EIP close to the top surface of alum sludge in the systems
 236 suggests that the inorganic P may be loosely bound to organic matter, bound to solid
 237 phases such as CaCO_3 , or present as recently precipitated, amorphous, monocalcium
 238 phosphate [5]. However, of the readily cycled/plant-available forms of P, the EIP
 239 accounted for the largest proportion stored in the used alum sludge samples. The
 240 readily cycled or plant available forms of P are classified into exchangeable inorganic P
 241 (extracted with NaHCO_3) herein referred to as EIP; the organic microbial tissue-P
 242 (chloroform released and extracted with NaHCO_3) and the exchangeable organic P
 243 (extracted with NaHCO_3), herein referred to as EOP [4]. The EIP represents free SRP
 244 in soil water, plus the weakly adsorbed inorganic P on surfaces. This pool also
 245 represents an immediately available pool of P, and a buffer regulating solution

246 concentration. In comparison, the microbial P accounted for only a very little
247 proportion of the readily cycled/ plant-available P as can be seen in Fig. 4. Furthermore,
248 has been noted that the microbial biomass P is not deposited in the same sense as other
249 forms of organic P, but it may represent a larger pool in enriched areas [4]. In the two
250 laboratory scale constructed wetland systems studied, the only contribution to
251 microbial-P would have come from the biomass, but this amount is very small,
252 compared to the EIP, and it does not represent a substantial pool for P storage under the
253 current conditions.

254 Also from Fig. 4, it can be observed that the concentration of EIP and the
255 microbial-P forms decreased with increasing depth from the surface in most cases. The
256 results also indicate that the high concentration of soluble P used in the influent
257 wastewater into the systems promoted the diffusion of initially adsorbed P into the
258 pores/interior in the alum sludge. Therefore, any labile inorganic P would likely be
259 converted from short-term to long-term storage by either precipitation or incorporation
260 into the interior structure of the alum sludge. Thus, most P would be in the forms that
261 are not easily decomposed compared to the readily cycled/plant-available form.
262 However, it should be noted that negative values were obtained for the exchangeable
263 organic P (EOP). This is because, analytically, EOP is calculated as the difference
264 between TP in solution extracted with NaHCO_3 from samples subjected to chloroform
265 fumigation, and P_i extracted from soils without chloroform [5, 12]. However, the
266 spiking of the influent wastewater with synthetic inorganic P greatly increased the P_i ,
267 such that its value was higher than the TP obtained by extraction with NaHCO_3 and
268 chloroform fumigation. This consequently led to the negative values.

269 **3.2.2 P forms that are not easily decomposed**

270 Fig. 5 (a & b) shows the relative percentage of each of the P forms that are not
271 easily decomposed across the sampling depth and also the total amount of these P
272 forms stored in each system and across the depths. The figure shows that across depth,
273 most of the P forms are stored in the inorganic form. In system 1, the range of
274 concentration of the inorganic P form across depth was 2.6-4.3 mg-P/g (for Fe- and Al-
275 bound P) and 2.2-6.9 mg-P/g (Ca- and Mg-bound P) while for inorganic P, the range
276 was 3.3-12.6 mg-P/g (for humic and fulvic acids bound P) and 1.4-2.0 mg-P/g (for
277 residual P). For system 2, the range across depth was 3.4 mg-P/g (relatively unchanged
278 for Fe- and Al-bound P) and 6.5-7.3 mg-P/g (for Ca- and Mg-bound P) while for
279 inorganic P, the range was 8.0-9.5 mg-P/g (for humic and fulvic acids bound P) and
280 0.6-0.8 mg-P/g (for residual P). Being that the alum sludge is chiefly composed of
281 aluminium, one may presumably expect a significant fraction of the inorganic P forms
282 that are not easily decomposed to be associated with the Fe and Al fraction.

283 According to Qualls and Richardson [4], four major processes can cause the
284 deposition of calcium bound-P: (a) chemical precipitation of calcium phosphates, (b)
285 coprecipitation with calcium carbonate, (c) adsorption and precipitation of phosphate
286 on the surfaces of calcium carbonate, or (d) precipitation by calcareous algae. It should
287 be noted that the influent wastewater used in these systems contains a significant
288 amount of calcium, which has been shown to be removed onto the alum sludge [17].
289 Consequently, the relative concentration of calcium in the sludge increased
290 significantly, while that of aluminium decreased. Although it is noted that acidic
291 conditions may favour formation of P bound to Fe and Al secondary minerals, while
292 more alkaline conditions favour the formation of Ca bound P [4], this is not likely to be
293 case in this study, as the pH values were circumneutral. In the case of the humic/fulvic
294 acid bound-P, there was no definitive trend observed. For instance, in system 1 (See Fig

295 5, b), humic/fulvic acid bound-P increased with increase in depth and distance from the
 296 topmost alum sludge surface up to the 20-30 cm depth, and then decreased and
 297 increased again between the 30-40 and 40-50 cm depth respectively.

