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1	Are enzymes transported in soil by water fluxes?
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14	Highlights
15	Hydrolytic enzymes are transported convectively, attached to soil colloids
16	• This transport is soil and enzyme specific
17	• Soil colloids released jointly with ions which alter ionic strength in effluents
18	• Enzyme activity demonstrated a bell-shape relationship with the ionic strength
19	
20	ABSTRACT
21	Transport of extracellular hydrolytic enzymes in soils has always been a subject of doubt. The
22	considerations against its importance are that (i) enzymes benefit their producers the most when
23	they remain in close proximity; and (ii) enzymes are large molecules with low mobility due to
24	high affinity to fine soil particles and organic matter. However, soil mineral colloids (SMC), to
25	which extracellular enzymes also have an affinity and which are known to facilitate transport of
26	a broad variety of chemicals and microorganisms in soils, can serve as vehicles for enzyme
27	transport as well. Since current literature lacks information on enzyme transport in soils, our goal
28	was to determine whether enzymes are transported and, if so, whether they are transported in a
29	free- or in a colloid-associated form. We conducted column transport experiments with four
30	hydrolytic enzymes, namely, β -glucosidase, acid-phosphatase, cellobiohydrolase, and
31	xylosidase, in soils with contrasting textures. The eluents containing enzymes were applied on

top of soil columns, while enzyme activities, SMC, and electrical conductivity were measured in the effluents from the columns. Our results provided evidence of joint enzyme transport with soil colloids. The enzymes associated with the coarse SMC (1 μ m < Ø) contributed 52 - 88% of the total enzyme activity in the effluents. The remaining enzyme activity was attributed to the enzymes associated with organic colloids, fine SMC (Ø < 1 μ m) and free enzymes in solution. This study suggested a dual effect of ionic strength in the soil suspension on enzyme activity and their release from soils with soil colloids.

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Keywords: Soil hydrolytic enzymes, colloid-facilitated transport, column experiments, ionicstrength.

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Abbreviations: IS, ionic strength; EC, electrical conductivity; POM, particulate organic matter;
SMC, soil mineral colloids; CMC, coarse mineral colloids; FMC, fine mineral colloids; TN, total
nitrogen; TC, total carbon; SL, Sandy loam; SL-S, sandy loam soil from summit; SL-D, sandy
loam soil from depression.

47

48 INTRODUCTION

49 Most plant and microbial cell debris present within soil as polymeric molecules are quickly transformed by extracellular enzymes to oligo- and monomers, which then become readily 50 51 available to microbial decomposers. Like most proteins, soil extracellular enzymes are capable to 52 diffuse away from their parent cell in free solutions due to Brownian motion (Burns et al., 2013). 53 Moreover, several studies in artificial solutions have reported a self-propelled diffusion of enzymes, which enhanced their movement by 30-80% during substrate catalysis (Yu at al., 2009; 54 Muddanaet al., 2010; Riedel et al., 2015; Jee et al., 2018; Günther et al., 2018). Still, there is no 55 56 consensus on the enzyme diffusivity. On one hand, diffusivity of free enzymes should increase as 57 substrate availability decreases, thus the enzyme producer can potentially access more distant substrates (Allison et al., 2011). On the other hand, competition for products between enzyme 58 producers suggests relatively low enzyme diffusivity (Burns et al., 2013). The amount of reaction 59 products captured by a microbial cell per unit of enzyme produced declines with increasing 60 distance between the cell and the produced enzymes due to diffusion losses of the product to the 61 62 environment, cell competition for reaction products, and decreasing enzyme and substrate

concentrations. These product losses increase exponentially with the distance between the enzymes and microbial cells, and less than 4% of the product can reach the microbial cells located at a distance > 200 μ m from the enzymes (Guber et al, 2021). Therefore, low enzyme diffusion, and thus lower diffusion losses of the product might represent a beneficial strategy for microbial cells.

The enzyme's capacity to move within an unrestricting volume of liquid does not imply that 68 enzymes can similarly freely diffuse through soil pores. Among factors restricting such diffusion 69 70 are: (i) enzyme attachment to soil particles and soil colloids (Nannipieri et al., 2003); (ii) 71 relatively low water retention and fast draining of large pores resulting in reduction of pore volume available for enzyme movement (Allison et al., 2011); (iii) hydraulic discontinuity of 72 water pathways in partly saturated pores and layers of extracellular polymeric substances, 73 created by microorganisms on surface of soil particles, restricting free enzyme motion (Or et al., 74 75 2007). Moreover, most free enzymes survive only briefly in pore solutions due to fast denaturation by physical and chemical factors or consumption by proteolytic microorganisms 76 (Sarkar and Burns, 1984; Burns, 1986). However, adsorption confers protection against 77 microbial degradation (Lähdesmäki & Piispanen, 1992; Kedi et al., 2013). Therefore, most 78 79 viable enzymes and microorganisms in soil are bound to organic surfaces (Ahmed and Oades, 1984; Christensen and Bech-Andersen, 1989; Jocteur Monrozier et al., 1991) and fine soil 80 particles (Singh and Singh, 1995; Schulten et al., 1993; Stemmer et al., 1998; Kandeler et al., 81 1999a). Association of microbial cells and enzymes with soil particles and organic surfaces 82 precludes their free diffusive transport due to relatively large size of these particles and low pore 83 volume available for the diffusion in partly water-saturated soil. However, rapid water 84 85 infiltration after heavy rainfalls or overland irrigation causes both physical and chemical perturbations in the soil and results in a release of colloids from soil matrix and their transport 86 via large pores (Ryan and Elimelech, 1996). These colloids are composed of organic and mineral 87 particles (including clay and partly silt soil fractions) with effective diameters $< 10 \,\mu m$ 88 (McCarthy and Zachara, 1989; Sposito, 2016). The colloids within size range 0.01-10 µm are 89 90 regarded as the most stable (Buffle and Leppard, 1995) and are common carriers of soil microorganisms, organic substances (Smith et al., 1985; McCarthy & Zachara, 1989; Natsch et 91 al., 1996), and environmental contaminants (Ryan & Elimelech, 1996; de Jonge et al., 2004). 92 Strong adsorption of enzymes on fine soil particles and their low extractability in free form 93

94 (Fornasier et al., 2011), suggests possibility of their transport by moving colloids. However,

95 despite extensive study of enzyme interaction with clay minerals and soil colloids during last 4

96 decades, surprisingly, enzyme transport in soils has never been reported. Therefore, the goal of

97 this study was to explore the possibility of enzyme transport in intact soils with contrasting

98 textures under water flow conditions that mimic those during heavy rainfalls.

