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Comparison of organic materials for the passive treatment of synthetic neutral mine drainage contaminated by nickel: Adsorption and desorption kinetics and isotherms

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# Abstract

Sorption is an effective process for the remediation of mine water with low metal concentrations. To identify promising low-cost organic sorbents for nickel (Ni), adsorption and retention properties of peat, compost, brown algae, sawdust, and wood ash were compared. Batch adsorption and desorption experiments were conducted at pH 7 in 0.05M NaNO<sub>3</sub> solutions to simulate the ionic strength and pH of a contaminated neutral drainage. Results of adsorption kinetic experiments were best represented by the Elovich model and the fastest rates were obtained with peat (796 075 mg g<sup>-1</sup> min<sup>-1</sup>) and compost (791 mg g<sup>-1</sup> min<sup>-1</sup>). Results of equilibration adsorption experiments were fitted to Langmuir and Freundlich isotherms and the

highest adsorption capacities were observed for peat (around 22 mg g<sup>-1</sup>) and compost (around 9 mg g<sup>-1</sup>). Desorption experiments revealed that peat and compost adsorbed more Ni and also released a lower percentage of the adsorbed metal upon exposure to Ni-free solutions.

**Keywords:** Neutral mine drainage, sorption, adsorption, desorption, kinetics, isotherms, nickel, organic materials.

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#### Introduction

Sorption of metals onto organic materials is a rapid process that can be used in passive treatment systems to lower the metal concentrations of contaminated neutral drainage (CND) below regulatory threshold values (Calugaru et al. 2018). The financial cost of transporting these materials to remote mining sites can be substantial and, thus, studying the sorption capacity of locally available materials is of great interest. Over the past few years, the sorption of metals onto various natural materials and industrial waste products has been investigated, and numerous solid substrates with good sorption capacities have been identified (Das et al. 2008; lakovleva and Sillanpaa 2013; Westholm et al. 2014; Zhou and Haynes 2010). Unfortunately, operational parameters such as pH, ionic strength and sorbent: sorbate ratios vary greatly between different studies, making comparisons difficult. To establish a proper comparison, different solid substrates must be tested under identical experimental conditions. In this study, the sorption properties of five low-cost organic substrates were compared with regards to their affinity for nickel. The substrates were selected for their sorption properties and potential local availability to eastern Canadian mining sites.

The main metal(loid) contaminants of CND are Cd, Cr, Co, Cu, Fe, Hg, Mn, Ni, U, Zn, As, Sb, and Se (Calugaru et al. 2018) and their concentrations are variable and mostly contingent on the mineralogy of each particular mine site. Ni is often present in CND and, in some instances, its concentration can exceed regulatory threshold values. In the province of Quebec (Canada), the mean monthly and maximum acceptable concentrations for nickel in mining effluents are, respectively, 0.5 mg L<sup>-1</sup> and 1 mg L<sup>-1</sup> (MDDELCC 2012). Metal toxicity in the environment depends on speciation, itself dictated by the local geochemical conditions. The ecotoxicity of nickel is mainly attributed to its free cation, Ni<sup>2+</sup> (Stokes 1988), which is the predominant species

at circum-neutral pHs. Other soluble nickel species include hydroxides, carbonates, sulfates, nitrates, chlorides and Ni-organic complexes. In unpolluted freshwater, an important fraction of the total nickel is bound to strong organic ligands (Xue et al. 2001). In mining effluents, competition with other cations such as Ca<sup>2+</sup> and Mg<sup>2+</sup> affects nickel complexation. Under these latter conditions, Ni-organic complexes tend to be more labile and the free Ni<sup>2+</sup> fraction more important (Guthrie et al. 2005; Mandal et al. 2002). The sorption of nickel onto organic substrates takes place at functional surface sites, such as carboxylic, phenolic and thiolic groups. The availability and structure of these sites depends on the type of organic matter. The sorption capacities of these substrates increase with pH (Stumm and Morgan 1996; Zhou and Haynes 2010) as their surface charge becomes more negative. Sorption of metals onto organic materials takes place at functional surface sites, such as carboxylic, phenolic and thiolic groups. The availability and structure of these sites depends on the type of organic matter. The sorption capacities of these substrates increase with pH (Stumm and Morgan 1996; Zhou and Haynes 2010) as their surface charge becomes more negative. Sorption of metals onto organic materials takes place at functional surface sites, such as carboxylic, phenolic and thiolic groups. The availability and structure of these sites depends on the type of organic matter.

In the present study, three different types of organic substrates were tested: biosorbents (brown algae and maple sawdust), humified materials (peat and compost) and biochar (wood ash). Biosorbents are defined in this paper as non-degraded inert organic matter. They have regularly spaced and organized surface functional groups disposed in repeatable patterns according to their cellular structure. One disadvantage of biosorbents in their field application is their high labile organic matter content. The latter may increase the chemical and biological oxygen demands in treated waters at the detriment of aquatic species (Wan Ngah and Hanafiah 2008; Zhou and Haynes 2010). A second disadvantage is the unknown impacts of biodegradation on the sorption capacity and specific affinity of these substrates. The vast majority of information on biosorbents originates from short-term experimental studies and their long-term behavior at the field-scale has yet to be investigated (Zhou and Haynes 2010).

Humified materials are organic substrates having undergone some degree of decomposition, polymerization and loss of volatiles. They are characterized by the presence of complex, high molecular weight, aromatic substances as well as inorganic solids (e.g., clay) or non-degraded plant material. Although highly aromatic, the structure of humic substances is poorly defined and the position of surface functional groups is heterogeneous and dependent on their chemical environment (Stumm and Morgan 1996). This complex arrangement and variable chemical properties of functional groups in humified materials can lead to multiple and possibly simultaneous sorption mechanisms such as adsorption, complexation, chelation, and coprecipitation. Metal sorption onto humified substances mostly occurs at carboxyl and phenolic functional groups (Stumm and Morgan 1996; Tan 2014). Humified substances are more recalcitrant to degradation than biosorbents and, thus, should theoretically be less affected during long-term field applications.

