

Elemental composition of natural nanoparticles and fine colloids in European forest stream waters and their role as phosphorus carriers

N. Gottselig, W. Amelung, J.W. Kirchner, R. Bol, W. Eugster, S. J. Granger, C. Hernandez-Crespo, F. Herrmann, J.J. Keizer, M. Korkiakoski, et al.

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90	Key points:
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01	Company where the mass is larger by the material as a specific base of the 11 state
91	• Stream phosphorus is largely bound to natural nanoparticles and colloids

92 • The chemical composition of colloids varies systematically from Northern to Southern
 93 European streams

94 Abstract

95 Biogeochemical cycling of elements largely occurs in dissolved state, but many elements may also be bound to natural nanoparticles (NNP, 1–100 nm) and fine colloids (100-450 nm). We 96 97 examined the hypothesis that the size and composition of stream water NNP and colloids vary systematically across Europe. To test this hypothesis, 96 stream water samples were 98 99 simultaneously collected in 26 forested headwater catchments along two transects across 100 Europe. Three size fractions ($\sim 1-20$ nm, >20-60 nm, >60 nm) of NNP and fine colloids were 101 identified with Field Flow Fractionation coupled to inductively coupled plasma mass-102 spectrometry and an organic carbon detector. The results showed that NNP and fine colloids 103 constituted between $2\pm5\%$ (Si) and $53\pm21\%$ (Fe; mean \pm SD) of total element concentrations, 104 indicating a substantial contribution of particles to element transport in these European 105 streams, especially for P and Fe. The particulate contents of Fe, Al and organic C were 106 correlated to their total element concentrations, but those of particulate Si, Mn, P and Ca were 107 not. The fine colloidal fractions >60 nm were dominated by clay minerals across all sites. The 108 resulting element patterns of NNP <60 nm changed from North to South Europe from Fe- to 109 Ca-dominated particles, along with associated changes in acidity, forest type and dominant 110 lithology.

111

112 1. Introduction

113 Networks of small streams and large rivers transport mobile compounds over long distances 114 and drive their export from the continents to the oceans [Bishop et al., 2008; Dynesius and Nilsson, 1994]. Environmental water samples contain a wide variety of chemical species, 115 116 including simply hydrated ions, molecules, colloidal particles and coarser grains [Stumm and 117 Morgan, 1981]. The partitioning between these species controls elemental cycling, transport 118 and loss processes [Stolpe et al., 2010]. Understanding the distribution of the elements 119 between different physicochemical binding forms is thus an important prerequisite for 120 understanding the mechanisms of aquatic and terrestrial ecosystem nutrition [Benedetti et al., 1996; Hasselloev et al., 1999; Tipping and Hurley, 1992; Wells and Goldberg, 1991]. This is 121 122 especially important for those nutrients that are frequently limiting like phosphorus (P) 123 [Jarvie et al., 2012]. Indeed, the recent review by Bol et al. [2016] highlighted the 124 (unexpected) scarcity of data on colloidal P fluxes in temperate forest ecosystems, which 125 severely limits accurate quantification of forest P nutrition and losses.

126 Research on nutrient acquisition and cycling processes in stream waters and terrestrial 127 ecosystems has often focused on the 'dissolved fraction'. This fraction is frequently 128 operationally defined as the aqueous phase that passes $a < 0.45 \mu m$ filter [Marschner and Kalbitz, 2003 and references therein]. However, it is increasingly recognized that naturally 129 130 occurring nanoparticles (NNP, d = 1-100 nm), and also larger particles belonging to the 131 overall term 'colloids' (d = $1 \text{ nm}-1 \mu m$), can be substantial components within this 132 operational definition of elements present in the 'dissolved fraction'. Colloids smaller than 133 0.45 μ m (=450 nm) are in the present study defined as fine colloids. In natural aqueous phases 134 up to 100% of the total elemental concentrations of metals, and also of specific nutrients like 135 P, can be associated with such particles [Gottselig et al., 2014; Hart et al., 1993; Hill and Aplin, 2001; Jarvie et al., 2012; Martin et al., 1995]. Hence, identifying NNP and colloids in 136 137 water samples is necessary to better understand the cycling and transport of elements in 138 catchments and to determine their biological availability. Headwater catchments are 139 specifically interesting for this analysis because their input variables can be closely defined, 140 thereby facilitating data interpretation. However, it is unknown how the composition and size 141 distributions of NNP and fine colloids vary between headwater catchments on a continental 142 scale. Large-scale studies are advantageous in this context to identify more overarching and 143 broadly applicable principles of NNP and fine colloid composition, and their variations, in 144 natural waters.

145 Prior studies have performed pioneering investigations on particulate P pools in aqueous and 146 terrestrial systems [e.g. Binkley et al., 2004; Espinosa et al., 1999; Sharpley et al., 1995], yet 147 through the applicability of modern particle analysis/fractionation techniques, these 148 functionally defined particulate fractions could be examined more closely and thus allow a 149 more accurate subcategorization of elements in the 'particulate' phase. Further, the focus of 150 dissolved elements as being ions of (hydr)oxized elements in aqueous solution needs to be 151 reconsidered due to the presence of particles within the operationally defined dissolved range 152 [cf. e.g. Gimbert et al., 2003; Lyven et al., 2003; Regelink et al., 2011]. In this regard, Field 153 Flow Fractionation [FFF; Giddings et al., 1976] is a viable tool for these analyses, because it 154 is a nearly non-destructive technique for the fractionation of NNP and fine colloids, thus 155 eliminating the need for pretreatments which can alter the particle composition or size range.

156 The specific reactivity of nanoparticles is high, in comparison to larger sized colloids, 157 [Hartland et al., 2013; Oafoku, 2010] rendering them potentially predominant carriers of 158 nutrients in ecosystems. It has already been shown that NNP can bind the majority of P 159 present in soil solutions [Hens and Merckx, 2001] and stream waters [Gottselig et al., 2014; 160 Gottselig et al., 2017], and that they can even support plant uptake of P from solution [Montalvo et al., 2015]. First results indicate that organic matter, Fe, and/or Al may be major 161 162 binding partners for P in NNP of an acidic forest river system and that the binding of P varies 163 depending on the stream water composition [Baken et al., 2016b; Gottselig et al., 2014]. 164 Under the acidic conditions that characterize many natural settings (particularly many 165 coniferous forest soils), surfaces of metal (hydr)oxides are positively charged and thus act as 166 strong binding partners for negatively charged nutrients like P [Hasselloev and von der 167 Kammer, 2008; Richardson, 1985] and organic matter [Celi et al., 2005]. Adsorbed P and 168 organic matter can even act as stabilizing agents for colloidal suspensions of particles 169 [Ranville and Macalady, 1997; Six et al., 1999]. Organic matter associated with NNP and 170 larger colloids also contains P [Darch et al., 2014]. Some authors even assume that in the 171 smaller size ranges, organophosphorus compounds can also act as the primary building blocks 172 of NNP [Regelink et al., 2013], although these building blocks are more commonly thought to 173 be aluminosilicates, organic matter (org C), or oxides and/or hydroxides of iron (Fe), 174 aluminum (Al), and manganese (Mn) [Baken et al., 2016b; Hartland et al., 2013; Lyven et al., 175 2003; Regelink et al., 2011]. At elevated pH levels in stream water, calcium (Ca) is 176 increasingly present in the colloids and associated with P in the form of Ca phosphates. These 177 Ca phosphate colloids can either be in mineral form or associated on organic/organomineral 178 complexes or clays. In the case of Ca colloids, Ca is enriched in the colloidal phase in

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179 comparison to larger suspended matter [Ran et al., 2000], especially when surface waters 180 drain carbonate-rich soils [Hill and Aplin, 2001]. This highlights that Ca may also be a 181 building block element of colloids [Dahlqvist et al., 2004], depending on stream water 182 chemistry. The pH-dependent speciation of elements and their related affinity to particles give 183 rise to the hypothesis examined in our study, which states that the composition and size 184 distribution of NNP and fine colloids may not be uniform across large regions. If this is not 185 the case, at least the speciation of elements and their presence in particle fractions would be 186 expected to differ between acidic and alkaline stream waters.