99 The activity of four hydrolytic enzymes involved in C and P acquisition were studied in the 100 transport experiment: β-glucosidase, acid-phosphatase, cellobiohydrolase, and xylosidase. The 101 former two participate in the last step of decomposition, i.e., release of monomers (glucose and 102 phosphate) that are easily available for microorganisms. The latter two are involved in the early 103 stage of decomposition destroying long polymeric chains of cellulose and hemicelluloses.

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2. MATERIALS AND METHODS

106 *2.1. Soil properties*

107 Soil for the column experiments was collected at three experimental sites with contrasting soil texture located in Michigan, USA. Sandy soil of Riddles-Hillsdale series (fine-loamy, mixed, 108 109 active, mesic Typic Hapludalfs) was obtained from Michigan State University's (MSU) Sandhill farm site, East Lansing, MI. We further refer to this soil as Sand. Two sandy loam soils of Capac 110 series (fine-loamy, mixed active, mesic Aquic Glossudalf) were obtained from summit and 111 depression topographical positions at MSU's Mason experimental farm, East Lansing, MI. We 112 refer to them as sandy loam soils SL-S and SL-D, for summit and depression respectively. 113 Loamy soil of Kalamazoo series (fine-loamy, mixed, active, mesic Typic Hapludalfs) was from 114 Cellulosic Biofuel Diversity Experiment at Kellogg Biological Station Long-Term Ecological 115 Research site, Hickory Corners, MI, referred to further as Loam soil. At each site, undisturbed 116 soil cores (Ø=2.5 cm, height=10 cm) were taken from 5-15 cm depth and stored at 4°C for 2-3 117 days prior to experiments. In addition, disturbed soil samples in amounts of approximately 300 g 118 were taken from the immediate vicinity of the intact cores for basic soil analyses. At the time of 119 120 sampling, Sand soil was under corn, SL-S and SL-D soils were under corn-soybean rotation, and Loam soil was under long-term (>10 years) native prairie vegetation. We collected 3 cores from 121 each site and one additional core from Loam soil site, for a total of 13 cores. The number of soil 122 cores was selected arbitrarily solely for exploring purposes. 123

The following soil analyses were conducted using disturbed soil samples. Soil texture and 124 soil mineral colloids (SMC), operationally defined as mineral particles $\emptyset < 10 \,\mu\text{m}$, were 125 measured using the pipette method (Gee & Or, 2002). Total nitrogen (TN) and total carbon (TC) 126 127 were measured using an elemental analyzer ECS 4010 CHNSO (Costech Analytical Technologies Inc., Valencia, CA, USA). Soil particulate organic matter (POM) was measured by 128 129 wet sieving (Cambardella and Elliot, 1992). Soil pH and electrical conductivity (EC) were measured using SevenCompact Duo s213 meter (Mettler-Toledo LLC, Columbus, OH USA) in 130 131 Soil-DI water suspensions at 1:1 solid/liquid ratio. General characteristics of the studied soils are 132 given in Table 1.

133 *2.2 Applied suspensions*

The compositions of applied suspensions, referred to as eluents, for the enzyme transport 134 experiment were designed to minimize artificial effects of solution chemistry on the transport 135 136 and transformation processes within the soil cores. This composition mimicked soil suspensions 137 generated in the field during heavy rainfalls by kinetic energy of rain drops or runoff water. Therefore, the eluents were prepared individually for each soil by adding 1 g of fresh soil to 100 138 ml of DI water followed by 5 min low-energy sonication using Fisher Scientific FS20 Ultrasonic 139 Cleaner (Thermo Fisher Scientific Inc., Waltham, MA, USA). The sonication settings were 140 chosen to break up soil aggregates, while preserving SMC (Stemmer et al., 1998), soil organic 141 colloids, microorganisms and enzymes (Kandeler et al., 1999b). Soil suspensions were kept for 142 30 min to allow settling of sand particles, and the supernatant solutions separated from the 143 sediment were used as eluents. Thus, the prepared eluents contained dissolved chemicals, soil 144 145 particles, microorganisms, and enzymes native to the respective soils. The activities of β glucosidase, phosphatase, xylosidase and cellobiohydrolase were measured in the eluents before 146 and after precipitating the coarse mineral colloids (CMC) as described in Sections 2.3 and 2.4. 147 Soil mineral colloids were measured in the eluents as described in Section 2.2. 148

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2.3 Column experiment

Soil column experiments were conducted to quantify possible activity of the four hydrolytic enzymes in the suspensions passing through the intact soil cores, which we will refer to as effluents. We used undisturbed soil cores to preserve distribution of enzymes and water flow pathways unchanged in the soils. For Sand soil the entire soil cores were used as experimental

columns, that is, the column height was 10 cm. For finer textured SL-S, SL-D, and Loam soils, 154 due to their low water infiltration rates, the intact cores were cut to 5 cm experimental columns. 155 Longer Sand columns as compared with those for SL-S, SL-D, and Loam soils reduced the water 156 flow velocity in the Sand columns and prevented mechanical detachment of colloids from soil. 157 The eluents were applied by a pipette to the top of the columns in 1 ml increments to prevent 158 water ponding on the soil surface (Fig. 1). A pressure head of -30 kPa, which is an equivalent to 159 the field capacity, was maintained at the bottom of the columns during the experiment to keep 160 161 steady-state flow through the columns. Coarse porous filters with particle retention $\emptyset > 40 \,\mu m$ (Filter Paper Grade 417, VWR[®], Radnor, PA, USA) were installed at the bottom of the columns. 162 The filters, permeable for SMC, microorganisms, and enzymes, prevented detachment of soil 163 particles from the columns. The effluents were collected in 4 ml increments from the bottom of 164 Loam, SL-S and SL-D columns, and in 8 ml increments from Sand columns. Collected effluents 165 166 were analyzed for activity of the four enzymes, SMC contents, pH, and EC. The enzyme activities were measured in the effluents with all colloids and in the supernatant solutions which 167 contained only the colloidal particles smaller than 1 µm (Fig. 1). To precipitate the coarse 168 mineral colloids (CMC) size of large 1 µm, the effluents were centrifuged for 5 min at 5000 rpm 169 using a Heraeus Megafuge 16 centrifuge (Thermo Fisher Scientific Inc., Waltham, MA, USA). 170 171 Diameter of precipitated CMC was calculated (Gee and Or, 2002) as:

$$d = \frac{1}{60} \sqrt{\frac{18\eta \ln{(\frac{r_2}{r_1})}}{(\rho_s - \rho_l)\omega^2 t}}$$
(1)