Biochars are a mixture of organic carbon-rich solids and ash resulting from the pyrolysis of biomass. Common by-products of biorefineries, their chemical composition depends on the nature of the parent biomass (e.g. wood residue, crop residue, manure) and production conditions (e.g. temperature, heating rate and time) (Calugaru et al. 2018; Mohan et al. 2014). They possess a high specific surface area and porosity making them good candidates for metal adsorption and sequestration (Calugaru et al. 2018; Mohan et al. 2014; Xu et al. 2014). Sorption of metals onto biochar can take place at both organic and inorganic functional groups. The sorption of metals onto the organic fraction of biochar is attributed to the presence of oxygen-containing functional groups such as carboxyl and phenol (Tong et al. 2011; Uchimiya et al. 2011). The role of the inorganic fraction in the sorption process is linked to the presence of calcite. The dissolution of calcite increases pH, favoring either sorption or precipitation of new metal-bearing solid phases (Arán et al. 2017; Xu et al. 2014; Zhou and Haynes 2010). For many

authors, the term adsorption refers strictly to 2D, non-specific or physical adsorption and the general term sorption is often used to include both specific (chemical) and non-specific adsorption as well as surface precipitation and co-precipitation. In this paper, the terms adsorption and desorption are meant to include all surface mechanisms and exclude only true precipitation.

Passive treatment systems designed for the final rehabilitation of closed or abandoned mine sites should require minimal maintenance and sorbent materials are typically used in a single, long-term sorption cycle (years ideally) before being disposed of. Hence, substrates need to be stable over equally long periods of time and metals need to be strongly bound to their surfaces (Zhou and Haynes 2010). In this study, the Ni adsorption and retention capacity of five low-cost organic substrates (brown algae, maple sawdust, wood ash, compost, and peat) is compared by conducting short-term kinetic and equilibration experiments in the absence of competing ions. The term "equilibration experiments" is used here for experiments in which a steady state with regards to adsorption-desorption reactions is presumably reached and does not necessarily refer to the attainment of a true equilibrium. The objective of the present study are to test different materials under identical operational conditions in order to help select the best candidates for long-term treatment studies of CND and to evaluate if certain physicochemical properties of substrates could be used to predict the outcome of sorption tests.

# Materials and methods

Substrate, stock solutions and equipment

The cold-water brown algae *Ascophyllum nodosum* (ASCO-PURE kelp) were provided by Organic Ocean Inc. (Rimouski, Quebec). They were air dried and gently crushed to less than 5mm. This substrate was used both in its original state (algae) and after being washed with hydrochloric acid (HCI-algae) to remove salts (Na, K, Mg, Ca), as described by Fourest and Volesky (1996). Briefly, the algae were stirred in a 0.1mol L<sup>-1</sup> HCI solution at a 1:50 solid:liquid ratio for 24 h with a magnetic bar. The HCI solution was then removed and the algae was rinsed extensively with deionized water and dried at 60 °C. The sphagnum peat moss was purchased from a local retailer (Home Depot, Montreal, Quebec). The compost was provided by P.W.I. Industries (Saint-Hyacinthe, Quebec). The wood ash (particles < 5 mm) was provided by Wood Ash Industries (Kirkland Lake, Ontario). It is a mixture of bottom and fly ash from a wood and gas co-generation plant and contains a small amount of gravel. All substrates were air dried and the compost and peat moss were sieved to less than 5 mm. All substrates were kept at room temperature in plastic containers or bags until use.

Experiments were performed at pH 7.0 in 0.05 M NaNO<sub>3</sub> solutions. The ionic strength and pH of this solution were chosen to reflect those of a nickel containing mine effluent ( $l \approx 0.029$  M and pH 7.31 ± 0.22) near Havre-Saint-Pierre (Quebec, Canada). Stock solutions of 0.05 M NaNO<sub>3</sub> and 0.175 M Ni(NO<sub>3</sub>)<sub>2</sub> were prepared in deionized water with pure chemicals (ACS grade). Initial nickel concentration in batch adsorption kinetic experiments (7.45 mg L<sup>-1</sup>) was much higher than measured in the field and was chosen as a worst-case scenario as well as to ensure analytical detection for all measurements. Nickel was analyzed by flame atomic absorption spectrometry (AAS; Perkin-Elmer AAnalyst-200). Calibration curves were built by sequential dilution of an AAS standard (~1000 ppm) solution (Perkin-Elmer). The detection limit was 0.01 ppm and the reproducibility of the measurements was better than 2 %, based on replicate analyses of this

solution. All pH measurements were carried out with a combined glass electrode (model Orion 9156BNWP, ThermoScientific), calibrated using three NIST-traceable buffer solutions (4.01, 7.00, and 10.00 at 25 °C), and a pH meter (model Orion Star A211 benchtop, ThermoScientific).

#### Substrate characterization

Results of substrate characterization are presented in Table 1. The water content (water weight/humid solid weight) of each solid substrate was determined prior to the adsorption experiments by drying 5-10 grams of substrate at 60 °C until constant weight was obtained. Specific surface areas (SSA) of the substrates were measured with a Micromeritics Gemini III 2375 surface analyzer using the N<sub>2</sub>-BET method BET PE2 AM-05 (Brunauer et al. 1938). The pH of a slurry of each substrate (2 g substrate: 100 mL deionized water) was measured after mixing thoroughly and letting stand for 1 hour. The point of zero charge (PZC) and cation exchange capacity (CEC) of each substrate were determined respectively by the salt addition method in 0.05 M NaNO<sub>3</sub> and by the ion adsorption method in 0.05 M NaCl at pH 4, 5, 6, 7, 8 and 9 (Bakatula et al. 2018). The total carbon ( $C_{TOT}$ ) and total organic and graphitic carbon ( $C_{org+g}$ ) contents of the substrates were measured by combustion at 1400 °C in an induction furnace (model CS744, LECO Corporation) according to Ma. 310-CS 1.0 (CEAEQ 2013). The Corg+g samples were pre-treated by immersion of the solid for at least one hour in a 50 % v/v HCl solution to dissolve carbonate phases. Dissolved organic carbon (DOC) was analyzed on filtered (0.45 µm) extracts of the slurries (20 g substrate (wet weight): 200 mL deionized water) shaken for 2 h at room temperature. Total organic carbon (TOC) of these extracts was determined at 680°C using a DOHRMAN, model DC-190 TOC analyzer, according to Standard Method 5310 B (APHA 2005).

