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3 **Acid Phosphatases from the Breadfruit**

4 ***Artocarpus communis* Seeds as Novel Plant**

5 **Phosphorylating Biocatalysts**

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10 **ABSTRACT**

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Aims: Investigation on the phosphotransferase activity of two non-specific acid phosphatases (EC 3.1.3.2) designated as AP1 and AP2, previously isolated from breadfruit (*Artocarpus communis*) seeds for further biotechnological and industrial applications.

Methodology: Transphosphorylation reactions were tested with sodium pyrophosphate as the phosphoryl donor and phenol as its acceptor. Transfer products were quantified by using high performance liquid chromatography.

Results: The two acid phosphatases were able to catalyse phosphoconjugates synthesis using pyrophosphate as the phosphoryl donor and phenol as acceptor. The optimal conditions of transphosphorylation reactions indicated that this synthesis was highly dependent on pH, temperature, time course, donor and acceptor concentrations and enzyme amount. A very short period (1.25 h) was observed for these synthesis reactions catalysed by acid phosphatases isolated from breadfruit (*Artocarpus communis*) seeds. This suggested energy saving during biotransformation processes. The high average yields of 84.20 and 66.78% obtained for AP1 and AP2, respectively, made them useful to phosphorylate a wide range of nucleophile compounds such as nucleotides often used as food additives and pharmaceutical intermediates.

Conclusion: The acid phosphatase AP1 would be the most promising on the basis the better synthesis product yield (84.20%). The two biocatalysts could be considered as new valuable tools for bioprocesses.

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13 *Keywords: Plant acid phosphatases, transphosphorylation, breadfruit Artocarpus communis*

14 *seeds, phosphoconjugate synthesis, biocatalysts*

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17 **1. INTRODUCTION**

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19 Transphosphorylation is an efficient process for the large-scale production of new phosphorylated conjugates with improved properties [1, 2, 3]. This reaction is essentially reversible, and the equilibrium position depends on the conditions and the amount of reagents and products present in the reaction mixture [1]. Kinases are well-known phosphorylating enzymes which transfers a phosphate unit from ATP to a variety of acceptors, but the large-scale application is impeded by the need of regenerating ATP and in addition these enzymes are specific for the substrate to be phosphorylated [4]. Nowadays, it has been shown that acid phosphatases (orthophosphoric-monoester phosphohydrolase (EC 3.1.3.2) catalyzing the hydrolysis of a broad and overlapping range of phosphomonoesters are also able to carry out transphosphorylation reactions in which a phosphate unit is transferred from a donor (phosphomonoesters or pyrophosphate P₂i) to an acceptor alcohol [5]. This enzymatic phosphorylation process shows more advantages than

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31 the chemical one that uses phosphoryl chloride (POCl_3) [6]. It is well known that enzymes
32 tolerate environmental stress due to their broad substrate specificity, they are able to accept
33 a wide range of cheap phosphoryl donors and acceptors [6, 7]. In contrast, the chemical
34 methods should be explored with caution due to high number of by-products obtained in
35 typical reactions. Current chemical routes to phosphate esters synthesis often proceed via
36 the corresponding phosphate di or triesters and require harsh reagents and conditions
37 leading to hardly separable product mixtures [8].

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39 Thus, Bacterial phosphatases from the class A nonspecific acid phosphatase family were
40 mostly used, and *Shigella flexneri* [7, 9], *Salmonella enterica* [7], *Morgenella morgani* [5, 1]
41 and *Escherichia blattae* [2] were prominent sources of the enzyme. The phosphoryl
42 transferase activities of nonspecific acid phosphatases from these microorganisms were
43 exploited to produce nucleotides as inosine-5 monophosphate (5-IMP) and guanosine-5
44 monophosphate (5-GMP) using pyrophosphate (PPi) as a phosphate donor. The production
45 of phosphoconjugates is another key area of interest in biotechnology, because they have
46 often used as food additives and as pharmaceutical synthetic intermediates [5].