172

173 where r_1 to r_2 are distances from the axis of centrifuge rotor to the particle [cm], η is the fluid viscosity [g cm⁻¹ sec⁻¹], $\rho_s \sim 2.6$ g cm⁻³ is the density of soil mineral particles, ρ_l is the liquid 174 175 density [g cm⁻³], ω is the centrifuge angular velocity [rpm], t is time for particle of diameter d to settle from r_1 to r_2 [sec]. It was confirmed in a preliminary study that selected centrifugation 176 settings had a negligible effect on the activity of colloid-free almond β -glucosidase and wheat 177 germ acid phosphatase (CALZYME Laboratories, Inc., San Luis Obispo, CA U.S.A.) dissolved 178 179 in DI water. Precipitating mineral colloids smaller than 1 µm (e.g., 0.45 mm) and organic colloids ($\rho_s < 2.6 \text{ g cm}^{-3}$) requires increasing of angular velocity of centrifugation proportionally 180 to ω or increasing the duration of centrifugation as t^2 . Such changes in the centrifugation settings 181 182 results in partial precipitation of bio colloids and free enzymes in the effluent suspension in

amounts that are generally unknown. Therefore, most organic colloids, bio colloids enzymes and
 mineral colloids size of smaller 1 µm remained in supernatants, while CMC were precipitated by
 centrifugation.

186 For comparison purposes the results of the column experiments were expressed in a normalized form. Specifically, volumes of eluent that passed through the columns were 187 expressed in total pore volumes, where the total pore volume in each soil column was calculated 188 from its bulk density, particle density [~ 2.6 g cm⁻³], and column's volume. The SMC contents, 189 enzyme activities and EC values in the effluents from the columns were normalized by the 190 corresponding values in the eluents. To evaluate the effect of ionic strength in the effluents on 191 the enzyme activity, the measured EC values were transformed using the Marion-Babcock 192 equation (Sposito, 2016): 193

194

$$log_{10}(I) = -1.841 + 1.009 \ log_{10} (EC)$$
 for $I \le 0.2 \text{ M}$ (2)

where *I* is the ionic strength in the solution [M], and EC is the electrical conductivity in the solution [dS m^{-1}].

197 *2.4. Enzyme assay*

The activity of β -glucosidase, acid-phosphatase, xylosidase and cellobiohydrolase were 198 measured in the soil, eluents, and effluents collected in the column experiments before and after 199 precipitating CMC as described in Section 2.3 (Fig. 1). For the activity measurements we used a 200 microplate fluorometric assay technique described in Saiya-Cork et al. (2002) and Deng et al. 201 202 (2011) with minor modifications of the substrate concentrations. Specifically, we used substrates based on 4-methylumbelliferone (MUF) fluorescent reagent (i.e. MUF-β-D-glucopyranoside, 203 MUF-Phosphate, MUF-b-D-xylopyranoside, and MUF-β-D-cellobioside) in quantities of 40 204 nmol well⁻¹ per 50 µl well⁻¹ soil suspensions with 5.0 µmol well⁻¹ of sodium MES buffer 205 (C₆H₁₃NO₄SNa_{0.5}, pH 6.1 at 25°C) solution. The calibration was performed using 0, 100, 200 206 and 400 pmol well⁻¹ MUF solutions with the same aliquots of soil suspensions. The fluorescence 207 intensity in the plates was measured using a Multilabel Plate Reader Victor³ (PerkinElmer Inc., 208 Waltham, MA, USA) every 15 min for 2 hours. The enzyme activities were calculated from 209 linear parts of the intensity time series with correction for the substrate autohydrolysis (Deng et 210 al., 2011). 211

The enzyme activity associated with CMC in the effluents v_c^{ef} was calculated as:

$$v_c^{ef} = v_t^{ef} - v_s^{ef} \tag{3}$$

214 were v_t^{ef} is the total enzyme activity in the effluent [pmol min⁻¹ ml⁻¹], and v_s^{ef} is the enzyme 215 activity in the supernatant from centrifuged effluent [pmol min⁻¹ ml⁻¹].

Since the mass of SMC differed in the effluents from the four soils and changed over time, to evaluate the effect of ionic strength on the enzyme activity associated with SMC, the activity was normalized by the dry mass of colloids in the effluent:

 $v_c^S = v_c^{ef} / S_c$

where v_c^S is the activity of enzymes associated with CMC per mass of dry colloids [nmol min⁻¹ g dry colloids⁻¹], and S_c is the dry mass of CMC in the effluent [g L⁻¹]. Note, that the dry mass of colloids in supernatants from the effluents was within the accuracy range of analytical balance (0.1 mg). Therefore, the dry mass of SMC measured in the effluents was attributed to CMC.

224 2.5. Data processing and statistical analysis.

Differences among the soils in terms of the studied properties were conducted via analysis of 225 variance, with soils as the only studied factor, using *lm* function of R (version 4.1.2). To compare 226 the enzyme activities within the soils we fitted the data with a statistical model that consisted of 227 (i) the fixed effects of the soils and the enzymes and their interaction, and (ii) a random effect of 228 the replicated samples, nested within the soils, also used as an error term for testing the soils' 229 effect. Model fitting was conducted using *lmer* function from R's *lme4* package. When the 230 effects of the studied factors were found to be statistically significant (p < 0.05) we performed t-231 tests to conduct all pairwise comparisons among the means using *emmeans* package. 232

We used correlation analysis to explore possible relationships between enzyme activity associated with CMC and colloid contents in the effluents from the soil columns using Analysis tool of SigmaPlot software (Systat Software Inc., San Jose, California, USA). Pearson correlation coefficients were reported as statistically significant and p < 0.05 and marginally significant at p < 0.1.

238 3. RESULTS

239 *3.1. Properties of the soils and applied suspensions.*

The soils of the four locations differed in their properties. The bulk density was the lowest inthe sandy soil (Sand) due to its loose structure. The highest sand content was expectedly

(4)

observed in Sand, lowest in Loam, and intermediate in the sandy loam SL-S and SL-D soils

243 (Table 1). An inverse trend was observed for SMCs, which contents were the highest in Loam,

lowest in Sand, and intermediate, though close to Loam, in SL-S and SL-D soils (Table 1). The

POM content was rather low in all soils (0.11 % - 0.31%) but tended to be the highest in Sand.

246 TN and TC increased in the soils in the order of decreasing sand content. The pH values ranged

from 5.4 to 6.0. The EC values were 2 to 3 times higher in SL-S and SL-D as compared to Sand

and Loam, indicating overall higher contents of dissolvable ions in sandy loam soils.