Table 1 Physicochemical characteristics of organic substrates used in adsorption and desorption

#### experiments

	Brown algae	Sawdust	Compost	Peat moss	Wood ash
Water content (% w/w)	8.9 ± 0.9 <sup>a</sup>	5.0 ± 0.5	3.9 ± 0.3	11.4 ± 0.7	2.8 ± 0.7
SSA (m <sup>2</sup> g <sup>-1</sup> )	0.17	0.91	3.24	1.43	45.91
рН	5.47 ± 0.21	4.68 ± 0.36	7.30 ± 0.25	5.06 ± 0.39	9.32 ± 0.37
PZC	6.06 ± 0.11	4.72 ± 0.06	7.68 ± 0.01	3.42 ± 0.03	9.50 ± 0.07
CEC, pH 7 (meq 100 g <sup>-1</sup> )	242	74.8	46.5	124	61.7
C <sub>TOT</sub> (% w/w dry weight)	34.9 ± 0.1	50.1 ±0.4	17.3 ± 4.3	53.4 ±0.0	19.9 ± 2.2
C <sub>org+g</sub> (% w/w dry weight)	26.1 ± 0.4	49.3 ± 1.6	15.2 ± 0.6	53.3 ± 1.0	18.6 ± 2.3
DOC – water extract (mg L <sup>-1</sup> )	5680 ± 160	669 ± 6	80 ± 2	132 ± 7	6 ± 1

<sup>a</sup>Results are expressed as means  $\pm$  standard deviation using n = 3 (water content, pH, PZC), n=2 (C<sub>TOT</sub>,

C<sub>org+g</sub>), n=6 (DOC).

#### Adsorption and desorption kinetic experiments

For these experiments, 2 g of substrate and 200 mL of the 0.05M NaNO<sub>3</sub> solution were placed in 500 mL Erlenmeyers and were shaken (150 rpm) at room temperature (23  $\pm$  2 °C) on a MaxQ-2000 orbital shaker (ThermoScientific). Each experiment was conducted in duplicates and flask openings were covered by aluminum foil to limit evaporation. A third reactor without substrate was prepared in the same manner and indicated that the addition of nickel to the reaction vessels had no measurable effect on the pH of the solution. In order to prevent the pH adjustment phase and associated reactions from interfering with Ni sorption, the pH was adjusted to 7.0 twenty four hours prior to the addition of Ni to the slurries using dilute HNO<sub>3</sub> and NaOH solutions. Adsorption was initiated in each reactor upon the addition of 0.145 mL of a 0.175 M Ni(NO<sub>3</sub>)<sub>2</sub> solution to obtain an initial dissolved nickel concentration of 7.45 mg L<sup>-1</sup> (127 µmol L<sup>-1</sup>). At various time intervals over the next 24h, nine 6mL aliquots of the decanted liquid were collected. After 24 h, 100 mL of the liquid were decanted from the reactor. The solid and remaining liquid were transferred to a 50 mL polypropylene tube and centrifuged at 10000 x g for 10 minutes using a Heraeus Megafuge 8 centrifuge (ThermoScientific). Using a pipette, most of the liquid was discarded, except for the last 6 mL which were stored and later analyzed to determine the initial nickel concentration for the desorption experiment. The wet solid was transferred to an empty 500 mL Erlenmeyer flask and its wet weight was determined by weighing the polypropylene tube before and after this transfer. The dry weight of each solid transferred to the reactors was estimated from its wet weight and a pre-determined wet: dry weight ratio (detail of the ratio determination is presented in SI). Desorption was initiated in the reactor by adding 75-200 mL of a pre-adjusted Ni-free 0.05M NaNO<sub>3</sub> solution to the wet solid. The added NaNO<sub>3</sub> solution had been adjusted to a pH of 7.0 over a period of 24h with a separate aliquot of each substrate at a solid:liquid ratio of 1:100 and filtered through a 0.45 µm Millipore MCE filter. The volume added was adjusted according to the dry weight of the solid in each reactor in order to maintain a solid: liquid ratio of 1:100, like in the adsorption experiments. At different time intervals over a 24 h period, nine 6 mL aliquots of the decanted liquid were collected. For the HCI-treated algae, 4 mL aliquots were collected because initial liquid volumes were lower. Samples collected during the adsorption and desorption kinetic experiments were immediately filtered through 0.45 μm PVDF syringe filters, stored at 4°C and analyzed within 14 days.

# Treatment and modeling of experimental kinetic data

During adsorption experiments, the concentration of nickel adsorbed to the solids at each sampling point,  $q_{t_{ADS}}$  (mg g<sup>-1</sup>), was calculated using Eq.1:

$$q_{t_{ADS}} = \frac{[Ni]_{diss_0} \times V_0 - [Ni]_{diss_t} \times V_t}{m}$$
(1)

where V<sub>o</sub>, V<sub>t</sub>, are the volumes of liquid (L) in the reactor at the start of the experiment (time = 0) and at each sampling point (time = t) and m is the mass of the solid (g).  $[Ni]_{diss_0}$  and  $[Ni]_{diss_t}$  are the dissolved nickel concentrations (mg L<sup>-1</sup>) at time = 0 and time = t. During the course of desorption experiments, the concentration of nickel still adsorbed to the solids at each sampling point,  $q_{t_{ADS(des)}}$  (mg g<sup>-1</sup>), was calculated using Eq.2:

$$q_{t_{ADS(des)}} = q_{t_{ADS(f)}} - \left(\frac{([Ni]_{diss_t} - [Ni]_{diss_0}) \times V_t}{m} + \frac{([Ni]_{diss_{t-1}} - [Ni]_{diss_0}) \times V_s}{m}\right)$$
(2)

