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1	Are nanoplastics able to bind significant amount of metals?
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4 5 6 7	Mélanie Davranche ¹ , Cloé Veclin ¹ , Anne-Catherine Pierson-Wickmann ¹ , Hind El Hadri ² , Bruno Grassl ² , Laura Rowenczyk ³ , Aline Dia ¹ , Alexandra Ter Halle ³ , Florent Blancho ¹ , Stephanie Reynaud ² and Julien Gigault ¹
8	¹ Univ Repres CNRS Géosciences Repres UMR 6118 E35000 Repres France
9	² IPREM_LIMR 5254_CNRS-Liniversité de Pau et des Pays de l'Adour_E64000 Pau Erance
10	³ Laboratoire des Interactions Moléculaires et Réactivité Chimique et Photochimique (IMRCP), LIMR
10	CNDS 5622 Université Deul Schetier UDS, Detiment 201, 2 ^{ème} (tage 110, route de Norberge 21062
11	CNRS 5623, Universite Paul Sabatier-UPS, Batiment 2R1, 3 detage, 118, route de Narbonne, 31062
12	Toulouse Cedex 09, France
 13 14 15 16 17 18 19 20 21 22 23 24 25 26 27 28 29 30 	Corresponding author: Mélanie Davranche E mail: melanie.davranche@univ-rennes1.fr Tel: +33 223 235 769 Fax: +33 223 238 787 Main finding Nanoplastics are able to sorb high amount of Pb(II) through sorption and intraparticle diffusion processes. They could be important vector of metallic pollutant in the environment. Highlights • Pb(II) binding experiments onto nanoplastics extracted from environmental micro-plastics were performed. • Pb(II) was largely bound onto nanoplastics by specific adsorption and intraparticle diffusion. • Nanoplastics could be significant vectors of metals in the environment
31	Abstract
32	The nanoscale size of plastic debris makes them potential efficient vectors of many pollutants
33	and more especially of metals. In order to evaluate this ability, nanoplastics were produced from
34	microplastics collected on a beach exposed to the North Atlantic Gyre. The nanoplastics were
35	characterized using multi-dimensional methods: asymmetrical flow field flow fractionation and
36	dynamic light scattering coupled to several detectors. Lead(II) adsorption kinetics, isotherm and pH-

37 edge were then carried out. The sorption reached a steady state after around 200 min. The 38 maximum sorption capacity varied between 97% and 78.5 % for both tested Pb concentrations. 39 Lead(II) adsorption kinetics is controlled by chemical reactions with the nanoplastics surface and to a 40 lesser extent by intraparticle diffusion. Adsorption isotherm modeling using Freundlich model 41 demonstrated that NPG are strong adsorbents equivalent to hydrous ferric oxides such as ferrihydrite (log $K_{ads}^{freundlich}$ = 8.36 against 11.76 for NPG and ferrihydrite, respectively). The 42 43 adsorption is dependent upon pH, in response to the Pb(II) adsorption by the oxygenated binding 44 sites developed on account of the surface UV oxidation under environmental conditions. They could 45 be able to compete with Fe or humic colloids for Pb binding regards to their amount and specific 46 areas. Nanoplastics could therefore be efficient vectors of Pb and probably of many other metals as 47 well in the environment.

48 Keywords.

49 nanoplastics, Pb(II), adsorption, kinetics, DLS, A4F

50

51 Main finding

Nanoplastics are able to sorb high amount of Pb(II) through sorption and intraparticle diffusion
 processes. They could be significant vectors of metallic pollutants in the environment.

54

55 **1. Introduction**

56 Due to the exceptional properties of plastics materials such as their low density and high 57 durability, the industrial production of plastics has intensely and exponentially grown since 1940s. 58 The reason of their success is explained by their multiples uses (e.g. food packaging, building, 59 transport, agriculture, etc.). However, these properties result also in their exceptional persistence in 50 the environment. Whereas marine pollution by plastics was the most highlighted (low et al., 2010; 51 Eriksen et al., 2014), microplastic contamination of freshwater, urban water, urban dust, flood plain

62 soil and atmosphere were also pointed out in several studies (Gaspery et al., 2014; Dris et al. 2016;

63 Dehghani et al., 2017; Scheurer and Bigalke, 2018; Dris et al. 2018).

64 Among plastic contaminants, microplastics have been the most studied (Andrady, 2011; Wright et 65 al., 2013). However, the existence of nanoplastics, was quite recently demonstrated by Ter Halle et 66 al. (2017) and Gigault et al. (2018). They defined nanoplastics as particles unintentionally produced, 67 whose size was varying from 1 to 1000 nm and displaying a colloidal behavior (Gigault et al., 2018). 68 They are produced from the use of manufactured objects but especially from aged-microplastics in 69 response to the photo-, chemical and physical degradation mechanisms that take place under 70 environmental conditions (Bouwmeester et al., 2015; Ter Halle et al., 2017). Their occurrence was 71 demonstrated under laboratory conditions from degradation of plastics items (Lambert and Wagner, 72 2016), environmental microplastics collected in seawater (Gigault et al., 2018) and under 73 environmental conditions in seawater samples (Ter Halle et al., 2017).