The SMC content in the eluents increased in the order Sand \leq SL-D \leq SL-S \leq Loam. The pH values were slightly higher than in the soil suspensions and ranged from 5.9 to 6.7, while the EC values were much lower (6-9 μ S cm⁻¹) in the eluents as compared to those in the soils, as an expected result of a 100-fold dilution of the soil suspensions (Table 1).

The differences in the soil properties were mirrored by enzyme activities, which for β -253 glucosidase and phosphatase were the highest in Loam and SL-S, and the lowest in Sand (Table 254 1). For xylosidase and cellobiohydrolase the highest activities were observed in SL-S, and the 255 lowest in Sand. The enzyme activity in the eluents followed the same general trend as that in the 256 257 soils, with much smaller differences between β -glucosidase and phosphatase activities as compared with those in the soils. Among the four enzymes, the activities in most soil samples 258 and eluents increased in the order xylosidase < cellobiohydrolase < phosphatase < β -glucosidase 259 260 and were up to one order of magnitude higher for phosphatase and β -glucosidase than for xylosidase and cellobiohydrolase (Table 2). Precipitation of CMC by centrifugation considerably 261 262 reduced enzyme activity in the eluents. The enzyme activity associated with CMC calculated using Eq.(3) ranged from 33% to 100% in the eluents in the four soils. The lowest activities of 263 CMC-associated enzymes were observed in Sand (Table 2), while the highest activities were 264 observed in Loam, where the enzyme activities in the supernatants after eluent centrifugation 265 266 were below the detection limit.

267

3.2.Column experiment

The relative concentrations of SMC in the effluents were the highest in the first portions of effluent collected from the columns and decreased with the relative volumes of the solution passed through the columns (i.e. pore volumes) (Fig. 2 and Fig. S1). The recovered SMC differed in the four soils and ranged from 7% of that applied in Loam to 49% in Sand, with intermediate values for SL-S and SL-D (Table 2). The relative concentrations of SMC were the

highest for Sand and the lowest for Loam columns (Fig. 2ab). Noticeably, in the Sand columns

the relative concentrations of SMC from the first portions of the effluent were >1 (Fig. 2a),

indicating that the effluents contained more SMC than the applied eluents. For Loam (Fig. 2b),

277 SL-S (Fig. S1a), and SL-D (Fig. S1b) columns, the SMC contents in the effluents were always

smaller than in the eluents.

The EC dynamics in the effluent resembled that of SMC, though unlike SMC, the highest relative EC were observed in SL-S and SL-D columns (Fig. S1). The relative EC values in the effluents were >> 1 in all samples of all studied soils, indicating much higher concentrations of soluble chemicals in the effluents as compared to those in the applied eluents.

As was the case with SMC and EC, the relative activity of four enzymes in the effluents 283 284 decreased with the amount of eluent passing the columns (Fig. 3). Among the four enzymes, the absolute values of activities in the effluents were the lowest for xylosidase and cellobiohydrolase 285 286 and the highest for phosphatase (Table 3). Enzymatic activity in the first portions of the effluents 1.5 to 8 times exceeded those in the eluents (Fig. 3 and Fig. S2), with exception of β -glucosidase 287 in SL-S and SL-D columns (Fig. S2a) and cellobiohydrolase in the Loam column (Fig. 3b). 288 Among the four enzymes, the relative activity in the first portions of effluents was the highest for 289 phosphatase (Fig. 3b and Fig. S2b), the smallest for β -glucosidase (Fig. 3a and Fig S2a), and 290 intermediate for xylosidase (Fig. 3c and Fig. S2c) except the Loam soil, where xylosidase and 291 cellobiohydrolase were not detected in the effluents. Among the four soils, the relative activities 292 in the effluent were the highest for Sand, the lowest for Loam (Fig. 3), and intermediate for SL-S 293 and SL-D columns (Fig. S2), which was consistent with the dynamics of SMC outflow from the 294 columns (Fig. 3 and Fig. S1). For example, β -glucosidase activities in the effluent corresponding 295 to 1 pore volume in Sand was >2 times higher than that in Loam, while for phosphatase it was >4296 297 times higher than in Loam (Table 3).

For β -glucosidase and phosphatase the activity associated with CMC constituted a substantial portion of the total enzyme activity (Fig. 4, Table 3). For example, for β -glucosidase in the effluent corresponding to the first pore volume, the activity associated with CMC constituted 66% and 52% of the total activity for Loam and Sand, respectively (Table 3). For phosphatase, it was even higher and constituted 79% and 88% for Loam and Sand, respectively. In most tested enzymes, their activity associated with CMC was positively correlated with concentration of SMC in the effluents (Fig. 5 and Fig. S3). The correlation coefficients between v_c^{ef} and concentration of SMC in the effluents were high at p < 0.1 and p < 0.05 for all enzymes except xylosidase and cellobiohydrolase in the Loam column where the activities were below the detection level (Fig. 5), and except phosphatase, xylosidase and cellobiohydrolase in SL-D columns (Fig. S3).

The relationships between the activity of enzymes associated with SMC v_c^{ef} and ionic 309 strength in the effluents, as derived from EC using Eq.(2), were rather scattered. Still, a sharp 310 increase of β -glucosidase and phosphatase activity associated versus not associated with CMC 311 can be seen in the Sand and Loam columns with increasing ionic strength from 0.1 to 2.3 mM 312 (Fig. 6). The slope of linear regression between the enzyme activity and IS ranged from 4.5 to 43 313 pmol min⁻¹ g⁻¹ mM⁻¹ and was steeper for Sand as compared with that in the Loam columns. For 314 the SL-S and SL-D columns the activity of all enzymes was somewhat decreasing within the 315 316 ionic strength range 1.4 - 21.4 mM. This trend was less expressed but still visible for xylosidase (Fig. S4 a,c) and cellobiohydrolase (Fig. S4 b,d) in the SL-S and SL-D columns. 317

The pH values in the effluents moderately increased with pore volumes during the experiment from 6.0 to 6.6, 6.3 to 7.1 and 5.2 to 6.3 in the Sand, Loam, and both SL-S and SL-D columns, respectively. However, v_c^{ef} did not corelate with pH in the effluents.

321

322 4. DISCUSSION

The results of column experiments demonstrated possibility for soil hydrolytic enzymes to be transported by water fluxes during heavy rainfall events in the soils with different soil texture.