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48 Todate, there are few reports on the transphosphorylation abilities of plant non-specific acid
49 phosphatases. Nevertheless, due to their wider availability and lesser cost, plant
50 phosphatases could be explored as good substitutes for biotechnological phosphate ester
51 synthesis. Koffi et al. [3] isolated four non-specific acid phosphatases from the neglected
52 crop *Lagenaria siceraria* seeds exhibiting high phosphoryltransferase activities. In search of
53 new biocatalysts with improved transphosphorylation yield, two acid phosphatases from
54 breadfruit (*Artocarpus communis*) seeds [10] were investigated. In this paper, we report on
55 the phosphoryltransferase activity of these enzymes.

56 57 58 59 **2. MATERIALS AND METHODS**

60 61 **2.1 Enzymes**

62 The acid phosphatases **named** AP1 and AP2 (EC 3.1.3.2) used for transphosphorylation
63 reactions originated from the purified enzyme collection of Laboratoire de Biochimie et
64 Technologies des Aliments, UFR Sciences et technologies des Aliments, Université Nangui
65 Abrogoua (Abidjan, Côte d'Ivoire). These phosphatases were purified as described
66 previously [10].

67 68 **2.2 Chemicals**

69 Sodium pyrophosphate (donor), phenol (acceptor) and tyrosine (internal standard) were
70 purchased from Sigma Aldrich. All other reagents used were of analytical grade.

71 72 **2.3 Transphosphorylation reactions**

73 The ability of phosphatases AP1 and AP2 from *A. communis* seeds [10] to catalyze
74 transphosphorylation reactions was tested with sodium pyrophosphate as the phosphoryl
75 donor and phenol as its acceptor. In typical experiment, transphosphorylation reactions were
76 carried out at 37°C in a total reaction mixture of 250 μL containing 400 mM of sodium
77 acetate buffer pH 4.0. The reactions were stopped by immersion in boiling water for 3 min,
78 followed by cooling in ice bath. Prior to each HPLC analysis, the reaction mixtures were
79 filtered using Ultrafree-MC filter (0.45 μm) (Millipore). Tyrosine was used as the internal
80 standard to correct chromatographic product areas. Twenty microliter (20 μL) aliquots of

81 each reaction mixture always containing the internal standard (2 mM final concentration)
82 were analyzed quantitatively by HPLC. The column used was SPHERECLONE 5 μm ODS
83 (2) (250 mm \times 4.60 mm; Phenomenex) and phenolic compounds were detected at 254 nm
84 with a SPECTRA SYSTEM UV 1000 detector. The elution was done with a BECKMAN 114
85 M solvent delivery module pump, at a flow rate of 0.5 mL min^{-1} by using a degassed mixture
86 of acetonitrile/water in the ratio 50:50 (v/v) as mobile phase. The chromatograms were
87 obtained with a SHIMADZU C-R8A CHROMATOPAC V1.04 integrator. The detailed
88 experimental conditions for studying parameters likely to affect the transphosphorylation
89 reactions (pH, time, donor concentration, acceptor concentration and enzyme amount) are
90 given below. For the determination of each parameter, reactions were carried out in
91 triplicate.

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93 **2.4 Determination of optimum pH**

94 The pH values were determined at 25°C. For determination of optimum pH,
95 transphosphorylation reactions were performed by incubating at 37°C for 1 h each
96 phosphatase (25 μL) in a pH range of 3.6 to 5.6 (sodium acetate buffer, 400 mM), with 100
97 mM final concentration of sodium pyrophosphate and 5 mM final concentration of phenol.

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99 **2.5 Determination of optimum time**

100 To determine the optimum time of transphosphorylation, the optimum pH determined for
101 each enzyme was fixed. The transphosphorylation reactions were performed at different
102 times ranging from 0 to 6 h (at 37°C), in 400 mM sodium acetate buffer at appropriate
103 optimum pH, with 25 μL enzyme, 100 mM final concentration of sodium pyrophosphate and
104 5 mM final concentration phenol.