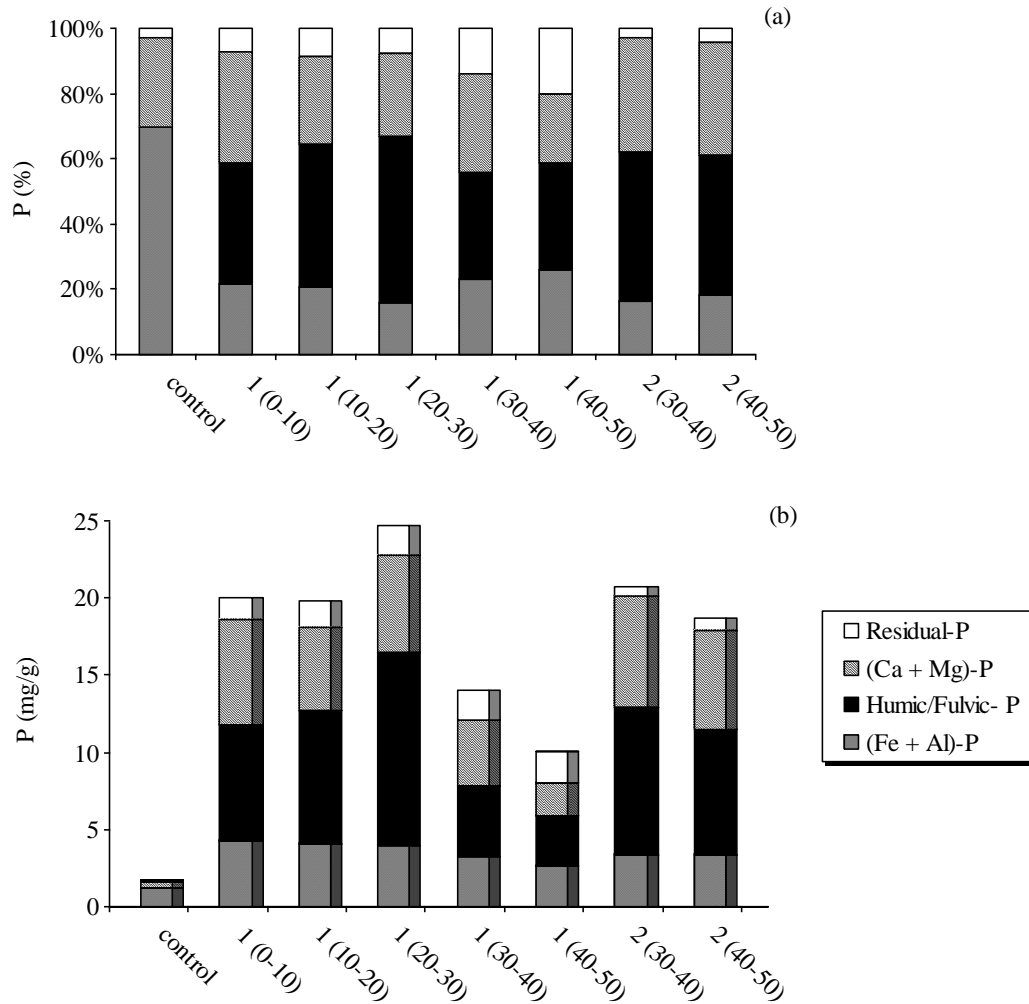


Fig. 5 Profiles of P forms that are not easily decomposed in the used alum sludge samples: (a) relative proportion in %; and (b) relative quantity in mg-P/g of the different proportions (note: 1 and 2 refers to systems 1 and 2, respectively, while values enclosed in bracket indicate the sampling depth range in cm. Control refers to the unused alum sludge).

298

299 However, for system 2, there were only two sampled points, and the humic/fulvic acid
 300 bound-P decreased with increasing depth from the topmost alum sludge surface.

301 Notwithstanding, the relatively high amount of P associated with the humic/fulvic acid

302 can be explained by the high organic content in the alum sludge. The percentage of the
303 residual P increased with depth away from the topmost surface (See Fig. 5, a). Again,
304 the fact that some amount of P is extracted as residual P shows that the systems
305 relatively functioned as a long-term P-sink.