74 Nanoplastics display a nanoscale size. They are polydisperse with an open structure and a 75 heterogeneous and asymmetrical shape. They present an inhomogeneous charged surface and they 76 are able to aggregate relative to the environmental physico-chemical conditions (Gigault et al., 2018). 77 All these properties make them formidable potential adsorbents of contaminants and especially of 78 trace metals. However, regards to their extremely recent highlighting, no studies have ever been 79 interested in their ability to bind trace metals and the involved governing mechanisms. Nevertheless, 80 several studies demonstrated that both plastics pellets collected from beaches and micro-plastics 81 recovered from seawater of the North Alantic gyre contain very variable amounts of trace metals 82 (e.g. Holmes et al., 2012; Turner and Holmes, 2015; Vedolin et al., 2018; Prunier et al., 2019). This 83 trace metal loading could result of the use of trace metals as additives in the plastics production and 84 also in sorption processes (Massos and Turner, 2017; Wang et al., 2017; Prunier et al. 2019). Wang et 85 al. (2017) suggested that the majority of metals carried by microplastics were derived from internal 86 load. However, Holmes et al. (2014) compared the sorption capacity of virgin pellets and pellets 87 collected from beaches. They provided evidence that trace metal adsorption was considerably

88 greater for the beach-sourced pellets. They attributed these results to the pellets weathering and to 89 the resulting specific adsorption onto charged mineral sites or to non-specific interaction between 90 the hydrophobic surface and the neutral or organic trace metal species. Adsorbed trace metals were 91 both cations and oxyanions. Ashton et al., (2010) suggested also that trace metals may co-precipitate 92 with or be sorbed onto hydrous oxides present onto the microplastic surface. Brennecke et al. (2016) 93 studied the sorption of Cu and Zn from antifouling paint onto virgin polystyrene (PS) pellets and 94 environmental aged polyvinyl chloride (PVC) fragments. They demonstrated that sorption of Cu was 95 higher onto aged PVC than virgin PS in response to the PVC higher surface aera and polarity.

In such context, the nano size of the nanoplastics and their exceptional surface properties (charge
and asymmetrical shape) make them better vectors of trace metals than pellets and microplastics.
Their precise role in trace metal transfer and the involved mechanisms have therefore to be
elucidated.

For this purpose, we studied the sorption of Pb(II) onto 'model' nanoplastics obtained from physical alteration of 'environmental' nanoplastics produced by sonication of microplastics collected on a beach exposed to the North Atlantics gyre in Guadeloupe (France). Lead was chosen regards to its high affinity for surface and to its toxicity. We developed an *in situ* polarography method allowing performing adsorption kinetics, isotherm and pH-sorption edge experiments in a single reactor. The evolution of the nanoplastic size and aggregation were monitored throughout the experiment using dynamic light scattering (DLS).

107 **2.** Materials and methods

All used chemicals were of analytical grade. The solutions were prepared with ultrapure water (Milli-Q system, Millipore). The containers used were (i) trace-metal cleaned with 10% (v/v) HNO₃ for 24 h at 45°C, (ii) soaked in and then rinsed with ultrapure water for 24 h at 45°C, and (iii) finally dried at 30°C. All solutions were prepared with ultrapure water (18.2 Ω , MilliQ, Millipore). The Pb(II) solutions were prepared using a 1000 mg L⁻¹ standard solution of Pb (NO₃)₂ (standard Metrohm

99.5% ion, VWR Chemicals). The analysis of Pb²⁺ in solution was carried out by polarography
(Metrohm 884 professional voltammetric meter).

115 2.1. Nanoplastics production

116 Environmental nanoplastics were produced by the sonication of 58 g of microplastics (5 mm to 2 117 cm) in 300 mL of ultrapure water. Sonication was performed during 5 days. The suspension was 118 filtrated at 0.8 µm (Sartorius). Microplastics were collected in 2017 on the beach of Baie Sainte-Marie 119 in Guadeloupe (France), which is exposed to the North Atlantic gyre. This sample corresponding to 120 Guadeloupean nanoplastics was called NPG. Microplastics were isolated from the sand by a sea 121 water extraction method relative to their density. Dynamic light scattering (DLS) displayed that both 122 samples were poly-dispersed with an average hydrodynamic diameter 150-450 nm. Zeta-potential 123 analysis demonstrated that they are negatively charged on the 2-to-14 pH range. At pH 6, zeta-124 potential was equal to -30.2 mV. Their final concentration in the suspension was determined by 125 measuring the dissolved organic carbon (DOC). The NPG sample displayed a DOC concentration of 50 mg L⁻¹. Note that it cannot be excluded that a few amount of the measured DOC corresponds to 126 127 organic matter.

128 2.2. Adsorption experiments

129 Kinetic experiments. Kinetic experiments of Pb(II) adsorption onto NPG were performed in situ, 130 namely directly in a polarographic cell. This in situ methodology avoids any external treatment of the 131 sample, Pb²⁺ concentration being continuously measured in the cell. Ten ml of the NPG solution were 132 introduced in a polarographic cell. The pH of the NPG suspensions was fixed at 7 using HNO₃ and/or NaOH at 0.1 M and their ionic strength was fixed at 10⁻² mol L⁻¹ using KCl. Before adding Pb(II), a first 133 polarographic measurement was performed to check (if any) the Pb²⁺ concentration release by NPG. 134 Then, Pb(II) was added to obtain a concentration of 2 and 6 mg g⁻¹ (g of dissolved organic carbon, 135 DOC). The concentration of Pb^{2+} in the suspension was measured every 2 min during 5 h. 136

