Sorption of Heavy Metals by Ferralic-Arenosol and Vertic-Luvisol: A Comparative Isothermal Study

Tlou Mosekiemang¹, Oagile Dikinya² & Elisha Toteng²

Correspondence: Tlou Mosekiemang, Department of Environmental Science, University of Botswana, Gaborone, Botswana. Tel: 267-355-2527. E-mail: mosekiemang@mopipi.ub.bw

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Abstract

Modeling of sorption data is essential for characterizing retention of heavy metals by the soil matrices as well as to predict the fate of heavy metals in the environment. We carried out a study to evaluate adsorption characteristics of metallic ions (Cr^{3+} , Pb^{2+} and Cu^{2+}) as sole sorbates as well as in coexistence onto clay extracts of Vertic-Luvisol and Ferralic-Arenosol, the generally under-researched yet wide-spread semi-arid soils. Additionally, sorption kinetics of these heavy metals was evaluated in these soils. Equilibrium sorption data for both non-competitive and competitive metal systems were found to obey the Langmuir isotherm. Overall metal loading capacity was higher in Vertic-Luvisol than Ferralic-Arenosol whereas metal-wise affinity to binding sites favoured sorption of Cr^{3+} over Pb^{2+} and Cu^{2+} in that order. Contrary to the hypothesis of this study, the preferential sorption process was found to be driven by cationic properties as opposed to soil properties. Adsorption of metallic ions onto soils was significantly reduced in competitive systems than non-competitive systems, a phenomenon attributable to the antagonistic effects of coexisting species. Cr^{3+} suppressed sorption of Pb^{2+} and Cu^{2+} on both soils as validated by the competitive adsorption capacity ratios which were < 1 in all instances where Cr^{3+} coexisted with either one of Pb^{2+} or Cu^{2+} . The adsorption kinetics of metal ions in both soils followed the pseudo-second-order rate law, suggesting chemical sorption as the rate-limiting step of the adsorption process.

Keywords: adsorption isotherms, adsorption kinetics, binary-metal system, mono-metal system

1. Introduction

The ability of soils to adsorb metal ions is of special interest and has consequences both for agricultural production and environmental remediation. With the current awareness of heavy metals as very toxic contaminants there have been many studies considering adsorption by solid surfaces as the most important mechanism for controlling metal content in soil solution and natural waters (Echeverria, Morera, Mazkiaran, & Garido, 1998; Bradl, 2004; Appel, Ma, Rhue, & Reve 2008; Javaid, Bajwa, Shafique, & Anwar, 2011). The most important chemical processes affecting the behavior and bioavailability of heavy metals in soils are those concerned with sorption mechanisms of metals from liquid phase onto solid phase (Rybicka, Calmano, & Breeger, 1995; Covelo, Andrade, & Vega, 2004). Moreover, clay minerals and to a lesser extent organic matter have been identified as among the major soil constituents that interact with almost all soil contaminants especially positively charged metallic species. Thus, clays often represent a short-term sink of heavy metals in soils, a trait attributable to their adsorptive properties. The heavy metal bearing component of the soil, which in the perspective of this study is the clay mineral, can be a good and reliable indicator of environmental pollution. Clay minerals play an important role in accumulation, adsorption/desorption, as well as exchange processes of metal ions (Wahba & Zaghloul, 2007). In general, heavy metal retention depends on the type of clay minerals contained in the soil. It is well known that basic soil characteristics influencing the sorption of metals, amongst

¹ Department of Chemistry, University of Botswana, Gaborone, Botswana

² Department of Environmental Science, University of Botswana, Gaborone, Botswana

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others, depends on contents of organic matter, clay, Fe and Mn oxides and hydroxides, and carbonates (Bradl, 2004; Usman, 2008; Irha, Steinnes, Kirso, & Petersell, 2009).