187 Despite their important role in element binding, NNP and fine colloids were not considered as 188 a substantial contributor to nutrient cycling in the past [e.g. Vitousek, 1982] and are still often 189 neglected in pioneering studies on the analysis of influential factors on terrestrial nutrient 190 availability [e.g. Fernández-Martínez et al., 2014]. Examining the significance of NNP and 191 fine colloids as well as their composition as a function of forest stream water pH on a 192 continental scale can provide insights into their ecological relevance, particularly if one can 193 use basic water quality parameters to estimate the total elemental concentrations that are 194 associated with NNP and colloids. Better information on the chemical form and reactivity of elements in the putative "dissolved" fraction, in turn, can improve estimates of nutrient 195 196 availability to plants and microorganisms.

For this study, it was hypothesized that NNP and fine colloids are ubiquitous carriers for 197 198 elements in forested European river systems, but that their composition varies along 199 continental-scale gradients. The respective variations in climate, vegetation, soil and 200 freshwater characteristics (e.g. pH) substantially influence the physico-chemical forms of P 201 and other ecologically important constituents in forested freshwaters, thus, potentially 202 revealing systematic controlling effects on NNP and fine colloid concentrations and 203 composition. To test this hypothesis, 96 stream water samples from 26 forested sites from 204 across Europe were collected. These 26 sites cover a wide range of soil types and parent 205 materials, under deciduous or coniferous forests, over very differently sized catchments, and 206 spanning different elevations and topographic gradients (Table 1). FFF [Giddings et al., 1976; 207 Hasselloev et al., 1999; Regelink et al., 2014], coupled online to high-precision elemental 208 detectors, was used for the fractionation of NNP and colloids. To untangle the importance of 209 single factors, a correlation matrix between all particulate elemental concentrations, total 210 elemental concentrations, and the basic site parameters such as MAT, MAP, forest coverage, 211 slope inclination, and runoff was also investigated.

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212 2. Materials and methods

213 2.1 Sampling

214 Twenty-six sites throughout Europe were selected along two transects, one from northern 215 Finland to Portugal and the other from Scotland to Greece (Figure 1, Table 1). Each site is a 216 forested headwater catchment with low-intensity forest management practices, high tree 217 coverage and without inflows from urban or agricultural settings (for site abbreviations, see 218 Table 1). Mean annual temperatures ranged between -1.4 °C (PA, Finland) and +15.9 °C (FR, 219 CO and PR, Italy) and mean annual precipitation ranged between 450 mm (BPC, Spain) and 220 2426 mm (LÜ, Switzerland) (Table 1). It was possible to assign a clear dominant forest type 221 for 25 of the 26 sites (either coniferous or broadleaf), while the BPC site had 50% coniferous 222 and 50% broadleaf tree species. Additionally, data on catchment size, average elevation, 223 average slope, percentage forest cover and mean annual runoff were collected where available 224 (Table 1). All data were provided by the current site operators. During sampling, the electrical 225 conductivity, pH and water temperature of the water samples ranged substantially from 226 13 µS/cm (SB) to 1775 µS/cm (BPC), from 4.2 (GS) to 9.5 (RS) and from 1.0 °C (PA) to 227 19.9 °C (RS) (for complete data of all main stream and tributary samples, see supplementary 228 material, Table S1). Up to six samples were taken per site in May 2015 in order to derive a 229 snapshot of NNP and fine colloid composition and their relevance for P transport across 230 Europe at a given time of the year. The sampling locations were defined in consultation with 231 the site operators to best reflect main and tributary streams of each catchment, similar to the 232 approach of Gottselig et al. [2014]. Sampling was conducted during base flow conditions. The 233 sampling resulted in a total of 96 samples taken in duplicates in pre-cleaned PE and glass 234 bottles and shipped cooled to the lab; analysis of NNP and fine colloids was completed within 235 1 week after sampling. Own unpublished data indicate that during one week there is no 236 significant alteration in the size distribution of NNP and fine colloids, despite selected risks of 237 minor particle re-aggregation in the larger size range. To minimize the risk of systematic 238 errors due to sampling storage and pretreatment, all samples were handled, treated and 239 analyzed in equal manner.

240

Figure 1

241

Table 1

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242 2.2 Asymmetric Flow Field Flow Fractionation

243 Fractionation of the particles was performed with Asymmetric Flow Field Flow Fractionation (AF⁴). Briefly, a 0.5 mm spacer, a 1 kDa polyether sulfone (PES) membrane and a 25 µM 244 NaCl eluent solution were applied. 5 mL of the PE bottled samples were injected into the AF⁴ 245 246 system at 0.3 mL/min tip flow and 3 mL/min cross flow and focused for 30 min with 247 3.2 mL/min focus flow. Thereafter, a 30 min linear cross flow gradient down to 0 mL/min 248 with subsequent 40 min constant elution at a detector flow of 0.5 mL/min was applied. 249 Reference materials (Suwanee River NOM, Humic Acid Standard II and Fulvic Acid 250 Standard II, International Humic Substances Society, Denver, USA; Sulfate Latex Standards 8% w/v 21- 630 nm; Postnova Analytics, Landsberg, Germany) were used with the same AF⁴ 251 252 conditions used for the samples for calibration of the particle diameters included in each size 253 fraction. No reference material exists that covers the diverse particle morphologies and 254 elemental concentrations of environmental samples. Therefore, the specified hydrodynamic 255 diameters of the particles are equivalent sizes based on the elution time of the reference 256 materials [c.f. Neubauer et al., 2011; Regelink et al., 2013; Stolpe et al., 2010]. The lower size 257 range of the first fraction was estimated according to the molecular weight cut-off (MWCO) 258 of the membrane.

259 Online coupling of a UV detector (254 nm) served to initially determine peak elution and turbidity of the particle fractions. A dynamic light scattering (DLS) device was coupled for 260 261 online size measurements of the largest size fraction. Blank runs inserted between sample 262 runs in the measurement sequence showed no pronounced peaks. A quadrupole inductively 263 coupled plasma mass-spectrometer (ICP-MS) with helium collision cell technology (Agilent 264 7500, Agilent Technologies, Japan) and for size-resolved detection of organic carbon an 265 Organic Carbon Detector (OCD; DOC Labor, Karlsruhe, Germany) were coupled online to the AF⁴. The ICP-MS allowed size-resolved detection of Al, Si, P, Ca, Mn and Fe, and the 266 267 OCD allowed size-resolved detection of organic carbon. The ICP-MS system was calibrated 268 through a post-column [Nischwitz and Goenaga-Infante, 2012] multi-point linear calibration injected via a T-junction between the AF⁴ and the ICP-MS at 0.5 mL/min AF⁴ injection flow 269 270 (= detector flow; no cross flow). The standard solutions (0 μ g/L, 25 μ g/L, 100 μ g/L, 250 μ g/L 271 and 500 µg/L) and the internal standards Rh and Y were dissolved in 0.5 mol/L HCl. This calibration technique is more complex than injecting the calibration standards directly into the 272 273 AF⁴ system, but allows more precise correction of instrumental drift and calibration to higher 274 concentrations without potentially contaminating the following sample (because the standards