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106 **2.6 Determination of phosphoryl donor optimum concentration**

107 The influence of phosphoryl donor concentrations (0 to 300 mM) on the transphosphorylation
108 reactions was determined under the optimum conditions of pH and time. Phosphatases (25
109 μL) were separately incubated at 37°C, with different concentrations of sodium
110 pyrophosphate ranging from 0 to 300 mM final concentration and 5 mM phenol.

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112 **2.7 Determination of phosphoryl acceptor optimum concentration**

113 The influence of phosphoryl acceptor concentrations (0 to 10 mM) on the
114 transphosphorylation reactions was determined under the optimum conditions of pH, time
115 and phosphoryl donor concentration. The phosphatases (25 μL) were separately incubated
116 at 37°C, with different concentrations of sodium pyrophosphate ranging from 0 to 300 mM
117 final concentration and 5 mM phenol.

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119 **2.8 Determination of enzyme amount**

120 For this study, the optimal condition of pH, time, donor concentration and acceptor
121 concentration determined for each enzyme were fixed. Only the amount of each
122 phosphatase varied from 0 to 10 μg . In the determinations described above, 25 μL of
123 enzyme corresponded to 5 μg , and 2.1 μg of protein, respectively, for AP1 and AP2. All the
124 reactions were stopped by immersion in boiling water for 3 min and the products quantified
125 by HPLC as described in the typical transphosphorylation reaction.

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2.9 Transphosphorylation reactions in optimum conditions

130 Ultimately, the optimal conditions of pH, enzyme amount, donor and acceptor concentrations
131 and time were fulfilled to perform a unique transphosphorylation reaction with phosphatases
132 AP1 and AP2 purified from *A. communis* seeds. These reactions were also carried out in
133 triplicate with regard to the typical conditions described in the other experiments, and the
134 synthesized products were quantified by HPLC as described previously.

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2.10 Estimation of the yield of transphosphorylation

137 One mol of the synthesized product corresponds to 1 mol of phenol used as acceptor in the
138 reaction mixture. Therefore, the starting phenol concentration deficit at the end of each
139 reaction is typical of the amount of acceptor necessary for the product synthesis. After
140 adjusting areas with the internal standard (tyrosine), transphosphorylation yields were
141 determined as follows:

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$$\text{Transphosphorylation yield} = \frac{\text{Area of initial phenol} - \text{Area of remaining phenol}}{\text{Area of initial phenol}} \times 100$$

3. RESULTS AND DISCUSSION

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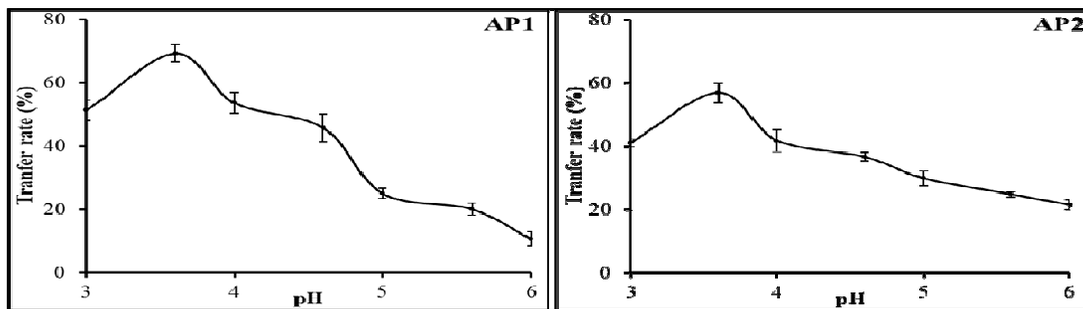
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The two non-specific acid phosphatases AP1 and AP2 purified from *A. communis* seeds [10] were tested in transphosphorylation reaction of pyrophosphate at pH 4. The retention time of the newly synthesized product was found to be similar to that of phenylphosphate (Data not shown). This shows clearly that the studied enzymes were able to catalyse the phosphorylation of pyrophosphate (PPi) to phenylphosphate. Pyrophosphate was earlier used as donor in many transphosphorylation reactions [3, 11] due to the fact that it is a safe and inexpensive compound which can be used in large excess for various manufacturing processes [12]. Also, this molecule is easily synthesized from phosphate groups [13]. Therefore, an efficient phosphorylation process could be achieved by recycling pyrophosphate to phosphate group by-products for further transfer in transphosphorylation reactions.