137

Adsorption isotherms. They were carried out in situ directly in a polarographic cell. Ten ml of 138 the NPG solution was introduced in a polarographic cell. The pH of the NPG suspensions was fixed at 7 using HNO₃ and/or NaOH at 0.1 M. The ionic strength was fixed at 10^{-2} mol L⁻¹ using KCl. Before 139 140 adding Pb(II), a first polarographic measurement was performed to check (if any) the initial Pb²⁺ 141 concentration. Then, Pb(II) was added every 4h (as assessed by the steady state) in order to obtain Pb(II) total concentration of 4, 8, 12, 16, 20 and 24 mg g^{-1} (g of DOC). The Pb²⁺ concentration was 142 143 measured by polarography.

pH-adsorption edge experiment. This experiment was carried out in situ directly in a 144 145 polarographic cell and not as classically performed in distinct batches. Fifteen mL of NPG solution 146 were introduced in a polarographic cell. Its initial pH was fixed at 2 using HNO₃ at 0.1 M. The ionic 147 strength was fixed at 10⁻² mol L⁻¹ using KCl. Lead(II) was added in order to obtain a concentration of 6 mg g^{-1} (g of DOC). After 4h, the Pb²⁺ concentration was measured by polarography. Then, every 4 h, 148 149 the pH was increased using an automatic titrator in SET mode (Titrino 719, Metrohm). The pH was successively fixed at 3, 4, 5, 6 and 9. The Pb²⁺ concentration was measured at steady state for each 150 151 pH.

152 2.3. Chemical Analysis

153 Dissolved organic carbon (DOC). DOC concentrations were determined using a Total Carbon 154 Analyzer (Shimadzu TOC-V CSH) by a standard solution of potassium hydrogen phthalate (Sigma 155 Aldrich). The uncertainty of the measurement was 5%.

Pb(II) measurement. Pb(II) concentrations were determined as Pb²⁺ species concentrations 156 157 using polarography directly in the NPG suspensions. They were determined using a Methrohm 158 polarogaph (Methrom 884 professional ASV) under Anodic Stripping Voltammetry (ASV) mode using 159 a hanging dropping mercury electrode (HDME). The analytical conditions are summarized in Table S1 160 (supplementary file). Deoxygenation of the medium was carried out using a continuous N₂ bubbling. 161 The analyses were carried out under stirring at 3000 rpm.

6

162 All experiments being performed directly in a polarographic cell, the metered addition method could not be carried out. A calibration curve was therefore performed for a Pb²⁺ concentration 163 ranging in between 0 and 400 μ g L⁻¹. To evaluate the impact of the NPG presence on the 164 165 measurement, calibration curve were carried out with and without NPG. Moreover, to avoid any Pb(II) sorption onto NPG and ensure that all the Pb(II) was under Pb^{2+} species, the pH was fixed at 2. 166 167 None influence of the NPG presence was observed. The calibration curve realized with NPG at pH 2 was therefore used to determine the experimental Pb^{2+} concentration (supplementary file Fig. S1). 168 The detection limit of the method was 0.05 μ g L⁻¹ for Pb. 169

170 *2.4. Nanoplastics size measurement.*

171 Dynamic Light Scattering (DLS). DLS was used to determine the size of the nanoplastics both in 172 the initial suspension and during the adsorption experiments. The in situ DLS measurements were 173 performed using a Vasco Flex model of nanoparticle size analyzer (Cordouan Technology, Pessac, 174 France). The probe was placed in front of the polarographic cell and measured directly in the 175 solution. The detection limit of the DLS instrument was investigated with PSL 100 nm standards within the range of 2×10^{-5} g L⁻¹ to 2×10^{-1} g L⁻¹. Although detection remains possible over the entire 176 concentration range, an average size analysis is only possible over the concentration range of 2×10^{-2} 177 to 2×10^{-4} g L⁻¹ using the Pade-Laplace algorithm. Each measurement corresponds to a statistical 178 179 average of six measurements. In addition, each of the six measurements is composed of six 180 acquisitions of light scattered for 80 s.

Asymmetrical Flow Field Flow Fractionation (A4F) coupled to static light scattering (SLS) and inductively coupled plasma-mass spectrometry (ICP-MS). A4F-SLS-ICP-MS was used to fractionate and size sort nanoplastics with metallic composition associated to. The A4F fractionation running conditions are summarized in table S2 (supplementary file).

185 The detection consisted in a static light scattering (SLS, Multi Angle Light Scattering - MALS, Dawn 186 Heleos 2, Wyatt Technology) to characterize the particles size. The incident laser wavelength of the 187 SLS is λ =658nm. A Berry formalism that allows giving the more suitable fit of the light scattered

7

188 according to the angle was used. UV detector was fixed at 254 nm, which corresponds to the 189 absorption peak of the colloidal materials. Finally, an inductively coupled plasma-mass spectrometer 190 (ICP-MS 7700x, Agilent technologies) was directly coupled at the end of the detection chain without 191 using the nebulizer pump. No online quantification was performed. Here ICP-MS was used to localize 192 the Pb(II) on the colloidal size range. The ICP-MS was operated without the collision cell, because of the lack of interferences for the mass isotope of interest (²⁰⁷Pb). The sample and skimmer cones were 193 194 in nickel. The RF power and the plasma gas flow rate were fixed at 1500 W and 15 L min⁻¹, 195 respectively.

196 **3. Results**

197 *3.1. Adsorption kinetics*

Lead concentration decreased extremely rapidly (Fig. 1a). After 2 min, the concentration decreased from 2 to 0.43 mg g⁻¹ (78.5 % of binding) and 6 to 0.17 mg g⁻¹ (97.2 % of binding) for both concentrations, respectively. Lead was therefore significantly adsorbed to NPG. The kinetic profile consisted of a rapid adsorption following by a subsequent approach to steady state, which was reached after around 200 min (Fig. 1a).