- do not pass through the AF⁴). The variations of the ICP-MS peak areas for triplicate measurements of a representative sample were calculated to be 5.9% for P, 7.6% for Al, 14.0% for Si, 5.3% for Mn and 15.6% for Fe. The limit of detection was 0.1 μ g/L for P, 0.01 μ g/L for Al, 3.3 μ g/L for Si, 0.01 μ g/L for Mn and 0.02 μ g/L for Fe. Quantitative atomization of the particles in the plasma has already been shown by *Schmitt et al.* [2002].
- 280 For the OCD coupling, 1 mL sample volume from the glass bottles was injected and focused 281 for 10 min; the remaining parameters were the same as for ICP-MS coupling. The OCD 282 system was calibrated using dilutions of Certipur® liquid TOC standard (EN 1484-H3/DIN 283 38409-H3, Potassium hydrogen phthalate in water, stabilized, 1000 mg/L; Merck Millipore 284 109017) in double-distilled water at concentrations of 0.05 mg/L, 0.1 mg/L, 0.5 mg/L, 285 1.0 mg/L, 3.0 mg/L and 5.0 mg/L. The same calibration standards were also used in 286 determining the total organic carbon in the stream water samples. For this determination, the 287 AF^4 channel was bypassed by connecting the tip inflow tubing to the detector outlet tubing of the channel. The runtime of the AF^4 method for this data acquisition was 20 min at 288 0.5 mL/min tip flow. The relative standard deviation of the organic carbon concentration for 289 290 triplicate measurements of a representative sample was calculated to be 2.2%. The limit of 291 detection for organic C was 0.01 mg/L.

292 2.3 Quality control

293 First investigations on the stability of NNP and colloids were conducted prior to the sampling 294 campaign to elucidate which sampling, storage and transport procedures best reflect natural 295 conditions at the time of measurement. This resulted in a sampling of unprocessed stream 296 water with polypropylene containers; only samples for organic carbon analysis were taken 297 with pre-cleaned and pre-equilibrated glass vials. Samples were always taken in order moving 298 upstream, from the catchment outlet to the headwaters. Containers were preconditioned in 299 triplicates with stream water before the sample was taken from the center of the flowing 300 stream without disturbing the sediment. Larger-sized particulates (e.g., visible parts of leaves) 301 were not included in the water sample. For transport and storage, the samples were kept at a 302 cool to ambient temperature, but neither were they frozen nor did their temperature ever 303 exceed the stream temperature at sampling. Sample analysis was conducted as soon as 304 possible after sampling, especially for organic carbon analysis. The analyses were performed 305 in the order the samples arrived. For a more detailed discussion of circumstances affecting 306 colloidal stability, see Buffle and Leppard [1995].

307 Immediately prior to analysis, samples were homogenized through agitation, then filtered 308 through 5 µm cellulose nitrate filters (GE Healthcare, Munich, Germany) to avoid clogging of the micrometer-sized AF⁴ tubing. No interference of cellulose and/or cellulose nitrate 309 310 compounds with the org C signal was expected because both are insoluble in water 311 [Hagedorn, 2006; Roth, 2011]. Still, the filters were pre-rinsed with 15 mL double-deionized 312 water to eliminate eventual bleeding compounds. Caking was prevented by filtering only 313 small sample volumes up to 15 mL. Additionally, more than one filter was available per 314 sample, but this was not needed due to the low turbidity of the samples. Filtration at 0.45 µm 315 was purposely not performed, to avoid the risk of excluding particles not specific to the given 316 size due to unknown morphological heterogeneity of the natural particles. Avoiding filtration 317 also avoided the risk of membrane clogging when filtering occurs close to target size ranges 318 of the analytes, which can result in a severe risk of underestimating NNP and fine colloidal 319 concentrations [Zirkler et al., 2012].

The recovery of NNP and fine colloids fractionated by AF^4 is greatly influenced by 320 interactions of the natural particles with the membrane, particularly during focus time and at 321 322 long and/or high cross flows. For natural samples, a portion of the sample was thus withdrawn 323 from the fractionation process and measured by independent ICP-MS analysis (cf. section 3.1), thereby allowing us to relate element yields after fractionation to those without AF^4 324 treatment. This analysis showed that the different elements associated to NNP and fine 325 colloids ranged up to 99.5% of total elemental concentration, showing that generally there 326 was no major particle loss. A more in-depth investigation of the AF⁴ recoveries with synthetic 327 iron oxyhydroxide colloids revealed recoveries between 70 and 93% [Baken et al., 2016a]. 328 329 These results are encouraging also for natural samples due to the similarity of the particle 330 constituents.

331 2.4 Analysis of raw data

ICP-MS raw data were collected in counts per second (cps) using the MassHunter 332 Workstation Software (Agilent Technologies, Japan) and OCD raw data were recorded in 333 volts detector signal (V) with the AF⁴ analytical software (Postnova Analytics, Landsberg, 334 335 Germany). Raw data were exported to Excel® (Microsoft Corporation, Redmond, USA) for 336 baseline correction, peak integration and conversion of peak areas to concentrations through 337 multipoint linear calibration. Different pools of elements were considered in this study: a) elemental concentrations assigned to the 1st, 2nd or 3rd size fraction of NNP and fine colloids 338 339 (see section 3.1 for explanation), b) all particulate elemental concentrations, reflecting the

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sum of (e.g.) Fe concentrations in the 1^{st} , 2^{nd} and 3^{rd} fractions combined, and c) the total concentration of an element in the sample prior to fractionation. The collective term 'colloids' is occasionally used in our study as an encompassing term for the whole size range of nanoparticles and colloids. Concentrations are primarily given in µmol/L for Al, Si, P, Ca, Mn and Fe, and in mmol/L for org C.

Stream water pH at sampling time was classified according to the Soil Survey Division of the Natural Resources Conservation Service, U. S. Department of Agriculture [*USDA*, 1993], in order to clarify the influence of stream water pH and associated covariates on the relationships between NNP and fine colloid concentrations and total concentrations. According to this classification, acidic is defined as pH<6.6, neutral pH between 6.6 and 7.3 and alkaline pH values >7.3. Thus, pH was transformed into a semi-quantitative variable, where each range of values is assigned to a given category.

352 Predictability of elemental concentrations through total concentrations was assessed through 353 log_{10} transformation of the elemental concentrations in the fractions and the total sample 354 concentrations. Here, we only considered regressions that achieved an $r \ge 0.71$. Further, sites 355 were classified according to location (North, Middle, South) in Europe. To test if this zoning 356 was able to differentiate element distributions of the NNP and other fine colloidal fractions, 357 we focused on particulate Fe as redox-sensitive element, and particulate Si as an indicator of 358 siliceous bedrock and clay minerals. Also, we classified the study sites by major soil types 359 (dystrophic, eutrophic and semi-terrestrial), and bedrock lithologies (siliceous, calcareous, and 360 flysch), as well as two main forest types (coniferous/needle vs. broadleaf vs. mixed, see Table 361 1). Dystrophic soils included Podzols, as well as coniferous Cambisols, Leptosols and 362 Regosols; eutrophic soils were deciduous Cambisols, Leptosols, and Mollisols; semi-363 terrestrial soils were Fluvisols and Gleysols (the sole Histosol was not included in this group 364 comparison). Siliceous bedrock included all bedrock types except calcareous stone, limestone 365 and flysch. Flysch was assigned to a specific bedrock group, because it usually contains both 366 carbonates and silicate minerals; the respective soils sampled were Gleysols (Table 1).