In order to describe the adsorption characteristics of adsorbates, experimental equilibrium data are most frequently modeled by the relationships developed by Freundlich and Langmuir (Echeverria et al., 1998; Limousin, Gaudet, Charlet, Szeknect, Barthes & Krimissa, 2007; Gomes, Cavaco, Quina, & Gando-Ferreira, 2010). Usually, the experimental approach involving sorption of sorbates onto sorbents is to gather data and produce adsorption isotherms, which are in turn interpreted to explain adsorption process. Analysis of sorption data is important for characterizing retention of contaminants by soil (Hinz, 2001) and in particular the Langmuir and Freundlich models are the most suitable for describing single solute adsorption of heavy metals to soils (Hinz, Gaston, & Selim, 1994; Bolster & Hornberger, 2007). The Langmuir and Freundlich isotherms are widely used since they have an ability to describe experimental results across a wide range of experimental adsorption data. These isotherm models can be transformed into linear forms to obtain adjustable parameters by graphical means or by linear regression analysis (Altin, Ozbelge, & Dogu, 1998). The analysis of isotherms may provide useful information about the retention capacity and the sorption strength by which sorbate is held onto the soil (Sipos, Nemeth, Kis, & Mohai, 2009).

The present contribution of this work is to provide information on heavy metal sorption characteristics of Vertic-Luvisol and Ferralic-Arenosol, the two dominant soil types in the semi-arid region of the world. These two soil types, especially Arenosols are among the under researched yet significantly wide-spread in the semi-arid region of the world (Ngole & Ekosse, 2008; Mosekiemang & Dikinya, 2012). According to Hartemink and Huting (2008), the total extent of Arenosols in the semi-arid regions covers approximately 273 million ha in the southern and western part of Africa. To this end, arenosols have not attracted much attention in research because of the many inherent attributes unfavourable for productive agriculture (Moroke, Dikinya, & Patrick, 2009). Thus the main aims of this study were (1) to investigate sorption characteristics of Cr³⁺, Cu²⁺ and Pb²⁺ ions as single adsorbates in solution as well as when they are in coexistence in solution, (2) to use sorption isotherms, particularly Langmuir and Freundlich models to determine the sorption capacity in the two types of clay extracts in view of assessing their vulnerability to pollution and (3) to evaluate the sorption kinetics of these selected heavy metals in these semi-arid soils. As a rule, soil properties and composition differ with soil type and it is logical to expect variations in heavy metal sorption characteristics of different soil types. Thus, the main hypothesis being tested is that heavy metal sorption may be influenced by soil type.

2. Method

2.1 Site Description and Sample Preparations

The soil samples used in this study were collected from Glen Valley Horticultural Project in Eastern Gaborone, Botswana lying between Latitudes (24°35'23.56"S and 24°37'01.14"S) and Longitudes (25°58'43.29"E and 25°58'16.74"E). Typical soils are predominately Ferralic-Arenosols (GARo) and Vertic-Luvisol (ALVv) characterized by reddish loamy sands and dark grey clays (FAO, 1998), respectively. The soil samples were collected from the plough layer (0-20 cm) and/or Ap horizon (FAO, 1998), air dried, and passed through a 2-mm sieve. To separate and remove the clay component from the 2-mm soil fraction, the sieved soil was mixed with double distilled water and stirred with a motorized stirrer at 1000 rpm overnight to concentrate the clay particles. The suspension was left to settle for 24 h and the top layer was subsequently decanted, filtered and dried in an oven at about 100 °C to constant weight to obtain the clay extracts.