where  $q_{t_{ADS(f)}}$  is the nickel concentration adsorbed to the solid at the end of the adsorption experiment. V<sub>t</sub>, m and  $[Ni]_{diss_t}$  are the volumes of liquid (L), mass of solid (g) and dissolved nickel concentrations (mg L<sup>-1</sup>) in the reactor at the time of sampling (t).  $[Ni]_{diss_0}$  is the initial dissolved nickel concentration in the reactor, calculated using the final dissolved nickel concentration in the adsorption experiment, the water content of the wet solid transferred to the reactor and the volume of NaNO<sub>3</sub> added to the reactor at the start of the desorption experiment. V<sub>s</sub> is the volume of sample (6 mL) removed from the reactor at the previous sampling (t-1) and  $[Ni]_{diss_{t-1}}$  is the dissolved nickel concentration of this sample. Finally, the concentration of nickel desorbed from the solid during the desorption experiment,  $q_{t_{DES}}$  (mg g<sup>-1</sup>), was obtained by subtracting the concentration of nickel still adsorbed to the solid at each sampling point,  $q_{t_{ADS(des)}}$  from the adsorbed nickel concentration at the end of the adsorption experiment,  $q_{t_{ADS(des)}}$  from the adsorbed nickel concentration at the end of the adsorption

$$q_{t_{DES}} = q_{t_{ADS(f)}} - q_{t_{ADS(des)}}$$
(3)

Results of the adsorption and desorption kinetic experiments were compared using three different reaction kinetics-based models: the pseudo-first order (PFO), pseudo-second order (PSO) and Elovich models. These empirical models were fitted to the experimental data using non-linear regressions, because linear regressions have been found to introduce a methodological bias and erroneously favor the PSO model (Tran et al. 2017; Xiao et al. 2018).

Non-linear regressions were carried out using the "Solver add-in" in Microsoft Office Excel (2010 version), as described by Tran et al. (2017). The PFO model was developed by Lagergren (1898) to describe the adsorption of oxalic acid and malonic acid onto charcoal. The PSO model, often referred to as Ho's PSO model (Ho 1995), was initially proposed by Blanchard et al. (1984) to describe the removal of heavy metals from water using zeolites. The differential and integrated, non-linear form of the PFO (Eq. 4 and 5) and PSO (Eq. 6 and 7) equations are:

$$\frac{dq_t}{dt} = k_1(q_e - q_t) \tag{4}$$

$$q_t = q_e (1 - e^{-k_1 t}) \tag{5}$$

$$\frac{dq_t}{dt} = k_2 (q_e - q_t)^2 \tag{6}$$

$$q_t = \frac{q_e^2 k_2 t}{1 + k_2 q_e t} \tag{7}$$

where  $q_e$  and  $q_t$  are the final and time = t (min) adsorbed nickel concentrations (mg g<sup>-1</sup>) respectively, and  $k_1$  (min<sup>-1</sup>) and  $k_2$  (g mg<sup>-1</sup> min<sup>-1</sup>) are the pseudo-first and pseudo-second order rate constants. The Elovich equation was initially proposed by Roginsky and Zeldovich (1934) for the adsorption of carbon monoxide onto manganese dioxide. The differential and integrated, non-linear form of the Elovich equation (Eq. 8 and 9) are:

$$\frac{dq_t}{dt} = \alpha e^{-\beta q_t} \tag{8}$$

$$q_t = \frac{1}{\beta} \ln \left( 1 + \alpha \beta t \right) \tag{9}$$

where  $q_t$  is the adsorbed nickel concentration (mg g<sup>-1</sup>) at time = t (min),  $\alpha$  (mg g<sup>-1</sup> min<sup>-1</sup>) is the initial rate constant and  $\beta$  (mg g<sup>-1</sup>) is the desorption constant. It must be noted that the Elovich model is physically unsound as adsorbed nickel concentrations increase endlessly, never

reaching a steady state. Thus, it is only suitable to describe kinetics far from equilibrium (Tan and Hameed 2017).

## Equilibration experiments

Six pre-weighted 50 mL polypropylene centrifuge tubes containing 0.4 g of substrate and 40 mL of the 0.05M NaNO<sub>3</sub> solution were stirred (150 rpm) horizontally at room temperature (23  $\pm$  2 °C) on a MaxQ-2000 orbital shaker (ThermoScientific). Each experiment was conducted in triplicate and, during the first 24 h pH adjustment period, pH was adjusted to 7.0 in all tubes using dilute HNO<sub>3</sub> and NaOH solutions. To monitor the evolution of pH in the absence of nickel, one extra blank tube was prepared in the same manner. After the pH adjustment period, adsorption was initiated in tubes by adding varying volumes of a 0.175 M Ni(NO<sub>3</sub>)<sub>2</sub> solution to obtain initial nickel concentrations of: 0.6, 7, 15, 29, 59 and 103 mg L<sup>-1</sup> (10, 119, 254, 494, 1000, 1750 μmol L<sup>-1</sup>). Tubes were shaken for at least 20 hours and, after allowing the solids to decant for 5 minutes, one 6 mL aliquot of the supernatant was collected from each tube. Subsequently, tubes were centrifuged at 10000 x g for 10 minutes and all the liquid was removed using a pipette. The tubes containing the wet solids were re-weighted and wet weights of the solids were determined. The dry weight of the solids in the tubes was estimated from their wet weight and a pre-determined wet: dry weight ratio (detail of the ratio determination is presented in SI). Desorption was then initiated in the centrifuge tubes by adding 38 mL of a pre-adjusted 0.05 M NaNO<sub>3</sub> Ni-free solution to the wet solid. The added NaNO<sub>3</sub> solution had been adjusted to a pH of 7.0 over a period of 24 h with a separate aliquot of each substrate at a solid:liquid ratio of 1:100 and filtered through a 0.45 µm Millipore MCE filter. Tubes were shaken for at least 20 hours and subsequently, one 6 mL aliquot of the liquid was collected from each tube. Samples collected during the equilibration experiments were immediately filtered through 0.45 µm PVDF syringe

filters, stored at 4 °C and analyzed within 14 days.