325

4.1. Associations between enzymes and colloids in the eluents

Precipitation of CMC in the eluents reduced considerably activity of the four enzymes 326 suggesting their association with colloids (Table 2). CMC are comprised of clay and silt 327 particles, which hydrolytic enzymes are mostly associated with (Feller at al., 1994; Turner et al., 328 2002; Sinsabaugh et al., 2008; Kandeler et al., 1999a; Kedi et al., 2013). Association of β -329 glucosidase with soil mineral colloids in paddy soil has been reported by Yan et al. (2010 a,b). 330 These authors found approximately 50% higher β -glucosidase association with fine (< 0.2 µm) 331 than with the coarse $(0.2 - 2.0 \,\mu\text{m})$ SMC. On the contrary, 86% to 100% of β -glucosidase 332 activity was associated with CMC in our study. The discrepancy likely results from Yan et al 333

(2010 a,b) using highly diluted (1:100 solid to liquid ratio) soil suspensions, where all added
enzymes freely interacted with all resuspended soil particles. Yet, in intact soils, as used in our
study, enzyme-colloid interaction occurs primarily in hydrologically active pores occasionally
and not completely saturated by liquids.

It should be noted that enzyme-CMC associations in eluents were enzyme and soil specific. 338 While activity of all enzymes was completely associated with CMS in Loam soil, in the coarser 339 textured soils the percent of CMC associated activity was lower. Of the four studied enzymes, β -340 341 glucosidase seemed to be the most associated with CMC, while activity of xylosidase and cellobiohydrolase tended to be the least associated, especially in Sand (Table 2). While it is not 342 clear what is the cause for the differences among the enzymes, the results suggest high 343 possibility of joint transport of enzymes with CMC, especially in fine-textured soils, though 344 transport of not associated enzymes cannot be excluded. 345

346 4.2. Colloid and enzyme transport through the soil columns

The transport experiments demonstrated high mobility and recovery of SMC in sandy and sandy loam soils and low mobility in the Loam (Fig. 2), likely caused by straining of colloids within small pores of the finer textured soil. This result is consistent with other studies (Bradford et al., 2002, 2003), where straining of colloids was shown to increase with increasing colloid size and with decreasing soil grain size (Bradford et al., 2003).

352 Soil colloids appeared to be involved in enzyme transport, the result that was consistently observed in all cases when the enzymes were present in the soil in appreciable amounts (Table 353 3). High significant correlation between the activity of these enzymes associated with CMC 354 (v_c^{ef}) and SMC contents in the effluent (Fig. 5, Fig. S3) imply that v_c^{ef} was proportional to the 355 mass of transported colloids for each portion of the solution passed the columns. Closeness of the 356 intercepts of the linear regressions to zero for all enzymes in Sand and Loam columns (Fig. 5), 357 358 and considerable (from 2 to10-fold) decreases in the enzyme activity after precipitation of CMC in the effluents (Table 3 and Fig. 5) suggested that large fraction of the enzymes was transported 359 360 convectively attached to SMC.

High correlation coefficients between enzyme activity and contents of SMC in the
effluents do not imply that all detected enzyme activity was associated solely with SMC. Nonzero activities of all enzymes were detected in the supernatants after centrifugation of effluents

from SL-D columns (Fig. S3). The centrifugation removed mostly CMC ($\emptyset > 1 \mu m$) from the 364 365 effluents. Fine mineral colloids, organic colloids, microbial cells and macromolecules, with density much smaller than that of soil mineral particles ($\rho_s < 2.6 \text{ g cm}^{-3}$), were only partly 366 precipitated by centrifugation, but could also act as colloidal carriers (Buffle and Leppard, 1995). 367 368 The number of fine colloid particles $< 0.2 \mu m$ (not necessarily mass) can constitute up to 70% of 369 the total number of mobile colloids during a heavy rainfall (Lehmann et al., 2021). Moreover, the 370 activity of β -glucosidase associated with fine (<0.2 µm) SMC can be higher than that associated 371 with coarse $(0.2 - 2.0 \,\mu\text{m})$ SMC (Yan et al., 2010 a,b). Therefore, enzymes remaining in the 372 effluents after centrifugation were more likely associated with not-precipitated FMC, viable microbial cells (Nannipieri et al., 2012), and/or organic colloids rather than being free floating 373 374 enzymes, whose lifetime in soil solutions is rather short (Burns, 1982; Ladd, 1985; Nannipieri, 1994, Nannipieri et al., 2002). Possible enzyme association with soil colloids suggest that in real-375 world systems enzyme transport depends on the source and composition of moving colloids. 376 Specifically for agricultural environments, colloid-facilitated transport of enzymes can be 377 expected after irrigation by reclaimed wastewater or heavy rainfalls followed manure 378 379 application.

380 Colloidal transport is known to affect soil hydraulic properties, e.g., conductivity of soil pores and their sizes (Miller and Baharuddin, 1986; McDowell-Boyer et al., 1986). Precipitation 381 and straining of colloids in soil pores during colloidal transport results in progressive straining of 382 colloids and in the associated decrease of pore volume available for transport. Therefore, it is 383 384 reasonable to suggest that, with time, the ratio between large and small colloids in the effluents 385 shifts toward small colloids. Since the proportion of organics and the surface area of colloids increases with decreasing sizes of colloidal material (Buffle and Leppard, 1995), higher 386 association of enzymes with fine colloids is expected. This fact can be illustrated by β -387 glucosidase (Fig. 4a) and xylosidase (Fig. 4c) activities in SL-S columns, where the ratio 388 between activities of enzymes associated (filled bars) and not associated (open bars) with CMC 389 390 decreased with the number of pore volumes.

The relationship between SMC content in effluents and activity of CMC-associated enzymes was enzyme specific (Fig. 5 and Fig. S3) possibly due to mineralogical composition of transported colloids and differences in enzyme-mineral interactions. The mineral composition of Ap horizon in fine, mixed, mesic, Typic Hapludalfs, used in our study, varies greatly among soil

fractions. According to Sparks et al. (1979), quartz and mica dominate (>50%) in > 5 μ m and < 2 395 μm mineral fractions of Typic Hapludalfs. The 2-5 μm soil fraction is composed of quartz, mica 396 397 and kaolinite in approximately equal amounts. Kaolinite contents vary from 7 to 28% in the soil fractions with approximately two-fold greater contents in $< 5 \mu m$ soil fractions. 398 Feldspar is present only in silt fractions, while vermiculite only in clay fractions, in the amounts 399 of less than 10%. Different enzyme affinity to the substrates in presence of soil minerals has 400 been reported in multiple studies (Ross, 1983; Makboul and Ottow, 1979; Haska, 1975; Pflug 401 (1982); Gianfreda and Bollag, 1994). Therefore, it is reasonable to suggest that SMC-enzyme 402 403 associations and enzyme activities in the effluents were affected by changing sizes and mineralogy of transporting colloids. For example, the share of quartz particles in the total SMC 404 405 mass passing the columns likely decreased with time, while the share of mica and kaolinite 406 particles increased. Due to differences in soil texture, it is expected that the ratios between contents of different minerals in the effluents was soil specific. Therefore, the differences in the 407 enzyme activities observed in Fig. 5 and Fig. S3 can likely be attributed to different association 408 409 of enzymes and their activity on SMC, which mineralogy changed in the effluents with respect to 410 the textural, mineralogical, and hydraulic properties of these soils. We realize that it is infeasible to conduct particle and mineralogical analysis in the effluents, but it is worth to note that the 411 412 relationship between the enzyme activity and quantities of transported colloids is more complicated than any adsorption isotherm used to model interactions between chemicals, 413 microorganisms, and soil particles. 414