The kinetics was modeled using a pseudo-first-order. However, the Lagergreen pseudo-firstorder equation was not applicable to our Pb(II)-NPG system, no linearization of the experimental data was obtained. The pseudo-second order was then tested. In this model, the rate-limiting step is the surface adsorption that involves chemisorption. The model is based on the solid phase adsorption capacity (Ho et al. 1996 a-b) and predicts the behavior over the whole range of adsorption (Ahsu et al. 1991). The corresponding equation is:

209
$$\frac{\mathrm{d}Q_{\mathrm{t}}}{\mathrm{d}t} = \mathrm{K} \left(\mathrm{Q}_{\mathrm{e}} - \mathrm{Q}_{\mathrm{t}} \right)^2 \qquad (\mathrm{eq.1})$$

With Q_t , the adsorbed Pb(II) concentration at time t (mg g⁻¹ of DOC), Q_e the adsorbed Pb(II) concentration at steady state (mg g⁻¹ of DOC), and K the rate constant of adsorption for the pseudosecond-order (g mg⁻¹ min⁻¹).

213 The integrated equation is:

214
$$\frac{1}{Q_t} = \left(\frac{1}{Q_e}\right) \cdot t + \frac{1}{K \cdot Q_e^2}$$
 or $\frac{1}{Q_t} = \frac{h}{1 + K \cdot Q_{e,t}}$ (eq.2)

215 Where h can be considered as the initial sorption rate. When t tends to 0, $h = KQ_e^2$

The linearity of the plots, the R² and the coherence between the theoretical and the experimental Qe indicated that chemical reaction is the main rate-controlling step of the binding process (Fig. 1b and Table 1). The rate constant (K) decreased with the increasing Pb(II) initial concentration as previously observed and discussed by Ho and Ofomaja (2006) and Ho et al. (2006) for Pb(II) adsorption onto palm fiber and peat.

Since the pseudo-second-order kinetic model did not allow identifying potential diffusion mechanism, the kinetic results were then analyzed using an intraparticle diffusion model. The following equation was used (El-Ashtoukhy et al., 2008):

224
$$Q_t = K_i t^{0.5} + C$$
 (eq. 3)

 K_i is the rate constant of the intraparticle diffusion, whereas C corresponds to the intercept and provides information on the thickness of the boundary layer.

The Qt was plotted relative to the \sqrt{t} in Fig. 1c and the kinetics parameters in Table 3. If the 227 228 intraparticle diffusion model is involved the plot of Qt relative to \sqrt{t} should be linear and if the 229 intraparticle diffusion model is the rate-controlling step, the plot should pass through the origin (C = 230 0). However, if C \neq 0, it indicates some degree of boundary layer control. The intraparticle diffusion 231 model is therefore not the rate-controlling step, but it could operate simultaneously with other 232 processes. The diffusion gradient can be linearized into four regions representing the external mass 233 transfer, and the intraparticle diffusion in the macro, meso and micropore structure of the adsorbent 234 (Allen et al., 1989; Tan et al. 2007; Ho and McKay, 1998). Multilinear lines were obtained for both 235 initial concentrations of Pb(II) (Fig. 1c). Moreover, any of them passed through the origin indicating 236 that the intraparticle diffusion process was not the rate-controlling step. However, the first linearity 237 can be considered as a first step of an instantaneous adsorption or adsorption on the outer surface 238 which was produced during the first 25 min of the sorption and corresponded to 83% and 89 % of

total bound Pb(II) for 6 and 2 mg g⁻¹ initial concentration, respectively. The second part represented a more slow and progressive part of the adsorption that could be driven by an intraparticle diffusion process that limited the adsorption process and corresponded to 4% and 5% of total bound Pb(II) for 6 and 2 mg g⁻¹ initial concentration, respectively.

Therefore, Pb(II) seemed to be rapidly sorbed by a chemisorption processes which was confirmed by the pseudo-second order model parameter, that may be followed by an intraparticle diffusion process lasting up to 25 min.

246 *3.2.* Adsorption isotherm

Experimental data of the adsorption isotherm are reported in Fig. 2. The adsorbed Pb(II) concentration increased with the increasing initial Pb(II) concentration. However, the proportion of adsorbed Pb(II) was higher for the low concentration than for the high concentration, suggesting a progressive saturation of the surface.

The isotherm was fitted using the non-linear Langmuir and Freundlich models. Since no linearity was obtained for Langmuir model, Freundlich model was therefore the only model described in this work. The Freundlich isotherm assumes a multilayer sorption onto a heterogeneous surface. The Freundlich isotherm corresponds to:

255

$$Q_e = K_{ads} C_e^n \qquad (eq. 3)$$

256 Where Qe is the adsorbed Pb(II) concentration (mg g^{-1}), K_{ads} is the Freundlich adsorption constant, C_e 257 is the concentration of Pb(II) in solution (mg g^{-1}) and n is the non-linearity coefficient.

In Fig. 2 were plotted the linearized data. Values for the Freundlich constant and coefficient are derived from the linear regression analysis of the log Qe versus log Ce. The obtained values are n= 0.34, Kads = 4.89. The correlation coefficient R² equal to 0.98 indicates that the Freundlich model is suitable to describe the sorption processes between Pb(II) and NPG. The coefficient n, ranging between 0 and 1, is indicative of a cooperative adsorption (here n=0.38).