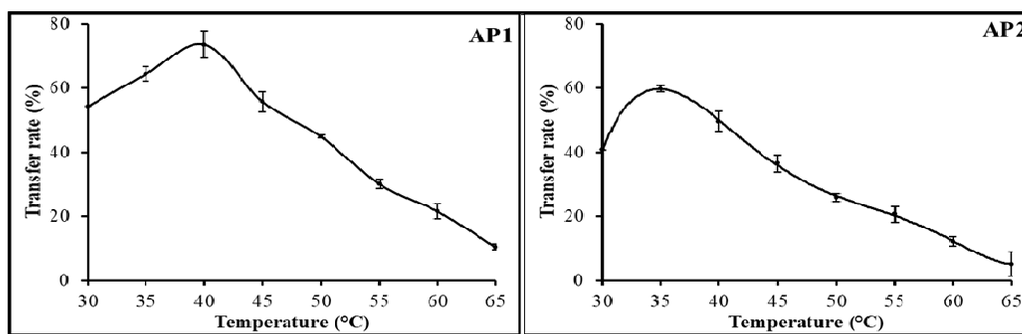
In order to optimize the phenylphosphate formation, five parameters were investigated namely the pH, time, donor concentration, acceptor concentration and enzyme amount dependency. In **Figure 1** the pH dependency of the phosphotransferase reaction is depicted. The highest yield of phenylphosphate (69.25 and 56.75% respectively for AP1 and AP2) was obtained at pH 3.6. It is noteworthy that there are few reports on phosphotransferase activity of plant acid phosphatases, but it seems that these biocatalysts exhibit better transfer activity at low pH compared to their hydrolysis activity (optima pH sited between 5 and 6). A similar behaviour has been observed by KOFFI et al. [3] for four non-specific acid phosphatases isolated from the cucurbit *L. siceraria* seeds. According to Huber et al. [14], the dissimilarity between the optima pH for phosphorylation and dephosphorylation could be due to ionized groups sited in the active site of the enzymes which would favour synthesis and disadvantage hydrolysis in more acidic conditions.