2.2 Characterization of Parent Soil Samples and Clay Extracts

The soil samples were analyzed for physico-chemical characteristics, including particle size distribution (sand %, silt %, clay %), soil colour, pH and cation exchange capacity (CEC) using laboratory standard methods (Breitbart, 1988; van Reeuwijk, 1993 in Dikinya & Areola, 2010). Clay extracts were further analyzed/scanned for morphological and chemical characteristics (Figure 1 and 2) using the Philips XL30 Environmental Scanning Electron Microscope/EDAX. Identification of primary and secondary minerals in the clay samples was achieved using a Philips PW 3710 X-Ray diffractometer operated at 40 kV with a Cu anode radiation and a graphite monochromator. Subsequently, interpretation of data and diffractograms was achieved by Philips X'PERT graphics and identify software (2001 version). The surface area of clay extracts (GARo and ALVv) were measured with a Micromeritics FlowSorb (III) 2310 instrument using the BET single point method and N₂ gas adsorption.

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2.3 Adsorption Experiments

Two equilibrium batch experiments were conducted to determine mono-metal and binary-metal adsorption characteristics of selected heavy metals (Cr3+, Cu2+ and Pb2+) onto clay extracts of GARo and ALVv. In mono-metal experiment, single-metal solutions were prepared by adding 20 mL of one metal solution to a 50 mL propylene tubes containing 20 mg of the soil mineral matrix (clay extract). In this experiment, concentrations of single-metal solutions were 0, 2.5, 5, 10, 25 and 50 mg/L of each of Cr³⁺, Cu²⁺ and Pb²⁺ and were carried out separately for one metal at a time. In binary-metal experiment, simultaneous competitive sorption of metals onto soil mineral matrix (clay extracts) were studied through batch equilibration. Binary-metal solutions were prepared such that each solution contained equal proportions of the two co-existing metals at a ratio of 1:1. The pairing of metals in a binary-metal solutions were aimed at achieving every possible combination option available to pair any two of the three analytes (Cr³⁺ with Pb²⁺; Cr³⁺ with Cu²⁺ and Cu²⁺ with Pb²⁺). Similarly, for monometal experiment, six concentration levels of 0, 2.5, 5, 10, 25 and 50 mg/L of each bimetal combination were maintained and sorption experiments were achieved by weighing 20 mg of the soil mineral matrix (clay extract) into 50 mL propylene tubes and equilibrating with a 20 mL of a binary-metal solution. In both experiments (mono and binary-metal), the tubes were shaken in a horizontal shaker for 3h at 25±2 °C, a previously determined optimal equilibration time (Alumaa, Kirso, Petersell, & Steinnes, 2002; Adebowale, Unuabonah, & Olu-Owalabi, 2006; Ghorbel-Abid & Trabeisi-Ayadi, 2011) followed by centrifuging at 3000 rpm for 20 min and subsequent filtration through a filter paper (Whatman No. 42) to obtain a clear filtrate. The filtrate pH was then selectively measured in the filtrate solution of mono-metal batch whereas the entire consignment was assayed for heavy metal concentration.

2.4 Sorption Kinetics

Batch equilibrium experiments described above were followed by another experiment setup to determine the sorption kinetics of Cr3+, Cu2+ and Pb2+ onto clay extracts of soils of interest (GARo and ALVv). In this experiment, just like in the batch experiments, both working temperature and solution-sorbent ratio were maintained at 25±2 °C and 1 g/L respectively, whereas the metal ion concentration and pH were kept constant at 100 mg/L and 6.5 ±0.02, respectively. To maintain solution's cationic composition and to avoid metal (hydro) oxide precipitation during the course of sorption experiments, solution's pH was kept within the desired range by adding either NaOH or HCl (Mettler Toledo 320 pH meter). Adsorbate-adsorbent contact time was varied at different times within 0-400 min range by shaking the solutions at 100 min time interval for the time range described earlier. Shaking was achieved by using a Labcon temperature controlled horizontal shaker followed by centrifuging, filtering and subsequent assaying for heavy metal content. In all cases described above, assaying of heavy metal content was determined using a using flame atomic spectrophotometer (Varian, SpetrAA FS-220, Australia). Whereas all chemicals used in this work were analytical reagent quality: Pb(NO₃)₂; Cu(NO₃)₂:2.5H₂O; CrCl₃·6H₂O NaOH and HCl were supplied by Sigma-Aldrich, South Africa.