These results showed that Pb(II) adsorption onto NPG was expected to be non-linear, namely produced through the formation of multilayer or through the formation of heterogeneous surface complexes. The surface complexes heterogeneity is the result of the variability of the binding sites (carboxylic, phenolic, hydroxyl, etc.) or of the denticity of the complexes (monodentates, bidentate, etc.).

268 3.3. pH adsorption-edge

The evolution of the Pb(II) adsorption relative to pH was non-conventionally plotted (Fig. 3). Since the experiments were both *in situ* and non-sequentially performed, it was not possible to evaluate, as done with the previous experiment, the Pb(II) concentration released by NPG at each pH. We therefore chose to present the evolution of the Pb²⁺ concentration (Fig. 3).

A Pb²⁺ concentration decrease following the increasing pH is displayed in Fig. 3. Several 273 mechanisms could explain such result and notably the complexation of Pb(II) by the OH- ligand, 274 275 whose concentration increased with the increasing pH up to the precipitation of Pb(II) as hydroxide at pH > 7. The Pb(II) speciation was calculated through VisualMinteq 3.1 considering the 276 277 experimental conditions but, without NPG (Fig. 4). To allow a better comparison between the 278 experimental and the modeled data, the Pb(II) concentration was presented in mg g⁻¹ (g of DOC even if no DOC was input in the calculation, namely to convert mol L^{-1} as mg g^{-1}). Without any NPG, the 279 modeling demonstrated that Pb²⁺ decreased from pH 6 in response to its complexation as PbOH⁺ and 280 281 from pH 7.5 to the precipitation of Pb(OH)₂s. By contrast, with NPG, Pb²⁺ decreased from pH>2, the largest decrease proceeding from pH 4. These results clearly demonstrated that the Pb²⁺ decrease 282 was provided in response to the NPG occurrence, namely in response to the Pb²⁺ adsorption onto 283 284 NPG.

285 3.4. Size and elemental composition characterization

286

DLS analysis was performed on NPG associated to Pb(II) on the studied pH range, concentrations
 and contact-time(kinetics) (Fig. S2, supplementary file). No significant changes were observed in the

289 auto-correlation function (ACF) and in the d_{zH} according to the different conditions. Nevertheless, 290 some difference on the polydispersity could have occurred and could not be identified by DLS. The 291 structural and physical changes of environmental nanoparticles are usually difficult to identify with 292 DLS due to their high polydispersity in size and composition (Gigault et al., 2018). To isolate, 293 fractionate and characterize, free-Pb(II) and Pb(II)-doped NPGs, A4F-SLS-ICPMS was further used (Fig. 294 5). The fractograms displayed a signal with a retention time covering the size range from 50 nm up to 295 400 nm (in diameter) based on A4F calibration (Gigault et al., 2017) (Fig. 5.a). The intensity of the UV 296 signal was slightly higher for Pb(II)-doped NPG than for free-Pb(II) NPG. Such results might be 297 produced in response to the Pb(II) adsorbed onto the NPG surface or to the NPG dispersion in 298 presence of Pb(II). The retention time was not modified indicating no modification of the size 299 distribution range. However, the UV signals showed a large colloidal fraction (from 10kDa to 60kDa) 300 at a retention time $t_R=2$ min. Residues of polymers, additives and organic matter remaining on the 301 microplastics surface, possibly released during the sonication. The SLS analysis validated these 302 observations. They displayed that NPG have a gyration radius ranging from 70 to 250 nm using Berry 303 formalism for both Pb(II)-doped and free-Pb(II) NPG. Angular dependence of the light scattered did 304 not fit using the spherical formalism indicating an indirect shape deviating from the ideal sphere (SI, 305 Fig. S2. The on-line coupling of ICPMS and A4F-UV-SLS allow obtaining the elemental composition 306 associated to the UV and SLS signals (Fig. 5b). The m/z 207 trace, corresponding to Pb(II) increased 307 from t_R=5min to t_R=17min for Pb(II)-doped NPG (black trace) as compared to free-Pb(II) NPG (blue 308 trace) indicating that Pb(II) was associated to NPG. Noted that for free-Pb(II) NPG, the m/z 207 trace 309 presented some spikes, starting especially at t_{R} =12min, namely at approximately 100 nm size. Such 310 spiked signal corresponds to the signal generally observed with ICP-MS in single particle mode. When 311 a significant amount is localized in a nanometer resolved surface, the quantity of ionized metal is 312 significantly higher, which corresponds to either metal nanoparticles trapped inside colloids or metal 313 accumulation within a cavity of a porous material. Nevertheless, for the Pb(II)-doped NPG, the spiked

- 314 signal was attenuated suggesting either, a Pb(II) surface adsorption or ,that the equilibrium time was
 315 long enough to allow a significant Pb(II) accumulation in the porosity.
- 316 **4.** Discussion
- 317 4.1. NPG is a significant Pb(II) adsorbent