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184 **Figure 1.** Effect of pH on transphosphorylation reaction catalyzed by the acid phosphatases
185 (AP1 and AP2) from breadfruit (*A. communis*) seeds.
186 Bars represent \pm SE
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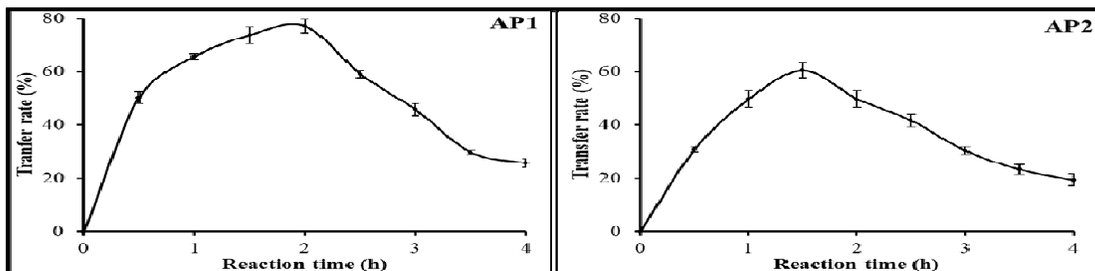
188 The enzyme activities were also measured at various temperatures from 30 to 65°C. AP1
189 had maximum activity at 40°C whereas AP2 shows maximum activity at 35°C. In these
190 conditions, optima yields of 73.58 and 59.75% were obtained for phosphatases AP1 and
191 AP2, respectively (**Figure 2**). The optimal temperatures of transphosphorylation were lower
192 than that obtained for dephosphorylation (55°C). So, these temperatures would be
193 advantageous since the studied enzymes have been earlier shown to be stable at
194 temperatures below 55°C [10]. Asano et al. [5] obtained their best transphosphorylation rate
195 at 40°C during the synthesis of Inosine-5'-monophosphate using a phosphorylating enzyme
196 isolated from bacteria *M. morganii*. The general observation would be that both the
197 phosphorylation and dephosphorylation temperatures do not coincide. It appears that the
198 optimum phosphorylation temperature is generally lower (ranging from 30 to 40°C)
199 compared to that of dephosphorylation (above 45°C).
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202 **Figure 2.** Effect of temperature on transphosphorylation reaction catalyzed by the acid
203 phosphatases (AP1 and AP2) from breadfruit (*A. communis*) seeds
204 Bars represent \pm SE
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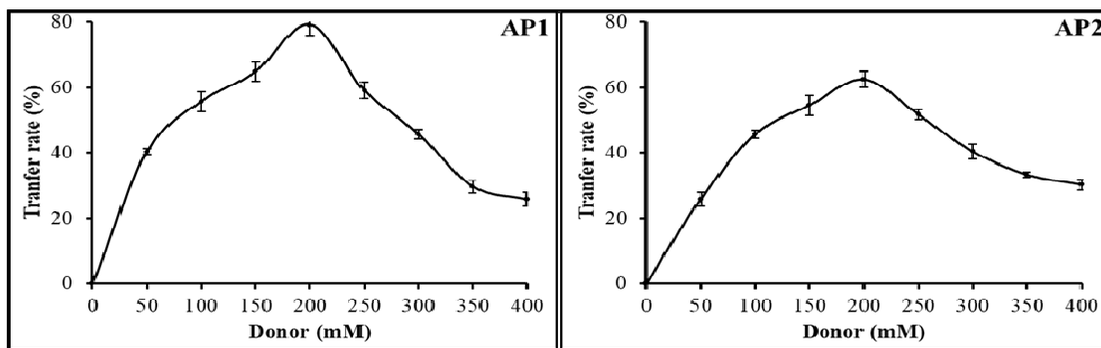
207 The time course of phenylphosphate synthesis by acid phosphatases AP1 and AP2 isolated
208 from bread fruit *A. communis* seeds is depicted in **Figure 3**. The two enzymes show
209 maximum transphosphorylation yield at 1.25 h. This time course of reaction is very low
210 compared with those obtained for 5'-inosinic acid synthesis as reported by Mihara et al. [2].