Further, the dependency of elements in the NNP and fine colloid fractions on the total elemental concentration as well as the potential predictability of NNP and fine colloidal composition were analyzed through correlation analysis with Pearson r coefficients, pairwise deletion of missing data (JMP 12.2.0, SAS Institute Inc., USA) and significance testing using non-parametric group comparisons with the Mann-Whitney U-test for comparisons between two sample sets and with the Kruskal-Wallis ANOVA for comparisons among more than two

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373 sample sets (Statistica, Version 13, Dell Inc., Tulsa, USA). To facilitate the examination of 374 controlling factors of NNP and fine colloid composition, correlation matrices followed by 375 principal component analysis (Statistica, Version 13, Dell Inc., Tulsa, USA) served as an 376 additional tool to reduce the number of variables and reveal a first structure in the 377 relationships between the variables. Through this, we aimed at an identification of the site 378 parameters, which are influential on particulate elemental concentrations. Varimax raw was 379 applied as the rotational strategy for the analysis, to maximize the variances of the squared 380 raw factor loadings across variables for each factor. Here, casewise deletion of missing data 381 was undertaken to ensure that the same number of cases entered into every analysis. Initially, 382 calculations of the Kaiser-Meyer-Olkin Measure of Sampling Adequacy (KMO value; IBM 383 SPSS Statistics 22) were performed to extract site parameters suitable for PCA analysis. 384 When using all data, KMO was 0.5 and the PCA result was not stable against random 385 elimination of input data; only with selected input variables a KMO > 0.6 was achieved and 386 the PCA was stable against variations in input data. As a follow-up tool, stepwise multiple 387 regression was used, but it did not yield meaningful results due to the remaining complex 388 interplay of the data.

389

390 3. Results and discussion

391 3.1 Fractionation of nanoparticles and fine colloids

Similarly to findings for a forested watershed in Germany [Gottselig et al., 2014], the 392 application of the AF⁴ technique to other forested sites across Europe revealed distinct 393 394 fractions of nanoparticles and fine colloids depending on the element investigated (Figure 2a, 395 b; cf. supplementary material, Table S2). In total, three fractions of NNP and/or fine colloids 396 were distinguishable for most (out of 96 samples, in 58.3% org C and Si were found in all 397 three fractions, in > 85% for Ca and P, and in > 95% for Al, Mn and Fe; cf. supplementary material, Table S2) samples. The fractograms included a peak of small-sized nanoparticles (1st 398 fraction), a second peak consisting of intermediate-sized nanoparticles (2nd fraction) and a 3rd 399 400 peak containing the largest-sized nanoparticles and fine colloidal matter (see also peak 401 separation by dashed lines on the x-axis in Figure 2). The peak pattern for each sample is 402 represented by the fractograms of each element (see Figure 2). The second peak as shown for 403 Krycklan, Sweden, was mainly defined through the Ca signal (violet; Figure 2a, b) due to the absence of other elemental peaks in the 2nd fraction. P was variably detected in one, two or 404

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three size fractions (Figure 2c, d; cf. supplementary material, Table S2). Hence, three
different size fractions were isolated in this work, in close agreement with *Stolpe et al.* [2010],
who also found three to four fractions of NPP and fine colloids in the lower Mississippi River,
the largest river in North America.

409 Nanoparticles were the exclusive constituent of the first two fractions and, based on 410 hydrodynamic diameters, accounted for approximately 20% in the third fraction. The first and 411 second fractions consisted of nanoparticles with standard equivalent hydrodynamic diameters 412 ranging from 1 kDa [equivalent to 0.66 nm, equation 2.2, *Erickson*, 2009] to 20 nm, and from 413 above 20 nm to 60 nm, respectively. The third fraction included nanoparticles larger than 414 60 nm up to fine colloids of approximately 300 nm in diameter. The DLS measurements 415 revealed this maximum particle size for all measured samples [c.f. Gottselig et al., 2014; 416 Gottselig et al., 2017]. Hence, all detected NNP and fine colloids fell into the operationally 417 defined 'dissolved phase' ($< 0.45 \,\mu\text{m}$). Substantial signals of the elements Fe, P, Mn, Al, C 418 and Si were recorded in all three size fractions with varying intensities, confirming 419 widespread occurrence of NNP and fine colloids in the size range < 450 nm.

Figure 2

421 The fractograms for P are markedly different among three sites spanning across one transect in Europe (Pallas in North Europe, Bode in Middle Europe, and Ribera Salada in South 422 423 Europe; PA, BO and RS in Figure 2c), consistent with the hypothesis that NNP and fine 424 colloid composition and size distribution vary across the continental scale. More generally, 425 we observed differences in the P distribution within NNP and fine colloids among the various 426 forested headwater catchments in Europe: i.e., sometimes the particulate P was associated 427 with clearly distinct fractions (e.g. PA; Figure 2c), whereas for other samples a less distinct 428 fractionation of P (e.g. BO; Figure 2c) was observed. This exemplary described variation of 429 elemental concentrations in the three fractions was observed for all recorded elements, 430 supporting the general hypothesis that there are differences in the elemental composition 431 between fractions and among the selected European streams. These patterns were found to be 432 related to site-specific properties such as climate, water chemistry, soil type and total stream 433 water elemental concentrations, i.e., we tested in the following to systematize the differences 434 according to element composition and site properties.

Version p

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436 The percentages of elements bound to NNP and fine colloids (i.e., percentage of all particulate 437 elemental to total elemental concentration) demonstrated the substantial contribution of NNP 438 and fine colloids to element fluxes in natural waters. The average percentages of elements 439 found in the NNP and colloid phases were 53% for Fe (within the bounds for an interquartile 440 range of 42-65%), 50% for P (36-70%), 26% for Mn (4-47%), 41%, for Al (28-56%), and 441 20% for organic C (5-28%), but only 2% for Si (0.1-0.4%) and 4% for Ca (0-5%; n=96). The 442 respective median values were 55% (Fe), 51% (P), 10% (Mn), 37% (Al), 11% (org C), 0.2% 443 (Si) and 1% (Ca). Further, up to 99% of Fe, 96% of P and Mn, 95% of Al, 92% of org C, 46% of Si and 27% of Ca were found associated with NNP and fine colloids, relative to the bulk 444 445 elemental concentrations measured offline in the untreated samples. Overall, the percentages 446 reflected a substantial contribution of the NNP and fine colloidal fractions within the 447 operationally defined 'dissolved' elements Fe, P, Al, org C and Mn (in descending order). 448 Previous research on the significance of colloid-bound elements within the operationally 449 defined 'dissolved' fraction also indicated maximum Fe binding in fine colloidal form 450 between 80 to 100% with averages between 50 and 90% [Hill and Aplin, 2001; Jarvie et al., 451 2012; Martin et al., 1995], for organic C between 40 and 80% with averages between 20 and 452 60% [Jarvie et al., 2012; Martin et al., 1995; Wen et al., 1999], and for Al around 40 to 50% 453 with averages around 45 to 55% [Hill and Aplin, 2001; Jarvie et al., 2012]. Hill and Aplin 454 [2001] determined that the fine colloidal fraction accounted for up to 50% of Mn (average 455 23%) and up to 30% of Ca (average 20%), but only up to 10% of Si (average 0%). Dahlqvist 456 et al. [2004] found an average of 16% fine colloidal Ca in an Arctic river and in Amazonian 457 rivers (also assessed with FFF). Data on total P are scarce, but Jarvie et al. [2012] reported 458 that a fraction of up to 90% (averaging 66%) of organophosphorus compounds was associated 459 with fine colloids in mixed land-use sites, and *Missong et al.* [2016] and *Jiang et al.* [2015] 460 also detected organophosphorus in the NNP and fine colloid fractions in soil samples. In 461 summary, the present data indicate that the particulate form of elements is substantial in 462 streams, and that there are varying contributions of NNP and fine colloids to overall element 463 fluxes across Europe.