318 Adsorption kinetic experiments displayed that 78.5 and 97 % of Pb(II) were sorbed onto the 319 nanoplastics at pH 7. These proportions are high and suggest that nanoplastics are strong Pb(II) 320 adsorbent. The Freundlich K_{ads} defined for Pb(II)-NPG in the present study was compared to the 321 Freundlich K_{ads} of various adsorbents. For the comparison, it was necessary to recalculate our 322 constants using the same calculation units than those of the literature. They are reported in Table 2. 323 Since in the literature, no data exist for the Pb(II) binding to nanoplastics, our results were therefore 324 compared to those obtained for plastics pellets collected on beaches on which Pb(II) was 325 experimentally adsorbed using sea and river water samples (Holmes et al., 2012; 2014). Note that the 326 beached pellets are micrometric and that they were submitted to sonication during 5 min before 327 being used. Although pH values were close to each other, the present Freundlich K_{ads} is 3 to 4 order 328 of magnitude higher than those of beached pellets (Table 2). The environmental nanoplastics (NPG) 329 are therefore stronger Pb(II) adsorbent than environmental micrometrics plastics. Note however, 330 that the sonication step submitted to the beached pellets could have released many nanometric 331 parts decreasing thus their binding capacity. Regards to these results, the present Freundlich K_{ads} was 332 compared to the K_{ads} calculated for Pb(II) binding to minerals known to be strong metals adsorbents, 333 namely the Fe oxyhydroxides. First, K_{ads} was compared to those calculated for Pb(II)-goethite 334 sorption. Goethite is a well-crystallized Fe oxyhydroxides with a low specific surface around 45 g m⁻². 335 The present Freundlich K_{ads} was 1 or 2 orders of magnitude higher than those of goethite (Table 2). 336 However, the binding experiments of Pb(II) to goethite were performed at lower pH. The metal 337 sorption onto Fe oxyhydroxides being strongly dependent to pH, the difference noted in between the 338 Freundlich K_{ads} could be involved by this pH difference, see notably the Olu-Owolabi and Ajayi (2011)

339 study where pH was 4.2. By contrast, Freundlich K_{ads} of Pb(II)-ferrihydrite sorption is in the same 340 order of magnitude than the present K_{ads}. Ferrihydrite is a Fe oxyhydroxide with a high specific surface varying between 250 to 600 g m^{-2} and is considered to be the strongest metal sorbent 341 342 amongst various particulate Fe oxyhydroxides. The similarity of the ferrihydrite and the present 343 Freundlich K_{ads} therefore suggests that NPG are strong adsorbent of Pb(II). Finally, the Freundlich K_{ads} 344 was compared to that obtained for nano-goethite, to eventually identify the role of the nano-size on 345 the sorption. Both Freundlich K_{ads} are in the same order of magnitude although they were highly 346 different for particulate goethite. Finally, our data were compared to that obtained for a purified 347 humic acid (PPHA). The Freundlich K_{ads} was calculated from data extracted from the Pb-humic acid 348 data compilation performed by Milne et al., (2001). Freundlich isotherm was not able to model Pb-349 humic acid binding data except that of Pinheiro et al. (1994), the Langmuir isotherm providing better modeling. Regarding the pH conditions, we can consider that both Freundlich K_{ads} are in the same 350 351 order of magnitude.

All these comparisons demonstrated that NPG are strong adsorbent of Pb(II), much stronger than beached micrometric plastics pellets. Their sorption capacity seems to be similar to strong metal sorbent such as ferrihydrite, nano-goethite and humic acid. Their sorption properties can thus be attributed to their nano-size and subsequent high specific surface area such as those of ferrihydrite, nano-goethite and humic acid.

357 4.2. Mechanisms involved in Pb(II) adsorption

358 We demonstrated that NPG is a strong adsorbent of Pb(II). Let us go further to elucidate which 359 are the mechanisms involved in the Pb(II) binding to NPG.

The first interesting results are provided by the adsorption kinetic study. The kinetic rate could not be modeled using the pseudo first-order model demonstrating that the adsorption kinetics was not controlled by the diffusion. By contrast, the pseudo-second order model allowed with success to model the kinetic data, indicating that the major rate limiting step was the surface adsorption involved by chemical interactions in between Pb(II) and NPG. Although this process dominates the

adsorption, intraparticle diffusion seems also to partly control the mechanism of Pb(II) adsorptiononto NPG.

367 Lead(II) sorption onto NPG was strongly dependent to pH as shown by the pH-adsorption-edge 368 experiment (Fig. 3). With the increasing pH, the surface electronegativity increased in response to 369 the surface functional groups deprotonation. In such conditions, Pb(II) affinity for the surface 370 increased, Pb(II) being able to be bound through electrostatic interactions (outer sphere complex) 371 and/or covalent binding (inner sphere complex). The formation of inner spheres supposes the 372 presence of surface sites able to exchange electron such as, hydroxyl, carboxylic and phenolic functional sites. Figure 3 shows that Pb^{2+} concentration strongly decreased from pH > 4 (i.e. Pb(II) 373 374 adsorption strongly increased). This pH corresponds to pH from which carboxylic groups generally 375 become significantly deprotonated (pKa $_{R-COOH} \approx 4.8$). If no data are available for nanoplastics, several 376 studies showed that plastics debris collected on beach or seawater displayed a higher oxidation state 377 than the reference sample (Ter Halle et al., 2017). Fourier Transform Infra-red Spectroscopy (FTIR) of 378 weathered plastics showed that oxygenated moieties such as, aldehydes, ketones, alcohol, 379 hydroperoxides and carboxylic acids groups increase in the plastics with the duration exposure 380 (Andrady, 2017). The increase of the oxygen moieties and oxygenated function is involved by the UV 381 irradiations weathering. The modeling of the adsorption isotherm data with the Freundlich model 382 demonstrated that the adsorption was not only favorable but, heterogeneous in response to either a 383 multilayer sorption or the formation of heterogeneous surface complexes (surface sites and complex 384 denticity heterogeneity). Since Pb(II) is a transition metal whose potential to form covalent binding is 385 high, the multilayer binding is therefore unlikely. However, regarding the oxidation state of the NPG 386 surface, the formation of heterogeneous complexes with various sites such as, carboxylic, hydroxyl or 387 phenolic sites is more than expected. Note, however, that Pb(II) sorption onto NPG surface does not 388 affect the aggregation/dispersion of the NPG, since no difference in their size and Rg variation was 389 observed with or without Pb(II).