#### Isotherm models

The concentrations (q<sub>ads</sub>) and percentages (%ads<sub>ads</sub>) of nickel adsorbed to the solids at the end of adsorption equilibration experiments were calculated using Eq. 10 and 11, respectively:

$$q_{ads} = \left( [Ni]_{diss_0} - [Ni]_{diss_f} \right) \times \frac{V}{m}$$
(10)

$$\%ads_{ads} = \frac{\left( [Ni]_{diss_0} - [Ni]_{diss_f} \right)}{[Ni]_{diss_0}} \times 100 \tag{11}$$

where V is the volume of liquid (L), m is the mass of solid (g) and  $[Ni]_{diss_0}$  and  $[Ni]_{diss_f}$  are the initial and final dissolved nickel concentrations (mg L<sup>-1</sup>), respectively. The concentrations (q<sub>des</sub>) and percentages (%ads<sub>des</sub>) of nickel adsorbed to the solids at the end of the desorption equilibration experiments were calculated using Eq. 12 and 13, respectively:

$$q_{des} = q_{ads} - \left( [Ni]_{diss_f} - [Ni]_{diss_0} \right) \times \frac{V}{m}$$
(12)

$$\%ads_{des} = \%ads_{ads} - \left(\frac{(q_{ads} - q_{des})}{q_{ads}} \times \%ads_{ads}\right)$$
(13)

where  $q_{ads}$  is the final adsorbed nickel concentration at the end of the adsorption experiment, V is the volume of liquid (L), m is the mass of solid (g) and  $[Ni]_{diss_f}$  is the final dissolved nickel concentration (mg L<sup>-1</sup>).  $[Ni]_{diss_0}$  is the initial dissolved nickel concentration in the reactor, calculated using the final dissolved nickel concentration in the adsorption experiment, the water content of the wet solid transferred to the reactor and the volume of NaNO<sub>3</sub> added to the reactor at the start of the desorption experiment. Equilibration (adsorption and desorption) results were fit to Langmuir and Freundlich isotherms. The Langmuir model was originally developed to describe the adsorption of a gas at the surface of a solid (Langmuir 1918). The underlying hypotheses for this model are that adsorption is driven by physical forces (electrostatic, London- van der Waals), that all surface sites have identical affinities for the sorbate and adsorption is limited to a monolayer coverage of the solid surface. The general form of the Langmuir equation is:

$$q = q_{max} \frac{K_{ads} [A]}{1 + K_{ads} [A]}$$
(14)

where q is the adsorbed species concentration,  $q_{max}$  is the maximal concentration of surface sites, [A] is the sorbate concentration in solution and  $K_{ads}$  is the adsorption constant. The Langmuir parameters were derived from a linear regression of the experimental data to Eq. 15:

$$\frac{[A]}{q} = \frac{1}{k_{ads}q_{max}} + \frac{[A]}{q_{max}}$$
(15)

The Freundlich model was originally introduced to describe the logarithmic decrease in adsorption energy with increasing surface coverage of the solid (Freundlich 1907). It was later interpreted as a model describing sorption on an heterogeneous surface whose sites have a variable affinity for the sorbate (Davis et al. 2003; McLean and Bledsoe 1992). The general form of the Freundlich equation is:

$$q = \mathbf{K}[A]^n \tag{16}$$

where q is the adsorbed species concentration, [A] is the sorbate concentration in solution and K and n are empirically determined constants. K is referred to as the Freundlich constant and is related to the maximum sorption capacity of the sorbent whereas n is a measure of the non-linearity of the adsorption and is related to the affinity or binding strength of adsorption sites

(Davis et al. 2003; Stumm and Morgan 1996). When studying the adsorption of metals on complex substrates such as the ones evaluated in this study, the underlying hypothesis of the Langmuir model are not respected. First, complex organic substrates can form chemical bonds with metals, and second, they typically host diverse, surface functional groups (e.g., carboxylic, phenolic, thiolic) that have varying affinities for these metals. In the present study, the Langmuir model is used empirically to evaluate and compare the sequestration potential of different sorbing materials in the neutral pH range. Hence,  $q_{max}$  and  $K_{ads}$  are empirically determined constants, not to be interpreted, respectively, as the maximal concentration of surface sites and the adsorption constant originally defined in Langmuir adsorption model.

# Geochemical modeling

Geochemical modeling was conducted to obtain the equilibrium nickel speciation under the experimental conditions of this study. Speciation calculations were performed in open system conditions ( $pCO_2 = 38.5 Pa$ ,  $E_H = 515 mV$ ) at 25 °C in the absence of a solid substrate (sorbent) using the freeware, chemical equilibrium model VMINTEQ version 3.0 (Gustafsson 2016). The VMINTEQ thermodynamic database is a revised version of the original USEPA MINTEQA2 database (Allison et al. 2011) that contains updated and expanded data from the NIST Critical Stability Constant database.

#### **Results and discussion**

#### Geochemical modeling

We first modeled the effect of pH on nickel speciation for the Ni-NaNO<sub>3</sub> open system solution used in the kinetic experiments (0.05 M NaNO<sub>3</sub>, 7 mg·Ni L<sup>-1</sup>). In this system, the solution reached

saturation with respect to an amorphous nickel carbonate phase (NiCO<sub>3(s)</sub>, Ksp =  $10^{-11.2}$ ) at pH values above 7.33, becoming the dominant species above pH 7.5 (Fig. S1a, SI). We then modeled the effect of dissolved organic carbon on nickel speciation in the same solution at pH 7.5 using the NiCA-Donnan formulation with the default settings (Fig. S1b, SI). Under these settings, 82.5 % of the DOC is composed of fulvic acids whose assumed carbon content is 50 %. We chose a fixed pH value of 7.5 for this model to evaluate the effect of DOC on the solubility of NiCO<sub>3</sub>(s). As DOC increases, more Ni-DOC complexes are formed and the concentration of all other Ni species decreases. For DOC concentrations above 68 mg L<sup>-1</sup>, all nickel species are dissolved at pH 7.5. Finally, we modeled NiCO<sub>3</sub>(s) saturation as a function of pH for the range of nickel concentrations investigated in the equilibration experiments (Fig. S2, SI).