211 These authors have obtained their better phosphotransferase activity at time courses
 212 ranging from 18 to 36 h. Ishikawa et al. [12] have reported phosphorylation times sited
 213 between 5 and 15 h for inosine-5'-monophosphate (5'-IMP) synthesis. Therefore, the very
 214 short period of synthesis reactions catalysed by the studied acid phosphatases may
 215 constitute an interesting feature insofar as this would make it possible to save energy in
 216 biotransformation processes using these biocatalysts. Also, it should be remembered that
 217 this reaction time is very well included in the stability time (fully active for more than 150 min)
 218 of these enzymes as describe by Konan et al. [10].
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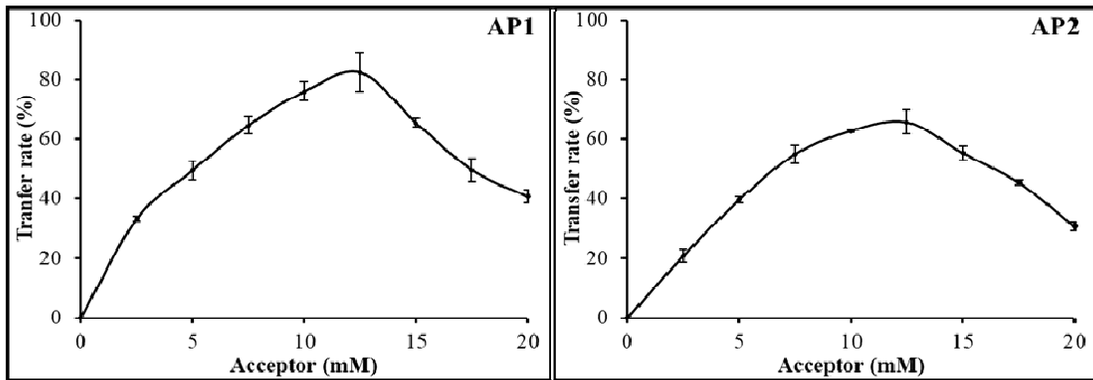


220 **Figure 3.** Time course of phenylphosphate synthesis by the acid phosphatases (AP1 and
 221 AP2) from breadfruit (*A. communis*) seeds.
 222 Bars represent \pm SE
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224 The yields of phenylphosphate produced by nonspecific acid phosphatases AP1 and AP2
 225 were highly dependent on donor and acceptor concentration (**Figures 4 and 5**). Similar
 226 behaviours were observed by Van Herk et al. [9]. In our study, the maximum concentrations
 227 of phosphorylated product were obtained from 200 mM pyrophosphate and 12.5 mM phenol
 228 for both AP1 and AP2. In these conditions, optimum yields of 82.39 and 65.71% were
 229 obtained for AP1 and AP2, respectively. As presented in figures 4 and 5, it seems that
 230 higher concentrations of donor and acceptor than those obtained in this study would inhibit
 231 the phosphoryltransferase activities. This would explain the decrease in the transfer rate
 232 since it is well known that high substrate loads are required to achieve high product titers,
 233 while kinetic control is crucial to keep the phosphorylation/hydrolysis ratio as high as
 234 possible due to the competition between hydrolysis and transphosphorylation in aqueous
 235 medium [15]. A common mechanistic feature of phosphate-transferring phosphatases is the
 236 formation of a covalent phospho-enzyme intermediate, which undergoes either nucleophilic
 237 attack by water (hydrolysis) or accepts a suitable nucleophile (transphosphorylation).
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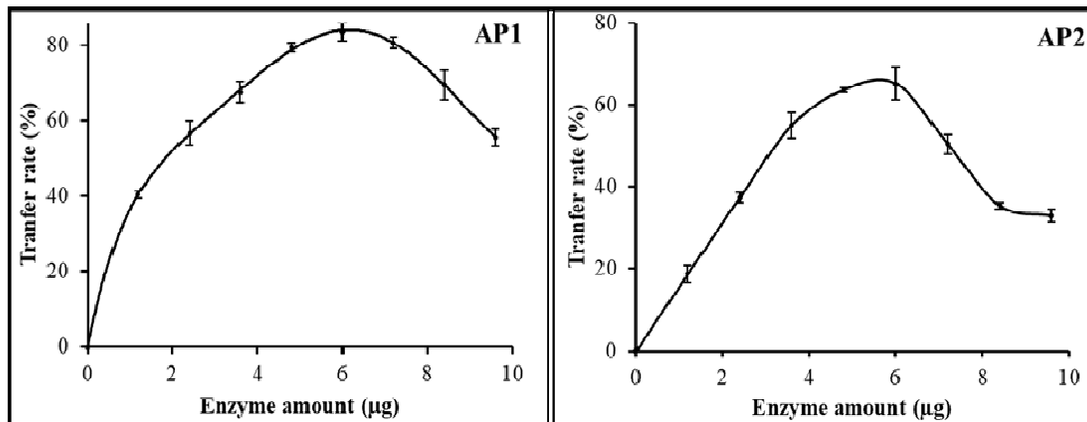