390 As previously explained, fractograms of free Pb(II) and Pb(II)-doped NPG obtained with A4F-391 ICPMS, exhibited spikes in the Rg range over 100 nm. Passing 100 nm in Rg, the frequency of high-392 intensity spikes increases. Nevertheless, the present A4F-ICPMS set up is not optimized to 393 characterize nanoparticle size using the single particle-ICPMS mode (Schwertfeger et al., 2016; 394 Bustos and Winchester, 2016). Dwell and settling times are not indeed sufficient to obtain a robust 395 correlation between the intensity of the spike and the exact size distribution. These spikes and 396 single-particle event in the fractograms could be explained by the absorption and accumulation of 397 Pb(II) in the NPG structure/porosity, confirming the intraparticle diffusion of Pb(II). Note that the 398 Pb(II) occurring in the porosity and in the residual structure of the free-Pb(II) NPG are strongly 399 trapped since very fewly released at acidic pH as shown by the polarographic measurements. The 400 number and intensity of spikes was higher for free-Pb(II) than for Pb(II)-doped NPG in response to 401 the Pb(II) adsorption in high proportion onto the NPG surface that partly hid the spike signal. Such 402 result confirms the large surface adsorption of Pb(II) to NPG as previously evidenced by experimental 403 data and modeling.

404 All this results demonstrated that Pb(II) is adsorbed on the NPG in high amount through its 405 binding as surface complexes with the oxygenated binding sites produced by the UV irradiation 406 under environmental conditions. This major process is, however, coupled to a physical 407 trapping/concentration of Pb(II) in the structure/porosity of the largest NPG in response to an 408 intraparticle diffusion mechanism.

409 4.3. Is this sorption mechanism environmentally significant?

The present results clearly demonstrate that nanoplastics have strong affinity for metals. The question becomes then to identify whether this affinity could impact significantly the metal environmental fate. The answer depends on two major parameters: (i) the nanoplastics environmental amount and (ii) their ability to compete with the other significant metal ligands such as, Fe(III) nano-oxides and/or organic matter.

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415 No thorough quantitative information exists on the true amount of nanoplastics occurring in the 416 environment. However, hypotheses regarding their presence within environment allow some 417 estimations to be performed as follows. Although across the oceans, the total mass of microplastics 418 was estimated, using circulation models, to vary in between 9.3 to 23.6 10³ tons, only 1% of this 419 estimated amount is actually found in the ocean (Jambeck et al., 2015). Several authors suggested 420 that this difference could be explained by the alteration of the micro-plastics as nanoplastics in 421 response to the ultraviolet radiation, physical wave forces and hydrolysis in ocean (Andrady et al., 422 2011; Gigault et al., 2016; Ter Halle et al. 2017). If nanoplastics would be equally distributed in the 423 whole ocean water (liquid water), their hypothetic concentration would vary from 7 to 18 pg L^{-1} , 424 which is rather low. However, accumulation zones of microplastics have been identified in several 425 regions such as, Artic water or oceanic gyres where microplastics can exceed 6x10⁵ pieces km⁻², which represent for the average depth of oceans 2.3 $\times 10^{18}$ pieces L⁻¹ (Law et al., 2010; Bergman et 426 427 al., 2016, Peeken et al. 2018). Considering that these microplastics represent 1% of the plastics present in ocean, the nanoplastics amount in such samples can potentially reach 2.3 10^{20} pieces L⁻¹ 428 429 which is rather high. In soils, the quantitative occurrence of microplastics and nanoplastics are 430 considered as scarce and inexistant, respectively (Rillig et al., 2012). However, the plastic particle 431 loading in agroecosystems should be high due to inputs of both recycled organic wastes and plastic 432 film mulching (Ng et al., 2018). Therefore, in some specific worldwide microplastic accumulation 433 areas, nanoplastics concentrations could be significant making them alternative actors to be 434 considered in the metal cycles and accumulation.

Concerning their ability to compete with strong soluble or solid ligands for metal binding a first answer can be found in Table 1. The Freundlich K_{ads} compilation indeed demonstrates that for equivalent mass, the Freundlich K_{ads} being in the same order of magnitude, nanoplastics could be potential competitors for Fe nano-oxides and humic substances. However, this competition will depend on the specific area and the binding sites density between the competitive ligands and the nanoplastics. Finally, the key points to be considered before starting regarding such competition are

the interactions that could be performed between these ligands and the nanoplastics (aggregation, coating, repulsion, etc.). These interactions could, indeed, increase the potential of nanoplastics to bind metals (through additive effects) or by contrast limit their sorption capacities by recovering their surface limiting as a consequence their binding sites availability.

445 **5.** Conclusion

446 Nanoplastics, recently identified and characterized, are emerging pollutants. Their size makes them potential vectors of other pollutants such as metals. This study provided evidence that 447 448 nanoplastics, formed in natural conditions under which they are mechanically degraded and 449 especially oxidized by UV irradiations, are able to adsorb Pb(II). Lead(II) adsorption seems to be 450 controlled by chemical and physical interactions with NPG. Adsorption is strongly dependent to pH. 451 A large increase in the adsorption was evidenced for pH > 4 that corresponds to the carboxylic sites 452 deprotonation. This observation suggests that Pb(II) could be bound to the surface oxygenated 453 binding sites through inner sphere complexes, which result in an heterogeneous adsorption process. 454 To a lesser extent, sorption seems to depend on a Pb(II) intraparticle diffusion as evidenced by 455 modeling and analytical results.