306 Fig. 6 shows a comparative plot of the P forms that are not easily decomposed and
307 the readily cycled/plant-available P in the used alum sludge samples. It can be seen that
308 more P is held in forms that are not easily decomposed which can be viewed as a long-
309 term storage. On the other hand, P stored in the readily cycled/ plant-available form is
310 comparatively smaller in all cases. The total amount of P held in forms that are not
311 easily decomposed in systems 1 and 2 ranged from 10.1–24.7 mg-P/g and 18.7-20.7
312 mg-P/g respectively, while the total amount of P held in forms that are readily
313 cycled/plant-available ranged from 1.6-2.3 mg-P/g and 1.6-1.7 mg-P/g for systems 1
314 and 2 respectively. The P held in readily cycled/plant-available form is exchangeable
315 and according to Craft and Richardson [18], this pool should only be viewed as a
316 temporary storage or an intermediate pool and not as a long-term storage mechanism.
317 In most cases, the total amount of P stored in forms that are not easily decomposed
318 increased with distance from the inlet. This is particularly due to the fact that the
319 amount of residual P increased with distance from the topmost surface, and this
320 contributed to the total P retained in forms that are not easily decomposed. The residual
321 P, which is organic, is highly refractory and it may not be bioavailable [5]. The relative
322 proportion of this residual P generally increased with depth in the systems as can be
323 seen in Fig 5 (a).

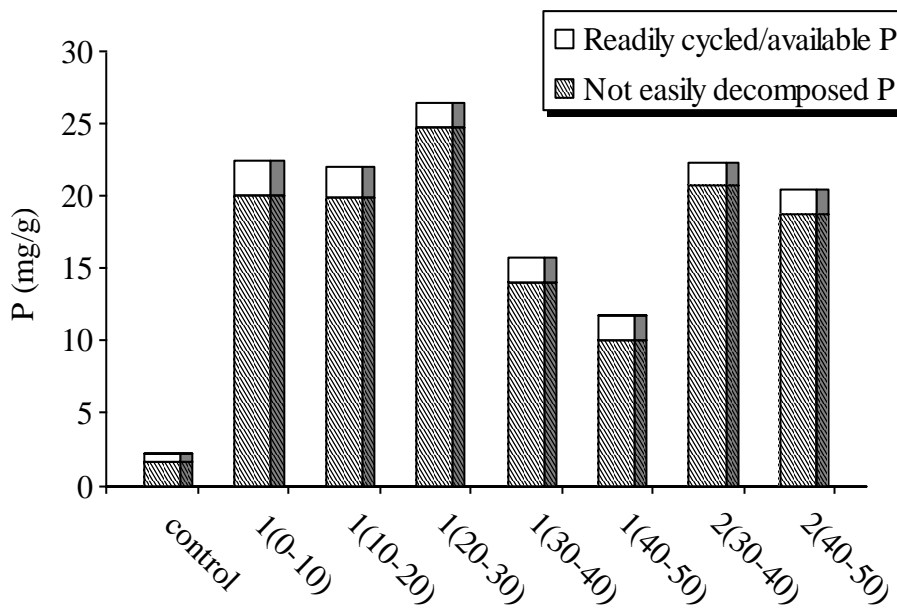


Fig. 6 Comparative profiles of P forms that are not easily decomposed and readily cycled/plant-available P in the used alum sludge samples (note: 1 and 2 refer to models systems 1 and 2, respectively, while values enclosed in bracket indicate the sampling depth range in cm. Control refers to the unused alum sludge)

324

325 **4 Conclusions**

326 The forms, patterns and extractability of dewatered alum sludge used as substrate
 327 in laboratory-scale constructed wetland system were examined in this study. Results
 328 obtained indicate that a major proportion of P retained in the used alum sludge samples
 329 were in forms that are not easily decomposed. The total amount of P held in forms that
 330 are not easily decomposed in the systems ranged from 10.1–24.7 mg-P/g while the total
 331 amount of P held in forms that are readily cycled/plant-available ranged from 1.6-2.3
 332 mg-P/g. Analysis of the P forms that are not easily decomposed across the two
 333 simulated constructed wetland systems indicate the following: In system 1, the range of
 334 concentration of the inorganic P form across depth was 2.6-4.3 mg-P/g (for Fe- and Al-
 335 bound P) and 2.2-6.9 mg-P/g (Ca- and Mg-bound P) while for inorganic P, the range

336 was 3.3-12.6 mg-P/g (for humic and fulvic acids bound P) and 1.4-2.0 mg-P/g (for
337 residual P). In system 2, the range across depth was 3.4 mg-P/g (relatively unchanged
338 for Fe- and Al-bound P) and 6.5-7.3 mg-P/g (for Ca- and Mg-bound P) while for
339 inorganic P, the range was 8.0-9.5 mg-P/g (for humic and fulvic acids bound P) and
340 0.6-0.8 mg-P/g (for residual P). Therefore, the results imply that P fractions associated
341 with Ca and Mg, and Fe and Al are the dominant forms of the P forms that are not
342 easily decomposed. The concentration of most of the P forms was also found to
343 decrease with increasing depth from the topmost surface of the alum sludge in the lab-
344 scale constructed wetland systems. The alum sludge showed an overall increase in its P
345 content and this could increase its agronomic value and potential for P recovery.

346

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352

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