464 *3.3 Predictability of NNP and fine colloid elemental composition*

The discrepancy between NNP and fine colloid and total elemental concentrations is potentially due to dissolved species in size ranges below the membrane MWCO when excluding the presence of larger particles. Two types of relationships between NNP and fine 468 colloid concentrations and total concentrations could be observed, independent of the fraction 469 in question. For Si, P, Ca and Mn, we found scattered relationships between total 470 concentrations and NNP/fine colloid concentrations (Figure 3, left panels). The concentrations 471 of these elements that were bound to NNP and fine colloids did not change systematically 472 with total elemental concentrations. All three size fractions of these elements showed such 473 scattered relationships, although their median concentration varied (Table 2). By contrast, for 474 Fe, Al and org C, we found positive log-log relationships between total concentrations and 475 colloidal concentrations (Figure 3, right).

476 The regression slopes of the log-log relationships between the particulate and total element 477 concentrations differed among the three size fractions of Fe, Al and org C (Table 2), which 478 could reflect differences in Fe, Al and org C speciation as stream water variables change, but 479 could also be related to site differences in the surface properties of mineral binding partners (which were not investigated here). Most interestingly, the log-log slope of the 1st size 480 481 fraction Fe was 1.0 (Table 2; Figure 3, right panels). This implied that across all European 482 sites, on average, a constant proportion of total Fe was present in this very fine 483 nanoparticulate fraction (<20 nm), and thus was independent of their stream water pH. This proportion was estimated to be 15%. Slopes for Fe in the 2nd and 3rd fraction, as well as for Al 484 and org C for of all three fractions, were not parallel to the 1:1 line (not shown), instead 485 486 indicating power-law relationships (rather than a fixed NNP or fine colloid fraction). In 487 contrast, for the elements that did not change systematically with total elemental 488 concentrations (Ca, Si, Mn, P) the median varied among the different size fractions as well as 489 the percentage of particulate elemental concentration present in the fractions (cf. Table 2). 490 The reasons for the different regression slopes and medians are still unclear and warrant 491 investigation in future studies, potentially in combination with measurements of discharge-492 dependent concentration variations as shown in e.g. Trostle et al. [2016].

493

494

Figure 3

Table 2

The majority of acidic sites were located in Northern Europe, neutral sites in Middle Europe and alkaline sites in Southern Europe (Table 3). These three geographical regions are shown in Figure 1, with borders at around 56 and 48°N. Most of the sampling sites in the Middle Europe region, i.e., between these latitudes, had neutral pH. A fairly equal amount of Middle Europe sampling sites could be involved among the three pH classes (Table 3), however, they may not constitute a grouping by themselves, but rather a transition zone between theNorthern and Southern European regions.

502 With some overlaps, the pH differences and the North-Middle-South groupings were reflected 503 in the distribution patterns of elements among NNP and fine colloids (Figure 3). The elements 504 with positive log-log relationships (e.g., right panels in Figure 3) exhibited some data point stratification, with lower elemental NNP and fine colloid concentrations at the Southern 505 506 European and alkaline sites (red dots in Figure 3), and higher concentrations at the Northern 507 European and acidic sites (green dots in Figure 3). Whether the main driver of this variation 508 was really pH, or a co-variate related to pH, warrants further attention (see also discussion 509 below). The total concentrations of Ca, Si, Mn and P increased from the Northern, acidic sites 510 to the more alkaline streams in the South (Figure 3, left panels). Such behavior could well be 511 expected for Ca but not necessarily for the other three elements [e.g. Song et al., 2002]. The 512 concentrations of Ca, Si, Mn and P within NNP and fine colloid fractions did not vary 513 systematically with total concentration, but were roughly constant in their median value as the 514 total concentrations increased from the Northern, acidic, sites to the more alkaline, Southern, 515 sites. This implies that the elemental proportions that were bound to NNP and fine colloids 516 decreased from North to South.

517 Eleven data points at three sites did not follow the clear regional stratification in the relationships between total and NNP/fine colloid concentrations (Figure 3); six Southern 518 519 water samples for Ca (scattered relationship; two at SC and four at LZ) and five Northern 520 water samples for Fe (all at AM; see Table 1 for site abbreviations). A concise explanation of 521 their anomalous behavior could not be found, especially because the other elements did not 522 exhibit this anomalous behavior, either in these samples or in any other samples. Site 523 characteristics (Table 1) and stream water parameters at sampling (supplementary material, 524 Table S1) showed no evident outliers in comparison to other sites within the same 525 geographical region, with the exception of the three sites closest to the coast.

526

Table 3

527 The presented data clearly showed the possibility to predict the concentrations of particle-528 bound elements, as some were independent of total concentrations (Ca, Si, Mn, P) while 529 others were linked (Fe, Al, org C). Future research should address potential temporal patterns 530 of NNP and fine colloid composition across an even larger, global scale.

531 *3.4 Controlling factors of NNP and colloid composition*

532 The variable composition of NNP and fine colloids between sites and between sampling 533 locations of one site can potentially be linked to differences in site parameters on larger (e.g. 534 MAT or forest cover) or smaller (e.g. pH value or electrical conductivity) scale. As would be 535 expected for large data sets, many statistically significant (p < 0.05) correlations for, e.g., all 536 particulate org C, Al, Ca and Fe concentrations were found, but few strong correlations (r >537 0.71) were found with site parameters (supplementary material, Table S3). Total Ca 538 concentrations were tightly correlated with electrical conductivity (r = 0.96), which is 539 unsurprising because Ca is often a significant fraction of total ionic strength, while all 540 particulate Ca showed strong correlations to all particulate org C (r = 0.84) followed by 541 catchment size (r = 0.81). In contrast, correlations between all particulate Si, P, Ca and Mn 542 and their total concentrations were weak (r < 0.02; partially negative values), as were the 543 correlations between total element concentrations of, e.g., Si and Mn and concentrations of other elements or site parameters (r < 0.49). However, concentrations of all particulate Si 544 545 were positively related to concentrations of all particulate Al (r = 0.72), and especially strongly (r = 0.98) for Si and Al in the 3^{rd} size fraction, reflecting the presence of both 546 547 elements in layer silicates such as clay minerals.

When particulate concentrations were correlated with site parameters, these correlations were rarely consistent among all size fractions. Marked differences were observed, for instance, between correlations of site parameters with the 1st and 2nd fractions vs. 3rd fractions for Al and Si (supplementary material, Table S4). The correlations thus showed that relationships between site parameters and NNP and fine colloid concentrations differed, depending on the fraction in question, and that NNP and fine colloid concentrations cannot be explained through linear relationships alone.

At a KMO criterion > 0.6, we did not observe significant alterations in factor loadings with random elimination of data, and PCA was run stably with all particulate elemental concentrations, pH, MAT, conductivity, water temperature and elevation, for instance. The PCA extracted four factors with eigenvalues > 1 that jointly explained 75% of the total variance. The resulting factor loadings revealed four distinct groupings of variables: 1) site parameters such as pH, water temperature and MAT, 2) particulate Al and Si, 3) particulate org C, Ca, and Fe, and 4) particulate P (see supplementary material, Table S5).