2.5 Data Analysis

In the case of batch equilibrium studies, the amount of adsorption at equilibrium Q_e (mg/g), was calculated by:

$$Q_e = (C_0 - C_e) \times \frac{V}{m} \tag{1}$$

 $Q_e = (C_0 - C_e) \times \frac{v}{m}$ (1) where C_0 and C_e (mg/L) are liquid-phase concentrations of adsorbates (Cr³⁺, Cu²⁺ and Pb²⁺) at initial and equilibrium, respectively. V is the volume of the solution (L) and m is the mass of adsorbent (clay extract) used (kg). In batch kinetic studies, the amount of adsorption of metal ions at time (t), q_t (mg/kg), was calculated by:

$$q_t = (C_0 - C_e) \times \frac{v}{m} \tag{2}$$

where C_0 and C_e (mg/L) are the liquid-phase concentrations of adsorbates (Cr³⁺, Cu²⁺ and Pb²⁺) at initial and any time t, respectively. Parameters V and m are similar to Equation 1.

2.5.1 Single-Metal Sorption Isotherms

The amounts of Cr³⁺, Cu²⁺ and Pb²⁺adsorbed were fitted to the Langmuir (Equation 3) and Freundlich (Equation 5), respectively. The general Langmuir sorption model is expressed as follows;

$$Q_e = \frac{Q_{max}bC_e}{1+bC_e} \tag{3}$$

where Q_e is the uptake of the metal ion by the clay extract (mg/kg), Q_{max} is the upper limit of Q_e which represents maximum uptake (adsorption maxima) of the clay extract (mg/kg). C_e is the equilibrium concentration of metal ions in solution (mg/L) and b is the Langmuir adsorption constant. This model implies that saturation of the adsorptive sites is possible, and ideally Q can eventually reach a maximum value (Q_{max}) . Therefore parameters

 Q_e and C_e can be experimentally determined whereas Q_{max} and b can either be calculated from least-square fitting or a simple plot of a linearized form of Equation 3 which yields;

$$\frac{C_e}{Q_e} = \frac{1}{Q_{max}b} + \frac{C_e}{Q_{max}} \tag{4}$$

 $\frac{c_e}{Q_e} = \frac{1}{Q_{max}b} + \frac{c_e}{Q_{max}}$ A plot of $\frac{c_e}{Q_e}$ vs c_e gives a linear plot with the slope of $\frac{1}{Q_{max}}$ and intercept of $\frac{1}{Q_{max}b}$

The general form of the Freundlich Equation is:

$$Q_e = k_F C_e^{1/n} \tag{5}$$

where Q_e and C_e are the same as in Equation 3, and k_F and n relate to the capacity and intensity of adsorption, respectively. The Freundlich model is based on sorption of metal ions on heterogeneous surface and can be linearized by taking logarithms:

$$\log Q_e = \log k_F + \frac{1}{n} \log C_e \tag{6}$$

 $\log Q_e = \log k_F + \frac{1}{n} \log C_e$ where the parameters k_F and n are determined similarly to the Langmuir case.

2.5.2 Binary-System Sorption

The mutual interference effects of binary-metal ions (Cr³⁺-Pb²⁺; Cr³⁺-Cu²⁺ and Cu²⁺-Pb²⁺) on sorption dynamics were probed using competitive adsorption capacity ratios (CACR) for the two adsorption isotherms under study. According to Qin et al. (2006); Mahamadi and Nharingo (2010) and Zhu, Hu and Wang, (2012), O'max/Omax and k_F'/k_F represents CACR for the Langmuir and Freundlich equations, respectively. The isotherm constants, Q_{max} and Q_{max} indicate the Langmuir adsorption capacity in single and binary system, while k_F and k_F are the Freundlich adsorption capacity of singe metal solution and in the presence of the other metal, respectively. The value of Q'_{max}/Q_{max} and k'_F/k_F has implications on the effect of mixing metals in a solution; synergistic $(Q)_{max}/Q_{max}$ and $k)_F/k_F > 1$; no net interaction $(Q)_{max}/Q_{max}$ and $k)_F/k_F = 1$; or antagonistic $(Q)_{max}/Q_{max}$ and $k)_F/k_F$