# Adsorption and desorption kinetics

During all adsorption experiments, the dissolved nickel concentrations decreased rapidly (within minutes) (Figure 1a). Invariant concentrations, presumably representing steady states between adsorption and desorption, were reached in all experiments with the exception of the wood ash slurry. In the presence of wood ash, dissolved nickel concentrations kept decreasing slightly even after 10 hours of equilibration. During the course of the adsorption experiments, recorded pH values (6.81 to 7.48) were below NiCO<sub>3(s)</sub> saturation (7.33) for all substrates except the HCl-algae (Fig. S3a SI). For this substrate, pH increased to nearly 7.5 within the first hour of the experiment before settling back below 7.33. It is unlikely that nickel carbonate precipitated during this pH variation as the brown algae slurry contains high DOC concentrations (Table 1).



Figure 1. Results of the kinetic adsorption (a) and desorption (b) experiments. Dissolved nickel concentrations with time of equilibration.

During desorption experiments (Figure 1b), the dissolved nickel concentrations increased rapidly for all substrates and reached constant concentrations for the compost, wood ash, sawdust and peat slurries after 0.5 to 4 hours. In the case of algae and HCl-algae, dissolved nickel concentrations kept increasing slightly beyond 10 hours of equilibration. While more nickel was adsorbed by peat and compost during adsorption experiment (Figure 1a), desorption trials indicated that they released three to seven times less nickel than the other substrates when exposed to a nickel-free solution. In desorption experiments, recorded pH values (6.70 to 7.63) were below NiCO<sub>3(s)</sub> saturation for all substrates (Fig. S3b, SI). The saturation pH was higher during desorption (7.66) than adsorption (7.33) experiments because dissolved nickel concentrations were lower, with maximum values of 1.6 mg L<sup>-1</sup> (27  $\mu$ mol L<sup>-1</sup>).

#### Treatment and modeling of kinetic experimental data

As the use of data points recorded after the attainment of a steady state is likely to lead to erroneous conclusions (Tran et al. 2017), partial data sets were used to fit the experimental data (Eq.1) to the kinetic models (Figure 2a, Table 2) (Fig S4, SI). The three reaction-kinetic models were compared on the basis of their coefficient of determination ( $R^2$ , Eq. 17) and chi-squared test ( $\chi^2$ , Eq. 18). An  $R^2$  value closer to unity and a  $\chi^2$  value closer to zero indicate the better fit.

$$R^{2} = 1 - \frac{\sum (q_{t(exp)} - q_{t(cal)})^{2}}{\sum (q_{t(exp)} - q_{t(mean)})^{2}}$$
 Eq. 17

$$\chi^{2} = \sum \frac{(q_{t(\exp)} - q_{t(cal)})^{2}}{q_{t(cal)}}$$
 Eq. 18

In these equations,  $q_{t(exp)}$  (mg g<sup>-1</sup>) is the experimental adsorbed nickel concentration at time = t,  $q_{t(cal)}$  (mg g<sup>-1</sup>) is the adsorbed nickel concentration achieved from the model using the Solver addin, and  $q_{t(mean)}$  (mg g<sup>-1</sup>), the mean of the  $q_{t(exp)}$  values.



Figure 2. Adsorbed (a) and desorbed (b) nickel concentrations (q<sub>tADS</sub>) as a function of equilibration time during HCl-algae experiment and Elovich, PSO and PFO model fits. Empty symbols identify data used for model fitting.

For the adsorption experiments, the estimated  $q_e$  as well as the R<sup>2</sup> and  $\chi^2$  values were always better with the PSO model, therefore PFO results will not be discussed. Experimental steadystate values,  $q_{e(exp)}$  (mg g<sup>-1</sup>), were estimated from adsorption experiments and values are presented in Table 2. For adsorption experiments,  $q_{e(exp)}$  were estimated as the mean value calculated from the last (wood ash), last two (compost, sawdust, HCl-algae, algae) or last six (peat) data points. The Elovich model provided a better fit to the selected adsorption data sets (Table 2). The steady-state adsorbed nickel concentration ( $q_e$ ) estimated by the PSO model was 6-12 % lower than the experimental value ( $q_{e(exp)}$ ) for all substrates except for peat for which the two values were identical within one standard deviation. The rate constants derived from the Elovich and PSO models ( $k_2$  and  $\alpha$ ) indicate that the adsorption kinetics increase in the following order: peat > compost > algae> HCl-algae > sawdust > wood ash. The ability of the model constants to reproduce the experimental data (Eq. 6 and 8) was evaluated and highlighted the presence of a fast initial nickel uptake which was not well described by either model. The presence of this fast initial uptake and its influence on the Elovich model fit have been discussed elsewhere (McLintock 1967).

Table 2. Comparison between experimental steady-state adsorbed [Ni]  $(q_{e(exp)})$ , pseudo-second order and Elovich model fitting parameters for adsorption kinetic experiments

		Pseudo-se			Elovich				
Material	$q_{e(exp)}^{a}$ (mg g <sup>-1</sup> )	<i>q</i> e (mg g⁻¹)	<i>k</i> 2 (g mg <sup>-1</sup> min <sup>-1</sup> )	R <sup>2</sup>	X <sup>2</sup>	β (mg g <sup>-1</sup> )	α (mg g <sup>-1</sup> min <sup>-1</sup> )	R <sup>2</sup>	X <sup>2</sup>
Peat moss	0.707 ± 0.015	0.69	9.70	0.85	0.003	28.4	796075	0.88	0.003
Compost	0.703 ± 0.004	0.65	2.91	0.79	0.06	21.9	791	0.93	0.02
HCI-algae	0.620 ± 0.016	0.57	1.01	0.84	0.17	17.1	7.65	0.96	0.02
Wood ash	≥ 0.565 ± 0.005	0.50	0.34	0.83	0.44	17.5	1.07	0.99	0.01
Algae	0.531 ± 0.009	0.47	3.87	0.66	0.10	26.5	110	0.96	0.01
Sawdust	0.502 ± 0.007	0.47	0.41	0.87	0.32	19.3	1.35	0.97	0.02

<sup>a</sup>Results are expressed as mean  $\pm$  standard deviation with n = 2 (wood ash), n=4 (compost, HCl-algae,

algae, sawdust), n=12 (peat).