Mattsson et al. [2009] showed a link between DOC and climatic and topographic factors 562 563 through the strong positive correlation between DOC and latitude; here, however, the 564 colloidal org C was rather related to that of Ca and Fe, this grouping even remained stable 565 when including other site parameters into PCA at worse KMO test results. Fe (and Ca) are 566 known to be key elements in reacting with soil C during microaggregation processes [see e.g. 567 Kögel-Knabner and Amelung, 2014], i.e., our data would be in line with particles being 568 released from riparian soils during riverbank erosion. Besides, our data showed that the NNP and fine colloid concentrations of Al and Si loaded highest at onto factor 2, thus supporting 569 570 the results from above-mentioned correlation analyses (supplementary material, Table S3) and 571 indicating that clay minerals are always present within the third fine colloid fraction. Finally, 572 the PCA analyses revealed that one factor can solely be assigned to particulate P. This is 573 congruent with the simple correlation analyses and confirms that P cannot be predicted by 574 simple linear statistical approaches across different geographic regions (see section 3.2). 575 Overall, the factor loadings thus confirmed that different processes controlled the fate of 576 different elements, while there was apparently no simple linear relation to site factors. Hence, 577 we elucidated the contribution of soil, geology, vegetation and pH class as additional controls 578 of NNP and FC properties.

579 The pH value is often a master variable controlling chemical forms in solution [Perry et al., 580 2008] and may also determine the size and elemental composition of NNP and fine colloids [Neubauer et al., 2013]. pH-dependent element abundances in a 'dissolved' state are well 581 582 understood in the context of nutrient availability [Perry et al., 2008], but not yet with respect 583 to NNP and fine colloids. The geographic zoning of the streams more or less coincided with three dominant stream water pH classes (Figure 1, Table 3). As shown for the 3rd size fraction 584 585 in Figure 4a, though also valid for all particulate Si (supplementary material, Figure S1), 586 plotting Fe against Si clearly separated the acidic (Northern) streams from the alkaline (Southern) ones. Similar results were also obtained when, e.g., plotting particulate Fe against 587 particulate Ca as an indicator of calcareous bedrock (Figure 4b; illustrated for the 2nd fraction; 588 valid also for the 1st fraction but not for the 3rd one; supplementary material, Figure S1). The 589 590 findings can be reconciled with a study on Fe in boreal catchments across different pH values 591 [Neubauer et al., 2013]. Fe entering stream waters is instantly oxidized and forms 592 Fe(oxy)hydroxides or complexes with organic matter [Ekstrom et al., 2016], while formation 593 of Ca- or Si-rich particles is likely independent from such processes. Overall, and despite the 594 many complex factors regulating specific element concentrations in NNP and larger colloids, 595 there are thus strong indicators of a geographic (or pH-dependent) zoning of the composition

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596 of these particles. Forest biomass and soils, for instance, co-vary along our geographic 597 transects, potentially affecting the pools and formation of humus and secondary minerals. 598 Relevant factors could also be any co-variates that induce non-linear variations in element 599 speciation across the pH range of 4.2 to 9.5 (Figure 4; supplementary material, Table S1).

600

Figure 4

601 3.5 Impact of specific soil types and land-use on element concentrations

Whengrouping the sites, e.g.,according to soils, bedrock, and dominant forest type, at least one particulate elemental concentration was affected (Table 4), and the effects found for total particulate elemental concentrations were largely reflected in the individual size fractions as well (supplementary material, Tables S6-8).

606 Sites characterized by semi-terrestrial soils with better hydraulic connectivity to the streams 607 showed significantly higher concentrations of colloidal Si than their counterparts with 608 terrestrial soils (Kruskal-Wallis ANOVA; p<0.05; Table 4). However, the more frequent 609 redox cycles at the semi-terrestrial sites [Blume et al., 2009] did not correlate with higher 610 concentrations of redox-sensitive elements, at least as this particular sampling time. Since semi-terrestrial soils are usually poor in soil structure [Blume et al., 2009], higher particulate 611 612 Si concentrations in the streams could be related to riverbank erosion, but more data at higher 613 temporal resolution would be needed to investigate this hypothesis.

614 Stream composition frequently correlates to the dominant bedrock [Krám et al., 2012]. For 615 the elements with positive log-log relationships (org C, Al, Fe; right panels of Figure 3), a 616 significant differentiation between siliceous and calcareous rocks was found. The 617 concentrations of org C, Al, Si, and Fe were higher for the siliceous sites (Table 4). Overall it 618 appeared that element release from siliceous sites was facilitated relative to the calcareous 619 counterparts, which was accompanied by the release of these elements in all fractions of NNP 620 and fine colloids (supplementary material, Table S6-S8). Particularly high concentrations of 621 Al and Si were found in the streams with flysch (Table 4), likely reflecting riverbank erosion of clav minerals (3rd colloidal fraction; supplementary material, Table S8) from the Gleysols 622 in vicinity of the rivers. Note, however, that among our sites, bedrock classification yielded 623 624 substantially more data points for siliceous than for calcareous rocks and flysch.

When the data were grouped according to the dominant forest type, the coniferous class differed significantly from the broadleaf class for all particulate org C, Al, P, Ca and Fe 627 concentrations, even for those of Si and Mn (Table 4; the one mixed stand was not included in 628 this analysis). Notably, the element concentrations were generally larger in the coniferous tree 629 stands than the broadleaved ones, consistent with the tendency for coniferous stands to have 630 more acidic stream waters [Allaby, 2006] and usually siliceous bedrocks (Table 1). In fact, 631 almost 90% of the acidic sites were associated with dominant coniferous forest types (cf. 632 Table 1 and supplementary material, Table S1). These conditions might generally favor NNP 633 and fine colloid formation because low pH increases the density of positive charges of metal 634 oxides and organic matter, which can then bind to negatively charged surfaces such as fine 635 clay minerals as also outlined before [see also Tombácz et al., 2004]. Indeed, when grouping 636 our sites according to main pH classes as outlined above, the concentrations of particulate org. 637 C, Al, and Fe were generally larger for the acidic sites (in Northern Europe) than for the 638 alkaline ones (in Southern Europe). Interestingly, however, P did not exhibit this pH effect despite pH is the main variable affecting P availability and mobility in soils; hence, further 639 640 analyses are necessary to help explain this effect.

Table 4

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642

643 4. Conclusions

644 Stream water sampling at 26 forested headwater catchments across Europe revealed 645 substantial binding of Fe, P, Al, Mn and org C to NNP and fine colloids. Overall, up to an average of 53±21% of the total content of these elements in the stream waters occurring in 646 three distinct particulate size fractions (1st: < 20nm, 2nd: 20–60nm, and 3rd size fraction: >60– 647 300nm). Particulate concentrations of org C, Fe and Al increased with total concentrations of 648 649 these elements from the South to the North, coincident with decreasing pH values and 650 increasing portions of coniferous forests and siliceous bedrock. The sampled sites could be 651 divided into sites that were characterized by the presence of Ca-containing NNP and fine 652 colloids in alkaline stream waters, and sites with an increasing predominance of Fe-NNP and 653 fine colloids (acidic stream waters) because the Fe concentration superimposed Ca. This distinction was found for the 1st and 2nd NNP fractions, whereas 3rd size fraction (larger NNP 654 655 and fine colloids) mainly consisted of Al and Si-bearing clay minerals at all sites.

656 Interestingly, substantial amounts of P, which previously had been assigned to the 657 operationally defined dissolved phase, were found to be associated with NNP and fine 658 colloids. While P was mainly bound to Fe-containing particles of the 1st size fraction in more

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659 acidic Northern European headwaters, it was associated with Ca-bearing particles of the 2nd

660 size fraction in Southern European headwaters. Also, variations of total P were not correlated

661 straightforwardly with variations of site characteristics across our sampling sites. Further

662 efforts will be needed to better understand the complex interplay between total and colloidal P

663 fluxes across the globe.