456 The sorption capacity of nanoplastics are high, with Pb(II) Freundlich binding constants close to 457 those of ferrihydrite, nano-goethite and humic acid, known to be strong adsorbents of metals. If no 458 evidence were provided that microplastics are able to load high amount of metals, by contrast the 459 present study demonstrated that the nanometric size and surface properties of nanoplastics 460 produced under environmental conditions, can allow nanoplastics to be efficient vectors of Pb(II) and 461 probably of many other metals. Moreover, both modalities of Pb binding to NPG raise the question 462 of the subsequent bioavailability of the bound metals. Are adsorbed Pb(II) more available than 463 porosity trapped Pb(II) or inversely?

464

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- 468

469 **References**

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Table 1:	Kinetic parameters calculated using the pseudo second order model and the intraparticle diffusion	
	model for the kinetic adsorption experiment at $[Pb(II)] = 2 \text{ mg g}^{-1}$ and $[Pb(II)] = 6 \text{ mg g}^{-1}$.	

Kinetic parameters	Kinetic experiment	Kinetic experiment		
	at [Pb(II)] = 2 mg g ⁻¹	at [Pb(II)] = 6 mg g ⁻¹		
	Pseudo-second order model			
Qe theorical (mg g ⁻¹)	1.89	5.44		
K (g mg ⁻¹ min ⁻¹)	0.24	0.072		
h (mg g ⁻¹ min ⁻¹)	0.88	2.14		
R ²	1	0.999		
Qe _{experimental} (mg g ⁻¹)	1.89	5.46		
1 1/2	Intra-particle diffusion			
$Ki_1 (mg g^{-1} min^{-1/2})$	0.07	0.16		
$C_1 (g mg^{-1} min^{-1})$	1.5	4.2		
$R_{1}^{2}(mg g^{-1} min^{-1})$	0.98	0.98		
$Ki_2 (mg g^{-1} min^{-1/2})$	0.008	0.023		
$C_2 (g mg^{-1} min^{-1})$	1.76	5.06		
$R_{2}^{2}(mg g^{-1} min^{-1})$	0.87	0.78		

Sample	K _{ads}	K _{ads} (present study)	Calculation units	References
Beached pellet in sea water pH= 7.8	2.1 x 10 ⁻¹	4.6 x 10 ²	y: μg g⁻¹ x: μg L⁻¹	Holmes et al. (2012)
Beached pellets in sea water pH= 7.8	6.5 x 10 ⁻¹	3. 5 x 10 ³	y: nmol g ⁻¹ x: nmol L ⁻¹	Holmes et al. (2014)
Beached pellets in river water pH= 6.3	3.7 x 10 ⁻¹	3. 5 x 10 ³	y: nmol g ⁻¹ x: nmol L ⁻¹	Holmes et al. (2014)
Goethite pH= 6, IS = 10 ⁻³ M	3 x 10 ⁻⁴	4.5 x 10 ⁻³	y: mol g⁻¹ x: mol L⁻¹	Christophi and Axe (2000)
Goethite pH = 4.2	1.710 ⁻¹	1.4 x 10 ¹	y: mmol kg ⁻¹ x: μmol L ⁻¹	Olu-Owolabi and Ajayi (2011)
Ferrihydrite pH = 5.5	11.76	8.31	y: mg g ⁻¹ x: mg L ⁻¹	Rout et al. (2012)
Nano-goethite pH = 5.5	10.33	8.31	y: mg g ⁻¹ x: mg L ⁻¹	Mohapatra et al. (2010)
Purified Humic Acid (PPHA) pH = 5	1.40	8.31	y: mg g ⁻¹ x: mg L ⁻¹	Pinheiro et al. (1994)

Table 2: Comparison of the present calculated Freundlich K_{ads} and the Freundlich K_{ads} of Pb(II) binding to various adsorbents.



Figure 1: a) Time-dependent Pb(II) sorption onto NPG for b) Pseudo second order kinetics modelling, c)Intraparticle diffusion kinetic model for $[Pb(II)]_{initial}=0.3 \text{ mg L}^{-1}$ and $[Pb(II)]_{initial}=0.1 \text{ mg L}^{-1}$, corresponding to red and green symbols, respectively



Figure 2: Adsorption isotherm of Pb(II) onto NPG at pH 7: a) experimental and modeled data and b) linearization of the experimental data using the Freundlich model.



Figure 3: Evolution of the Pb^{2+} concentration relative to pH in presence of NPG at 50 mg g⁻¹.

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Figure 4: Pb(II) speciation under pH-adsorption-edge experimental condition but without NPG: a) corresponds to the Pb²⁺ speciation in solution and b) the evolution of the saturation index of Pb(OH)₂s, the unique species that was identified to precipitate.



Figure 5: A4F fractograms of free Pb(II) and Pb(II)-doped NPG (a) the UV and SLS detection with the Rg variation (blue cross), (b) the ICP-MS detection with m/z ²⁰⁷(Pb(II)).

Highlights

- Pb(II) binding experiments onto nanoplastics extracted from environmental microplastics were performed.
- Pb(II) was largely bound onto nanoplastics by specific adsorption and intra-particle diffusion.
- Nanoplastics could be significant vectors of metals in the environment

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