415

4.3. Whether ionic strength in solution affect enzyme transport?

Soil's ionic strength affected recovery of SMC applied with eluent in the fine-textured soils 416 (Loam, SL-S and SL-D). The recovery was greater in the SL-S and SL-D columns (Table 2), 417 where soil EC and, respectively, IS were much higher than those in the Loam soils (Table 1). 418 Increasing IS causes coagulation, while decreasing IS causes disaggregation and mobilization of 419 420 soil colloids (Ryan and Elimelech, 1996). Rainfall water has typically much lower IS as compared with that in soil solutions (McCarthy and Zachara, 1989). Therefore, decrease of soil 421 422 IS, due to dilution of pore solution by rainwater, results in a release of colloids from the soil matrix at early stages of rainwater infiltration into the soil (Flury et al., 2002; Grolimund and 423 Borkovec, 1999). With time the difference between IS in soil pores and the applied solution 424 decreases slowing down soil disaggregation. However, soil, disaggregated by low IS, becomes 425

less conductive and filters out new infiltrated colloids via straining or physical-chemical 426 427 collection by attractive surfaces of the immobile soil matrix (McCarthy and Zachara, 1989) 428 Visually, this process manifested itself in this study via a noticeable reduction of the infiltration rate in the fine textured soils after 1-1.5 pore volumes of the effluent were collected. The high 429 recovery of colloids from the Sand columns (49.1%) was less affected by soil disaggregation due 430 to much lower clay to sand ration as compared with the other soils (Table 1). Therefore, the 431 effect of IS on colloidal outflow was less pronounced in the Sandy, and more pronounced in 432 Loam and Sandy loam columns. 433

The IS also affects the activity of transported enzymes. In batch experiments with pure clay 434 minerals and in solutions with controlled pH and EC the relationships between enzyme activities 435 and IS could be well described quantitatively (e.g., Quiquampoix et al., 1993; Leprince & 436 Quiquampoix, 1996). In real soil and transport conditions this relationship is rather scattered 437 (Fig. 6) and likely altered by: (i) possible differences in mineralogy of soil colloids of the four 438 soils and effluents from the columns (Jaber et al., 2018; Nannipieri et al., 1996); (ii) different 439 affinity of enzymes to different minerals and colloids (Makboul and Ottow, 1979; Ross, 1983; 440 441 Sarkar et al., 1989); (iii) and different and likely changing during the transport experiment ionic and colloidal composition of the effluents. Yet, we attempt to derive a relationship between the 442 443 activity of adsorbed enzymes and IS for out column studies using dependencies between phosphatase catalytic activity and pH measured for montmorillonite at different IS levels 444 445 reported by Leprince & Quiquampoix (1996, Fig. 2, central column). The original data were reorganized to depict relationships between the rate of the catalytic reaction (V) and IS at pH 446 447 levels within 5.0 - 7.0 interval (Fig. 7, symbols). The reorganized relationships were fitted using an empirical equation: 448

449

$$V = \frac{a}{IS} exp[-b(\ln(IS) - c)^2]$$
(5)

where *a*, *b*, and *c* are the fitting parameters. The regression wizard of SigmaPlot software (Systat Software Inc., San Jose, California, USA) was used to fit Eq.(5) to the data shown in Fig. 7. The similarity in shapes between the regression curves and experimental data was assessed using the two-sample Kolmogorov-Smirnov test in the Real Statistics Resource Pack software (Release 7.6, Copyright (2013 – 2021), Charles Zaiontz, www.real-statistics.com).

455 Despite a small number of experimental points (6 on each curve), Eq.(5) adequately (p < 456 0.05) reproduced shapes of the V(IS) curves (Fig. 7). Therefore, Eq.(5) was applied to reconstruct

the shape of experimental curves describing relationships between activities of enzymes (i.e., β -457 458 glucosidase and phosphatase associated (Fig. 6 a,b) and not associated (Fig. 6 c,d) with CMC) 459 and IS in effluents from the four soils. Based on the Kolmogorov-Smirnov test the fitted curves reproduced adequately (p < 0.05) the shapes of β -glucosidase and phosphatase activity curves in 460 the Sand and Loam columns for enzymes not associated with CMC (Fig. 6 c,d). The shapes of 461 fitted curves differed significantly (p<0.05) from those experimental for enzymes associated with 462 463 the CMC. Fitting Eq.(5) to xylosidase and cellobiohydrolase activity was not successful due to scattered data on these curves (Fig. S4). 464

Several mechanisms can potentially explain a bell-shape relationship between enzyme 465 activity and IS. The first one is a competition between enzymes and cations in the effluents for 466 467 adsorption sites on mineral and organo-mineral colloids. Large IS values imply higher contents of anions and cations in the effluents, and stronger competition with enzymes for the adsorption 468 469 sites on SMC. The increase in IS also alters repulsive electrostatic interactions and weakens 470 enzyme association with colloids. The competition mechanism explains the decrease of enzyme activity but does not explain its increase within the IS range 0.2-1.3 mM. It also does not explain 471 the decrease in activity of the enzymes not associated with CMC (Fig. 6 c,d) which were likely 472 partly associated with unprecipitated colloids and partly were in a free form in the effluents. 473

474 The second mechanism, affecting the relationship between the enzyme activity and IS, is based on enzyme activity association with pH, which commonly has a bell-curve shape with the 475 maximum activity at an optimal pH level (Leprince and Quiquampoix, 1996; Turner, 2010; Kedi 476 et al., 201'3). Importantly, the optimal pH level of enzyme activity narrows and shifts toward the 477 478 alkaline pH when enzymes are adsorbed on mineral surfaces of soil particles (McLaren and Estermann, 1957; Aliev et al., 1976; Leprince and Quiquampoix, 1996). Furthermore, the 479 enzymes activity peaks shift towards more acid pH values with increasing IS in the suspensions 480 481 (Goldstein et al., 1964). Given almost one unit difference in the pH between the surface of the colloids and the soil solution and changing chemical composition of the effluents, the pH optima 482 for particular enzyme species, as well as the activity of the colloid associated enzymes, can vary 483 strongly and deviate from that in chemically clean laboratory solution. Therefore, less adequate 484 reproduction of our experimental curves (Fig. 6) by Eq.(5) than those derived from Leprince and 485 Quiquampoix (1996) (Fig. 7) can be attributed to variations in mineral composition of soil 486 487 colloids and chemical composition of effluents.