664

665

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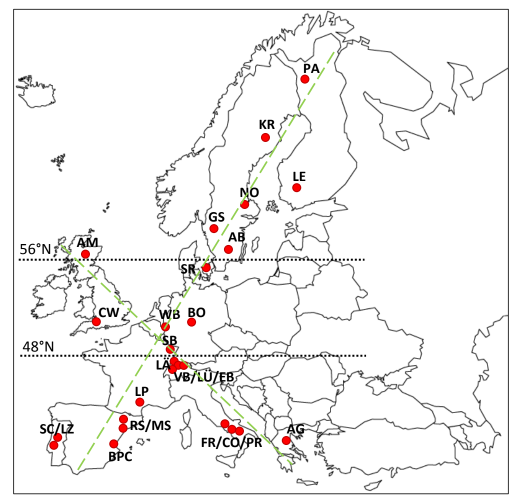
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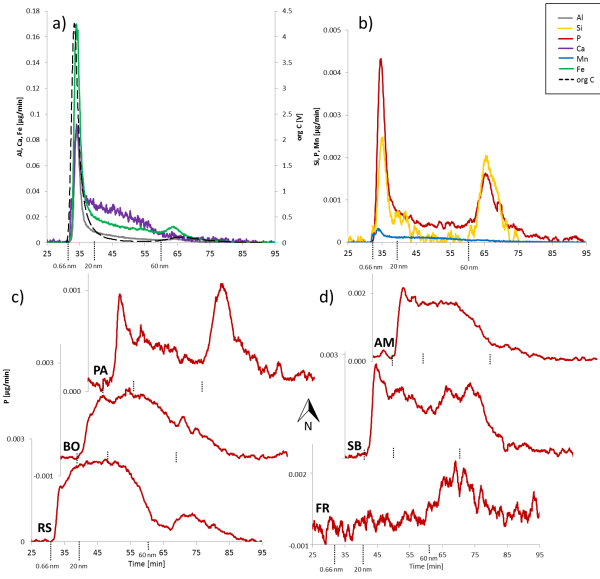
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867 Figure 1: Location of the 26 sampling sites in Europe along two transects (green dashed lines). Site 868 abbreviations are as follows: Pallas (PA) and Lettosuo (LE), Finland; Krycklan (KR), Norunda (NO), Gårdsjön 869 (GS) and Aneboda (AB), Sweden; Soroe (SR), Denmark; Allt a'Mharcaidh (AM) and Cotley Wood (CW), United 870 Kingdom; Wüstebach (WB) and Bode (BO), Germany; Lägeren (LÄ), Vogelbach (VB), Lümpenenbach (LÜ) and 871 Erlenbach (EB), Switzerland; Franchesiello (FR), Costiglione (CO) and Piano Rabelli (PR), Italy; Agia (AG), 872 Greece; Strengbach (SB) and La Peyne (LP), France; Ribera Salada (RS), Montseny (MS) and Baranco de 873 Porta Coeli (BPC), Spain; Sierra de Cima (SC) and Lourizela (LZ), Portugal. Black dotted lines indicate 874 geographical separation between northern (north of $56^{\circ}N$), middle (between $48^{\circ}N$ and $56^{\circ}N$) and southern 875 (south of 48°N) sites.

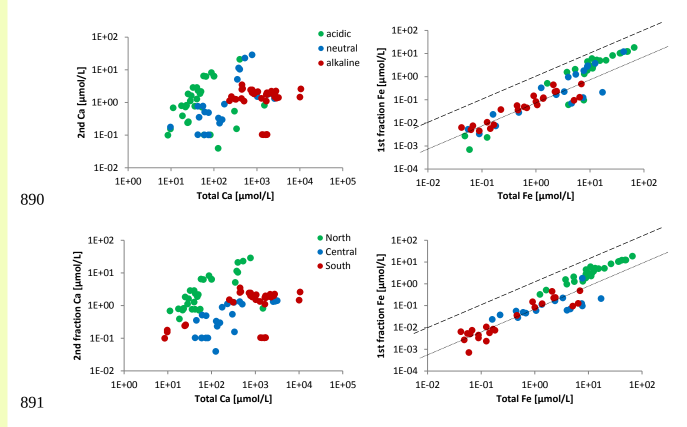
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878 879 Figure 2: AF^4 -ICP-MS and AF^4 -OCD raw data fractograms. a) Fractogram of Al, Ca, Fe and org C of one 880 sampling point at Krycklan, Sweden; b) Fractogram of P, Si and Mn of same sampling point as a) at Krycklan, 881 Sweden; c) Fractogram of P of three sampling points at sites in South, Middle and North Europe (increasing °N, 882 y-axis), North: PA = Pallas, Finland, Middle: BO = Bode, Germany, South: RS = Ribera Salada, Spain d) 883 Fractogram of P of three sampling points at sites in South, Middle and North Europe (increasing °N, y-axis), 884 North: AM = Allt a'Mharcaidh, Scotland, Middle: SB = Strengbach, France, South: FR = Franchesiello, Italy,. 885 X-axes represent the method time in minutes. Focus time was partially cut off. Y-axes for Al, Si, P, Ca, Mn and 886 Fe reflect mass flow in $\mu g/min$ and for org C detector signal in V. Fraction borders apply to the ICP-MS signal; 887 for the OCD evaluation these borders were modified because OCD peaks exhibit peak broadening due to the 888 high volume of the OCD reactor.

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892 Figure 3: Examples for the two relations of particle concentration as function of the total element 893 concentrations. Top left: Scatter diagram of log_{10} -transformed data of 2^{nd} fraction Ca, as an example of the 894 behavior of Si, P, Ca and Mn across all fractions; top right: Linear distribution of log₁₀-transformed data for 1st 895 fraction Fe, as an example of the behavior of Fe, Al and org C across all fractions; color coding represents pH 896 class of the site (cf. Table S1); acidic pH <6.6, neutral pH 6.6-7.3, alkaline pH >7.3. Bottom left and right: 897 Same data as top left and right, but with color coding according to geographic regions (cf. Figure 1). 2^{nd} 898 fraction Ca was chosen opposed to 1st fraction Ca for the left diagram because the 2nd fraction often exhibited a 899 Ca peak (cf. Figure 2a). Dashed line represents 1:1 line and dotted line average proportion of total Fe present 900 in the 1^{st} fraction. n = 96.

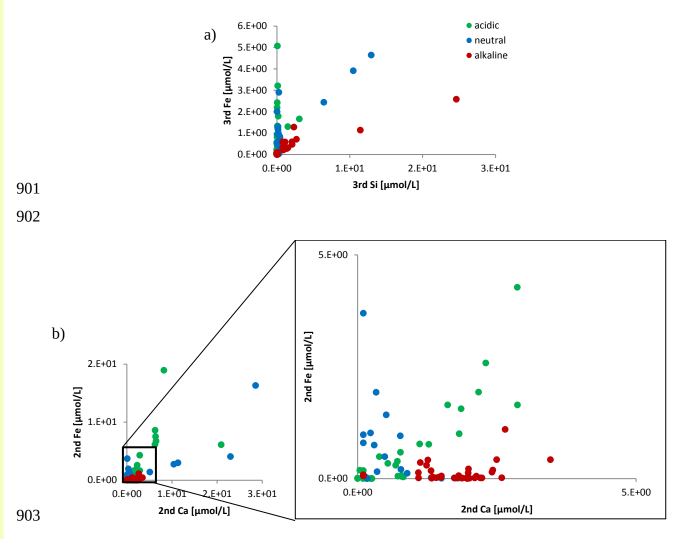


Figure 4: Example data plots of a) 3^{rd} fraction Fe over 3^{rd} fraction Si, and b) 2^{nd} fraction Fe over 2^{nd} fraction Ca with zoom window. Color coding according to site pH class, acidic pH = green dots (pH <6.6), neutral pH = blue dots (pH 6.6–7.3), alkaline pH = red dots (pH >7.3). n = 96.