239 **Figure 4.** Effect of phosphoryl donor (sodium pyrophosphate) on transphosphorylation
 240 reaction catalyzed by the acid phosphatases (AP1 and AP2) from breadfruit (*A. communis*)
 241 seeds. Bars represent \pm SE



243 **Figure 5.** Effect of phosphoryl acceptor (phenol) on transphosphorylation reaction catalyzed
 244 by the acid phosphatases (AP1 and AP2) from breadfruit (*A. communis*) seeds.
 245 Bars represent \pm SE.

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 247 **Figure 6** shows effect of enzyme amounts on transphosphorylation reactions. This result
 248 indicates that high amounts of enzymes would be unfavourable to the phosphoryl transfer
 249 reaction. It appears that these high enzyme amounts result in the release of high amounts of
 250 inorganic phosphate, which could inhibit the phosphatase [16]. The phosphotransferase
 251 activities were optimal with enzyme amounts of 6 μ g for both AP1 and AP2, and average
 252 yields of 83.60 and 65.10% were respectively obtained.
 253



254 **Figure 6.** Effect of enzymatic unit on transphosphorylation reaction catalyzed by the acid
 255 phosphatases (AP1 and AP2) from breadfruit (*A. communis*) seeds. Bars represent \pm SE.

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258 **Table 1** summarized the average yields of transfer at the different stages as well as those
 259 obtained under the optimal conditions. During this study, yields significantly improved (about
 260 10 to 15%). In optimal conditions yields of 84.20 and 66.78% were obtained for AP1 and
 261 AP2, respectively. These values are higher than those reported for nonspecific acid
 262 phosphatases from seeds of the orphan crop *L. siceraria* [3] and for the nonspecific acid
 263 phosphatase from *Salmonella typhimurium* LT2 [17]. Although they are plant acid
 264 phosphatases, the studied enzymes exhibit good transphosphorylation potentials compared
 265 to bacterial phosphoryl transfer enzymes which were often modified to improve their
 266 transferase properties. Regarding these interesting phosphoryltransferase activities, it is
 267 likely that nonspecific acid phosphatases from the breadfruit *A. communis* seeds would be

268 useful to phosphorylate a wide range of nucleophile compounds such as nucleotides often
269 used as food additives and pharmaceutical intermediates.

270

271 **Table 1.** Transfer product yields in optimal conditions of the transphosphorylation assay
272 catalyzed by acid phosphatases AP1 and AP2 from breadfruit (*A. communis*) seeds.

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Parameters	Transphosphorylation yield	
	AP1	AP2
pH	69.25 ± 0.8	56.75 ± 1.1
Temperature	73.58 ± 0.7	59.75 ± 2.2
Reaction time	76.98 ± 1.2	60.50 ± 0.9
Donor	79.08 ± 1.5	62.47 ± 0.9
Acceptor	82.39 ± 0.5	65.71 ± 0.7
Enzyme unit	83.60 ± 0.9	65.10 ± 1.0
Optimum conditions	84.20 ± 0.6	66.78 ± 0.5

274 Values given are the average from at least three experiments.

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277 4. CONCLUSION

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279 The data obtained in this study showed that acid phosphatases from seeds of the breadfruit
280 *A. communis* may be good alternative biocatalysts to form phosphoconjugates by applying
281 them in synthesis of various phosphorylated compounds such as nucleotides often used as
282 food additives and pharmaceutical intermediates. Shorter time course reactions (1.25 h at
283 the most) were observed for optimum synthesis, suggesting energy saving during
284 biotransformation processes. The acid phosphatase AP1 would be the most promising on
285 the basis the better synthesis product yield (84.20%). The two biocatalysts could be
286 considered as new valuable tools for bioprocesses.

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