For the desorption experiments, data points recorded prior to reaching a steady state were used to fit the experimental data (Eq. 3) to the Elovich and PSO kinetic models (Figure 2b, Table 3) (Fig. S5, SI). The PSO model generally provided a better fit to the selected desorption data set. The steady-state desorbed nickel concentrations ( $q_e$ ) estimated by the PSO model were identical, within one standard deviation, to the experimental value ( $q_{e(exp)}$ ) for all substrates but the algae. No correlation was observed between  $k_2$  and  $\alpha$  for the desorption models which may be caused by the limited range of concentrations represented by the desorption data sets. Table 3. Comparison between experimental steady-state desorbed [Ni] ( $q_{e(exp)}$ ), pseudo-second order and Elovich models for desorption experiments

	Pseudo-second order					Elovich			
Material	$q_{e(exp)^a}$ (mg g <sup>-1</sup> )	<i>q</i> e (mg g <sup>-1</sup> )	<i>k</i> 2 (g mg <sup>-1</sup> min <sup>-1</sup> )	R <sup>2</sup>	X <sup>2</sup>	β (mg g <sup>-1</sup> )	α (mg g <sup>-1</sup> min <sup>-1</sup> )	R <sup>2</sup>	X <sup>2</sup>
Peat moss	0.016 ± 0.002	0.017	41.1	0.93	0.001	316	0.030	0.89	0.001
Compost	0.025 ± 0.002	0.027	15.2	0.93	0.004	208	0.034	0.93	0.002
HCI-algae	0.092 ± 0.008	0.091	8.94	0.93	0.005	69.1	0.269	0.87	0.011
Wood ash	0.092 ± 0.006	0.094	6.83	0.93	0.01	91.7	0.732	0.94	0.005
Algae	0.078 ± 0.005	0.087	1.94	0.96	0.06	56.9	0.028	0.91	0.07
Sawdust	0.110 ± 0.002	0.111	1.38	0.99	0.007	66.5	0.097	0.93	0.02

<sup>a</sup>Results are expressed as mean  $\pm$  standard deviation using n = 2 (wood ash), n=4 (compost, HCI-algae,

algae, sawdust), n=12 (peat).

Kinetic-based models such as the PSO and Elovich models are empirical fits to kinetic curves that lack physical significance and combine many different models with different controlling mechanisms (Tan and Hameed 2017). Kinetic-based models are applicable when the mechanism at play is chemisorption on solids that are either non-porous or porous and exhibit high diffusion coefficients (Inglezakis et al. 2019). For porous solids (activated carbon, biochar, zeolites), diffusional mass transfer is the rate-determining step and adsorption processes may require days to weeks to reach a steady state (Inglezakis et al. 2019; Rees et al. 2014; Tran et al. 2017). Multiple adsorption mechanisms are bound to take place on the biochar, biosorbents and humified materials used in the present study. Diffusion-based models may help identify adsorption mechanism and rate-controlling steps, but these models are more complex and can be computationally demanding and time consuming (Inglezakis et al. 2019; Tan and Hameed 2017). Nevertheless, the PSO and Elovich models still allow us to compare the performance of these materials, but the derived kinetic parameters are solely descriptive and only apply to the experimental conditions investigated in this study. Both models describe the increase in

adsorbed nickel at the surface of the substrates, but they do not satisfactorily describe the attainment of a steady state or the fast initial nickel uptake.

# Adsorption equilibration experiments and NiCO<sub>3(S)</sub> saturation

Over the range of nickel concentrations investigated and in the absence of DOC, the pH at which NiCO<sub>3(s)</sub> saturation is reached varies from pH 6.77 ([Ni] = 103 mg L<sup>-1</sup> or 1750  $\mu$ mol L<sup>-1</sup>) to pH 7.88 ([Ni]= 0.6 mg L<sup>-1</sup> or 10  $\mu$ mol L<sup>-1</sup>) (Fig. S2 and S6, SI). These saturation pH boundaries are conservative as they apply to an open system with no DOC, whereas our samples were equilibrated in closed tubes and solutions contained various amounts of DOC. Ideally, the precipitation of a solid Ni phase should be avoided during equilibration adsorption experiments but, in the present study, field relevant pH conditions were used and thus oversaturation with respect to NiCO<sub>3(s)</sub> could not be avoided. During the adsorption equilibration experiments, recorded pH values were below the NiCO<sub>3(s)</sub> saturation value for samples with initial nickel concentrations of 15 mg L<sup>-1</sup> or more. Therefore, precipitation of a solid nickel carbonate phase may have taken place in samples with high initial nickel concentrations. During the desorption experiment, the pH values recorded were always below the NiCO<sub>3(s)</sub> saturation limit.

# Adsorption and desorption equilibration experiments

The final adsorbed nickel concentrations ( $q_{ads}$ , Eq. 10) were used to derive the empirical Langmuir and Freundlich isotherm parameters (Table 4). Data points from samples with initial nickel concentrations of 0.6 mg L<sup>-1</sup> (10 µmol L<sup>-1</sup>) were ignored as they were clear outliers in the linear regression. Moreover, results from the algae experiment could not be fitted to the Langmuir model as data points show no clear curvilinear trend. The results from the HCl-algae experiment follow a more curvilinear trend but the adsorption capacity of this substrate is not

significantly increased relative to the untreated algae (Figure 4). According to the R<sup>2</sup> coefficients, results for algae, HCI-algae, peat and wood ash are more appropriately represented by the Freundlich isotherm while those for the compost and maple sawdust could be fitted to both models (Table 4). When compared graphically, the model that best fits the experimental data is not readily identified (Figure 3). For example, the fit to the Langmuir model appears better for peat, but this is not reflected by the R<sup>2</sup> coefficients. Furthermore, it should be noted that the precipitation of nickel carbonate might have occurred in samples with higher initial [Ni], reducing their final nickel concentrations and apparently increasing their adsorbed nickel concentrations, thus favoring a fit to the Freundlich model.