904 Table 1: Characteristics of each European site. Climate data, bedrock and soil type and dominant forest type

- 905 provided by site operators. Abbr. = abbreviation, MAT = mean annual temperature [°C], MAP = mean annual
- 906 precipitation [mm], catch = catchment size [km^2], elev = average elevation [m], slope = average slope, forest =
- 907 proportion forest cover, MAR = mean annual runoff [mm].

site	abbr.	MAT	MAP	soil type	dominant forest	bedrock type	catch	elev	slope	forest	MAR
Aneboda	AB	5.8	750	Podzol	coniferous	Granite ^s	0.19	225	0.13	0.73	280
Agia	AG	15.8	691	Cambisol	broadleaf	Gneiss ^c	0.75	916	0.28		
Allt a' Mharcaidh	AM	5.8	1110	Podzol	coniferous	granite ^s	9.79	716		0.02	
Bode	BO	7.1	1600	Cambisol	coniferous	shale&greywacke	^s 1.27	515	0.07	1.00	
Barranco de Porta Coeli	BPC	14.5	450	Fluvisol	mixed	sandstone ^s	3.20	523	0.21	0.50	
Costiglione	CO	15.9	1183	Cambisol	broadleaf	Carbonatic ^c	11	563	0.32	1.00	225
Cotley Wood	CW	10.1	1044	Cambisol	broadleaf	siltstone	0.50	146	0.08	1.00	
Erlenbach	EB	6.0	2294	Gleysol	coniferous	flysch	0.73	1330	0.24	0.39	1778
Franceschiello	FR	15.9	1183	Cambisol	broadleaf	Carbonatic ^c	11	563	0.32	1.00	225
Gårdsjön	GS	6.7	1000	Podzol	coniferous	granite ^s	0.07	127	0.22	0.65	520
Krycklan	KR	1.8	614	Podzol	coniferous	greywacke ^s	679	260		0.87	311
Lettosuo	LE	4.6	627	Histosol	coniferous	gneiss	1.25	111	0.01	1.00	413
Lägeren	LÄ	8.4	930	Cambisol	broadleaf	Limestone ^c		680	0.35	1.00	
Montseny	MS	9.0	870	Inceptisol	coniferous	schist					
La Peyne	LP	12.0	818	Leptosols	broadleaf	schist ^s	110	305	0.25	1.00	
Lümpenenbach	LÜ	6.0	2426	Gleysol	coniferous	flysch	0.88	1260	0.15	0.19	2001
Lourizela	LZ	13.8	1300	Cambisol	coniferous	schist ^s	0.65	365	0.37	1.00	775
Norunda	NO	5.5	730	Regosol	coniferous	granite ^s	6084	45		1.00	250
Pallas	PA	-1.4	484	Podzol	coniferous	granite ^s	5.15	308	0.08	0.60	220
Piano Rabelli	PR	15.9	1183	Cambisol	broadleaf	Carbonatic ^c	11	563	0.32	1.00	225
Ribera Salada	RS	15.6	800	Cambisol	broadleaf	Carbonatic ^c		1483		1.00	
Soroe	SR	8.5	564	Mollisol	broadleaf	(glacial moraine) ^c		37	0.07	1.00	
Strengbach	SB	6.0	1400	Podzol	coniferous	granite ^s	0.80	1015	0.15	1.00	814
Serra de Cima	SC	13.8	1300	Cambisol	broadleaf	schist ^s	0.52	432	0.16	1.00	775
Vogelbach	VB	6.0	2159	Gleysol	coniferous	flysch	1.58	1285	0.23	0.63	1601
Wüstebach	WB	7.0	1220	Cambisol	coniferous	shales ^s	3.85	612	0.04	1.00	280

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^s: siliceous bedrock group; ^c : carlcareous bedrock group (see later Table 4)

909 Table 2: Fraction-specific median, slope and intercept values for predictability of elements (cf. Figure 3);

- 910 Si/P/Ca/Mn: fraction specific median concentrations represent midpoint of the data point distribution and
- 911 numbers in parentheses represent the percentage of e.g. 1st fraction Si as sum over all samples relative to all
- 912 particulate Si, Fe/Al/org C: linear regression slope (m) and intercepts (b) of log_{10} transformed data. n = 96;
- 913 unit: µmol/L, org C: mmol/L.

	Si P		P Ca Mn		Fe		Al		org C	
		m	b	т	b	т	b			
1st	0.03 (6.1)	0.04 (18.3)	0.67 (42.1)	0.001 (32.9)	1.00	-0.83	0.76	-0.55	1.71	-1.72
2nd	0.02 (3.1)	0.12 (53.2)	1.32 (49.5)	0.003 (37.3)	0.85	-0.47	0.65	-0.57	0.88	-1.55
3rd	0.08 (90.8)	0.08 (28.6)	0.24 (8.4)	0.002 (29.7)	0.72	-0.26	0.50	-0.07	0.56	-1.23
all part.	0.17	0.28	2.38	0.007						

914 Table 3: Number of sampling points in each pH class (acidic pH < 6.6, neutral pH 6.6-7.3, alkaline pH > 7.3) per $(1 + 1)^{-1}$

915 geographic region (Figure 1). n = 96.

	acidic	neutral	alkaline	total sampling points
North	24	7	0	31
Middle	5	10	8	23
South	8	2	32	42

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918 Table 4: Mean values of all particulate molar elemental concentrations (mmol/L for org C, µmol/L for Al, Si, P,

919 Ca, Mn) across classes representing groupings of site characteristics. Significantly different classes per category

920 and per element are marked by a, b and c; p < 0.05.

		org C	Al	Si	Р	Ca	Mn	Fe	number of sampling points
	dystrophic	0.44 a	0.59 a	0.36 a	0.32 a	6.14 a	0.03 a	3.72 a	45
soil class	eutrophic	0.03 b	0.04 b	1.12 b	0.21 a	1.80 a	0.03 b	0.54 b	34
	semi terrestrial	0.08 ab	0.05 ab	4.14 c	0.39 b	3.80 b	0.01 ab	1.04 ab	12
hadreat alars	siliceous	0.37 a	1.77 a	0.35 a	0.32 a	5.50 a	0.03 a	3.12 a	54
bedrock class	calcareous	0.03 b	0.15 b	0.08 b	0.24 a	2.06 ab	0.04 a	0.09 b	22
	flysch	0.08 b	3.36 c	4.96 c	0.30 a	3.89 b	0.01 a	1.24 a	10
dominant forest	coniferous	0.47 a	2.50 a	1.13 a	0.30 a	6.64 a	0.031 a	5.11 a	54
type	broadleaf	0.03 b	1.02 b	1.16 b	0.20 b	1.77 b	0.028 b	0.56 b	26
	acidic	0.59 a	2.46 a	0.25 a	0.24 a	5.65 a	0.03 a	6.79 a	31
pH class	neutral	0.39 a	1.71 ab	0.47 a	0.39 a	7.89 a	0.03 a	3.47 ab	24
	alkaline	0.04 b	1.62 b	2.11 a	0.29 a	2.47 a	0.03 a	0.76 b	41

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