488 *4.4. Implications of colloid-facilitated transport.*

The colloid-facilitated transport of microbial cells and enzymes is likely part of microbial 489 survival strategy. While microorganisms in soil are primarily attached to the solid surfaces and 490 491 form associations there (e.g., colonies, films, or flocks), their activity is mainly associated with soil solution. Most organic materials entering the soil (plant residue, dead roots, manure) are 492 insoluble or only partly soluble in water, and barely mobile. Therefore, soil microorganisms must 493 either populate most of available pore space or be highly mobile to access new organic inputs. 494 495 However, microbial cells and free enzymes are unlikely capable to travel far in soil pores in searching for new energy sources due to: (i) their relatively large size and slow diffusion rates 496 (i.e., for 1-10 μ m s⁻¹ for enzymes and 7–8 μ m s⁻¹ microbial cells according to Young et al., 1980; 497 Dechesne et al., 2010; and Zhang and Hess, 2019); (ii) small pore volumes available for 498 499 diffusion because of pore discontinuity and low thickness of water menisci in partly saturated 500 soils; (iii) overall short presence of free enzymes in soil solutions due to denaturation and proteolysis, or interaction with clay and organic surfaces (Burns, 1982; Ladd, 1985; Sarkar et al., 501 502 1989; Nannipieri, 1994, 2002). Moreover, most enzymes are irreversibly attached to soil solids, as evident through low extractability of enzymes from bulk soil (Vepsäläinen, 2001; Štursová 503 504 and Baldrian 2011). Therefore, only a small fraction of soil enzymes can potentially freely diffuse in soil pores and reach newly added organic materials. 505

506 Mobilization of colloids and their convective transport through soil macropores during fast 507 water flow events, e.g., irrigation or rainfall, as well as in partly saturated soils, are important for soil microorganisms in exploring new sources of energy. The benefits of colloid-facilitated 508 transport for soil microbes are obvious: (i) much shorter times and longer travel distances as 509 compared with the restricted diffusion; (ii) better protection from protozoa on colloid surfaces 510 (Sarkar et al., 1980; Nannipieri et al., 1982); (iii) energy savings for production of new enzymes, 511 since enzymes can be transported by the colloids; (iv) influx of new partly degraded organic 512 materials from soil surface and their joint transport with microbial cells. Such transport explains 513 appearance of new hotspots of microbial activity commonly observed in soil after heavy rainfalls 514 much better than the diffusion theory does. Soil hotspots and hot moments are defined based on 515 516 time and rates of microbial activity exceeding the average rates in bulk soil (Kuzyakov and Blagodatskaya, 2015). Input of labile substrates to the hotspots triggers microbial activity and 517 thus drives the hot moments. We suggest that, in addition to the substrate-triggered hot spot 518

activation, the colloid-facilitated transport enables enzymes and microbial cells to move quickly
and in relatively large quantities to or with the labile substrates, thus forming new transporttriggered hotspots during high precipitation and preferential flow events.

The results of this study and communication with anonymous reviewers have risen research 522 questions that require further in-depth exploration. Among them there are: (i) how to separate the 523 release of colloids and enzymes from soil matrix and their transport through pores; (ii) how sizes 524 and mineralogy of colloids present in soil affect enzymes association with them; (iii) how sizes 525 and mineralogy of colloids are changing in the effluents, and to what extend these changes affect 526 enzyme association and transport with colloids; (iv) what drives the differences between 527 enzymes in their associations with mineral and biological soil colloids and subsequent transport; 528 (v) are enzymes transported with their producers; (vi) how soil structure affects enzyme locations 529 within the soil matrix and their transport with colloids; (vii) which factors are the dominant 530 drivers of enzyme transport in soils? 531

532

533 CONCLUSIONS

This study revealed a possibility for hydrolytic enzymes (i.e. β -glucosidase, acid-534 phosphatase, xylosidase and cellobiohydrolase) to be transported through soil pores by water 535 fluxes. Strong association of hydrolytic enzymes with fine soil particles and mobility of soil 536 colloids results in their joint convective transport. This transport is affected by ionic strength in 537 pore solution via dissociation and release of soil colloids from soil and alteration of enzyme 538 activity in the transported suspensions. The former effect can be attributed to the shift of optimal 539 pH of enzyme activity near the surface of soil colloids. It remains to be seen how soil texture and 540 541 structure, colloid size and composition, enzyme properties and location in soil pores contribute to their release and transport in the field conditions. 542

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544 **Declaration of competing interest**

The authors declare that they have no known competing financial interests or personalrelationships that could have appeared to influence the work reported in this paper.

547

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Soil texture	Sand	Loam	Sandy loam (SL-S)	Sandy loam (SL-D)	
Sand %	87±2* a **	38±2 b	58±8 c	56±2 c	
Silt %	8±3 a	48±1 c	29±7 b	31±3 b	
Clay %	5±2 a	14±1 b	13±1 b	13±1 b	
SMC [£] %	7±2 a	29±2 c	24±1 b	26±1 b	
POM %	0.31±0.07 a	0.15±0.09 ab	0.15±0.10 ab	0.11±0.01 b	
BD, g cm ⁻³	1.41±0.09 a	$1.62{\pm}0.08$ b	1.73±0.05 bc	1.80±0.05 c	
pH	5.7 ± 0.2	6.0±0.2	5.4±0.3	5.4±0.2	
EC, µS cm ⁻¹	75±19 a	86±24 a	148±28 b	247±37 c	
TN %	$0.03{\pm}0.01~a$	$0.14{\pm}0.03~{\rm c}$	0.11±0.01 bc	0.10±0.01 b	
TC %	0.49±0.11 a	1.53±0.34 c	1.13±0.01 cb	0.98±0.01 ab	
	Enzyme activity in soil, pmol min ⁻¹ g ⁻¹				
β -glucosidase	2.23±0.42 aA	6.53±1.01 cA	6.90±1.05 cA	4.94±0.18 bA	
Phosphatase	1.09±0.26 aB	3.41±0.31 cB	2.77±0.16 bB	2.46±0.06 bB	
Xylosidase	0.20±0.04 aC	$0.30{\pm}0.08~{\rm aC}$	0.82±0.12 bC	0.66±0.12 bC	
Cellobiohydrolase	0.23±0.03 aC	0.46±0.29 aC	0.89±0.04 bC	0.55±0.13 aC	

Table 1. Selected properties of the soils collected from the four studied experimental sites.