Table 4. Nickel adsorption isotherm model fitting coefficients for the studied substrates

	Langmuir			Freundlich			
Material	R <sup>2</sup>	q <sub>max</sub> (mg g⁻¹)	K <sub>ads</sub> (L mg <sup>-1</sup> )	R <sup>2</sup>	n	K (L <sup>n</sup> mg <sup>n-1</sup> g <sup>-1</sup> )	
Doot mass	0.95	22 ± 0	$0.15 \pm 0.00$	0.09	0.96 ± 0.22	27+02	
Peal moss	0.85	22 ± 9	$0.15 \pm 0.09$	0.98	$0.80 \pm 0.23$	2.7 ± 0.2	
Compost	0.98	8.8 ± 2.8	0.13 ± 0.07	0.98	0.57 ± 0.07	$1.1 \pm 0.1$	
HCI-treated algae	0.65	6.9 ± 5.0	0.03 ± 0.03	0.97	0.68 ± 0.20	0.27 ± 0.15	
Wood ash	0.96	3.9 ± 1.4	0.04 ± 0.02	0.99	0.58 ± 0.07	0.24 ± 0.03	
Algae	-	-	-	0.88	0.97 ± 0.48	0.05 ± 0.04	
Maple sawdust	0.97	3.8 ± 0.9	0.03 ± 0.02	0.98	0.62 ± 0.15	0.21 ±0.11	



Figure 3. Compost and peat adsorption isotherms with Langmuir and Freundlich model fits

Langmuir model parameters derived from nickel adsorption studies onto organic materials similar to the ones used in this study can be found in the literature (Table S1, SI). The reported  $q_{max}$  values vary within one to two orders of magnitude of each other. These expected discrepancies may originate from the provenance or preparation of the tested organic materials as well as differences in experimental conditions. According to the published  $q_{max}$  values, brown algae display some of the highest nickel adsorption capacities. The highest values were obtained with brown algae of much smaller particle size (<0.5 mm) than used in our experiments (<5 mm). In contrast, under our experimental conditions, the adsorption capacities of the investigated substrates increased in the following order: sawdust/wood ash, HCl-algae, compost, and peat. This illustrates why different low-cost sorbent materials need to be compared under identical field-relevant experimental conditions.

During the adsorption experiments, pH was adjusted to 7.0 during a 24 h pH adjustment period and was then allowed to drift after the addition of nickel. It varied between the beginning (full symbols) and the end (empty symbols) of the experiments for all substrates except wood ash, for which the pH of all reactors remained stable at 7.0 (Fig. S7, SI ). For the other substrates, a decrease of up to 1 pH unit was observed in most reactors. For the algae and HCl-algae, the pH also decreased in the nickel-free control vessel, likely due to an incomplete pH equilibration prior to the beginning of the experiment. For sawdust, peat and compost, the pH decrease was proportional to the initial nickel concentration and could be interpreted as resulting from an exchange between H<sup>+</sup> and Ni<sup>2+</sup> ions on the solid surfaces.



Figure 4. Adsorption and desorption isotherms. Results are presented as average values from triplicate reactors, with error bars representing one standard deviation from the mean.

The amount of nickel released during the desorption experiments varied amongst the different substrates (Figure 4). Desorption results show that adsorption was not reversible and that peat and compost adsorbed and retained more nickel than the other substrates. They also show that

more nickel was retained by the HCl-algae than the untreated algae and that sawdust, wood ash and HCl-algae displayed a similar hysteresis upon desorption yielding linear isotherms.

# Organic materials properties and adsorption

In terms of nickel adsorption capacity, retention capacity and adsorption kinetics, the performance of humified materials surpassed that of the other substrates tested in this study. None of the physicochemical properties of the five investigated organic substrates could have alone allowed us to predict this outcome (Table 1). Humified materials have lower specific surface areas than wood ash, and yet they adsorb more nickel because of their organic surface functional groups. Furthermore, their cation exchange capacities (CEC) are not the highest, implying that a material's capacity to adsorb and release sodium ions cannot predict its behavior with regards to nickel ions. This being said, peat does possess the second highest CEC (124 meg 100 g<sup>-1</sup>) after brown algae, and the CEC measurement for brown algae could be biased by the presence of sodium alginate, a water soluble component in the algae's cellular membrane. The point of zero charge (PZC) of peat (3.42) is the lowest suggesting a negative surface charge at circum-neutral pH, supporting the highest Ni adsorption capacity. Regarding compost, despite having a high PZC (7.68) and, therefore, a mean positive surface charge at circum-neutral pH, it displays a strong Ni adsorption capacity. This may be interpreted as evidence of chemical interactions with Ni, rather than electrostatic binding, most likely bidentate binding to carboxylic groups, consistent with the proton exchange (decrease in pH upon adsorption).

Leaching of dissolved organic carbon also plays a role in the substrate's capacity to adsorb nickel. DOC forms soluble complexes with divalent nickel and reduces the concentration of free nickel ions in solution (Fig. S1b, SI). Therefore, as DOC increases in a given system, the amount of charged nickel species decreases, leading to less nickel adsorption. The humified materials

who performed better, release much less DOC than the other investigated biosorbents (e.g., algae, sawdust), but they release more DOC than the wood ash.

#### Conclusion

The Ni adsorption and retention capacity of five low-cost organic materials was compared by conducting short-term batch experiments at pH 7.0 in 0.05 M NaNO<sub>3</sub>. Results of adsorption and desorption kinetic experiments revealed that peat and compost adsorbed nickel more rapidly than the other substrates. A steady state was reached within 40 minutes in the presence of peat, whereas it took nearly 10 hours with compost, sawdust, algae and HCI-algae. For wood ash, a porous solid for which diffusional mass transfer is the expected rate-determining step, adsorption kinetics were slower and steady state was not reached within 23 hours. More nickel was adsorbed by peat and compost during kinetic experiments and desorption trials indicated that they released three to seven times less nickel than the other substrates when exposed to a nickel-free solution. In equilibration experiments, peat and compost also displayed a higher adsorption capacity than the other substrates and, irrespective of the initial nickel concentrations tested, retained a higher percentage of the adsorbed nickel when exposed to nickel-free solutions. Whereas humified materials performed better than other materials under the experimental conditions of this study, other factors that affect metal sequestration need to be evaluated before moving on to long-term column experiments. These factors include the presence of competing ions, pH and the Ni retention capacity of the substrates over periods of several weeks.

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