* Data presented as mean \pm one standard deviation.

^{**} Means within the same row followed by the same low case letter are not significantly different from each other (p<0.05); means of enzyme activity within the same column followed by the

same upper-case letter are not significantly different form each other (p < 0.05).

760 [£] Particle size $< 50 \ \mu m$

761 SMC denotes the soil mineral colloids, POM denotes the particulate organic matter, BD denotes

the soil bulk density, EC denotes the electrical conductivity, TN denoted the total nitrogen, andTC denotes total carbon.

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Soil texture	Sand	Loam	Sandy loam (SL-S)	Sandy loam (SL-D)
SMC, g l ⁻¹	0.68±0.16* a **	2.89±0.15 c	2.42±0.09 b	2.28±0.11 b
pH	5.9	6.7	6.2	6.4
EC, μ S cm ⁻¹	6	7	8	9
Enzy	yme activity in the	e applied suspensi	ons, pmol min ⁻¹ ml	-1
β -glucosidase	6.03±1.13 aA	9.66±1.22 bA	14.95±2.28 cA	14.06±0.52 cA
Phosphatase	3.13±0.75 aB	10.63±1.23 bA	10.26±0.60 bB	12.61±0.30 cB
Xylosidase	1.09±0.24 bC	0.28±0.08 aB	1.81±0.26 cD	1.60±0.30 cC
Cellobiohydrolase	0.69±0.09 aC	0.48±0.30 aB	3.43±0.16 cC	1.77±0.41 bC
Average enzyme activity associated with coarse mineral colloids (CMC) in the applied suspensions, %				
β -glucosidase	85.7 aA	100.0 aA	94.1 aA	97.3 aA
Phosphatase	97.6 bA	100.0 bA	57.4 aB	99.5 bA
Xylosidase	33.1 aB	100.0 bA	65.4 abB	57.3 abB
Cellobiohydrolase	32.9 bB	100.0 aA	99.5 aA	78.0 aAB
SMC recovered in the column experiment				
SMC recovered, % of applied	49.1±11.8 c	7.3±2.5 a	31.3±3.3 b	40.5±15.0 bc

Table 2. Selected properties of the applied suspensions generated from the soils from the fourstudied experimental sites

^{*} Data presented as mean \pm one standard deviation.

** Means within the same row followed by the same letter are not significantly different from each other (p<0.05); means of enzyme activity within the same column followed by the same upper-case letter are not significantly different form each other (p<0.05).

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Table 3. Estimated enzyme activities in the effluent from the column experiments in Sand and

Loam soils obtained for the eluent amount that replaced 1 pore volume in the columns. Shown

are averages from all columns for the total enzyme activity in the effluent and the proportion of

778	the total activity	from the enzymes	associated with CMC.
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Enzymes	Soil	Total enzyme activity in the effluent,	Activity of the enzymes associated with CMC, %
		pmol min ⁻¹ ml ⁻¹	of total
<i>Q</i> aluggaidaga	Loam	1.4	66
<i>p</i> -glucosidase	Sand	3.0	52
Dhaarhataa	Loam	1.8	79
Phosphatase	Sand	7.6	88
Verlagidage	Loam	0.0	-
Aylosidase	Sand	1.0	50
Callabiahadaalaaa	Loam	0.0	-
Centobionydrolase	Sand	0.9	30

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Figure 1. Experimental design and measurements conducted in the eluents and effluents from the

785 columns.



Figure 2. Concentrations of soil mineral colloids (SMC) and electrical conductivity (EC) in the effluents relative to those in the applied solutions for Sand (a) and Loam (b) soil columns for eluent volumes that passed the columns relative to the pore volumes. Data are combined from 3 columns from each soil. Horizontal line (a) marks the relative SMC in the effluent corresponding to that in the eluent. Different scales are used in (a) and (b) for better data visibility.



797

Figure 3. Activity of the four enzymes in the effluents from Sand and Loam soil columns relative to those in the applied solutions for eluent volumes that passed the columns relative to the soil pore volumes. Zero values indicate enzyme activities below the detection limit. Data are combined from 3 columns of each soil. Horizontal lines mark the relative activity in the effluent corresponding to that in the eluent.



Figure 4. Activity of enzymes associated (filled bars) and not associated (open bars) with CMC in the effluents from Sand, Loam, SL-S and SL-D soil columns for the four enzymes and different eluent volumes passing the columns. Sum of open and filled bars represents the total enzyme activity in the effluents.





Concentration of SMC in the effluent from columns, g L⁻¹



contents in the effluents from Sand and Loam soil columns. Zero values indicate concentration
of colloids and enzyme activities below the detection limit. Lines denote linear regressions

between enzyme activity and SMC contents. Data are combined from 3 columns for each soil.

819 The numbers show the Pearson correlation coefficients at p < 0.1 (*) and p < 0.05 (**),

820 respectively.



822

lonic strength in the effluent solutions (IS), mM

Figure 6. β -glucosidase and phosphatase activities associated and not associated with CMC as a function of ionic strength in the effluents from four soils. The activities are expressed per 1 g of SMC dry mass (a,b) and per 1 ml of effluent (c,d). Black lines show Eq.(5) fitted to the

- 826 experimental data.
- 827
- 828 829
- 029
- 830

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Figure 7. Rate of the catalytic reaction (V) of phosphatase adsorbed on montmorillonite as

function of ionic strength in solutions with different pH values (symbols) reconstructed from

836 Leprince and Quiquampoix (1996) (Fig. 2, Central column). Lines show lognormal distribution

837 fit to the experimental data.

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842 Figure S1. Concentrations of soil mineral colloids (SMC) and electrical conductivity (EC) in the

effluents relative to those in the applied solutions for SL-S (a) and SL-D (b) soil columns for

844 eluent volumes that passed the columns relative to the pore volumes. Data are combined from all845 columns of each soil.



Figure S2. Activity of the four enzymes in the effluents from SL-1 and SL-2 soil columns
relative to those in the applied solutions for eluent volumes that passed the columns relative to
the pore volumes. Data are combined from all columns of each soil. Horizontal lines mark the

850 relative activity in the effluent corresponding to that in the eluent.







858

lonic strength in the effluent solutions (IS), mM

Figure S4. Xylosidase and cellobiohydrolase activities of enzymes associated and not associated with CMC at different ionic strength in effluents from four soils. The activities are expressed per 1 g of SMC dry mass (a,b) and per 1 ml of effluent (c,d).

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