2.5.3 Sorption Kinetics

In order to study adsorption kinetics, various rate equations (first-order, second-order and inter-particle diffusion) were applied to the time-dependent kinetic data in order to obtain rate coefficients. The pseudo-first order (Lagergren 1898, in Argun, Dursun, Ozdemir, & Karatas, 2007), Equation (7); pseudo-second order, Equation (8) and inter-particle diffusion, Equation (9) are expressed as follows:

$$\log(q_e - q_t) = (\log q_e) - k_1 t \tag{7}$$

$$\frac{dq_e}{d_t} = k_2 (q_e - q_t)^2 \tag{8}$$

$$q_t = k_i t^{1/2} + C (9)$$

where q_t is the metal uptake per unit weight clay (mg/g) at the time t, q_e is the metal uptake per unit weight of clay (mg/g) at equilibrium, and k₁ (min⁻¹) and k₂ (g/mg/min) are the rate constants of the first-order and pseudo second order kinetic equations, respectively. C (mg/g) and k_i (mg g⁻¹ h^{-1/2}) are the equation's constant and intra-particle diffusion rate constant, respectively. Linearized plots of Equations. (7-9), $\log(q_e - q_t)$ vs t (pseudo-first order), t/q_t vs t (pseudo-second order) and q_t vs $t^{1/2}$ (intra-particle diffusion) were used to identify the best kinetic model that describe sorption kinetics of Cr^{3+} , Cu^{2+} and Pb^{2+} on clay adsorbents. The choice of the best kinetic model was based on superiority of the r^2 correlation coefficient for linearity of the plots.

3. Results

3.1 Soil Morphology, Classification and Physico-Chemical Properties

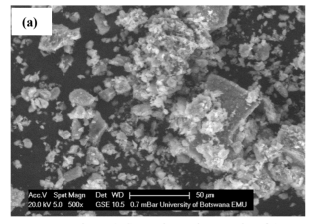
The morphological features of the soil samples are presented in Table 1. These soils were classified according to the FAO (1998) - world reference base for soil resources as Vertic Luvisol (ALVv) and Ferralic Arenosol (GARo), respectively. For GARo, the colour of the top (Ap) horizon was 5YR hue with value 4-6 and chroma 4 to 8 and predominantly red. While ALVv was predominantly grey along with a 10YR hue with value 2 to 3 and chroma 1 to 4. Laboratory analyses supported field observations. Texture of the soil samples ranged between sandy clay loam and sandy loam for ALVv and GARo, respectively. ALVv had the highest clay content and the least pH value whereas GARo had the least clay content and highest pH value. The CEC values ranged between 33.08 to 61.54 cmol/kg for GARo and ALVv, respectively. The high CEC values were contributed largely by 2:1 swelling clays (Sposito, 1984).

Table 1. The physico-chemical properties of the soils

Clay type	Parameter	Description
	Colour	Red (5YR 4/6 – 4/8)
GARo	Mineral Composition (%)	Ferrosilite magnesian (26); hematite (58); Quartz (16)
	Specific surface area	$212 \text{ m}^2 \text{ g}^{-1}$
	Cation Exchange Capacity	33.08 cmol/kg
	$pH-(H_2O)$	7.49
	Clay(%) / Silt(%) / Sand (%)	28.2% / 14.5% / 57.3%
	Colour	Grey (10YR 2/3-1/4)
ALVv	Mineral Composition (%)	Smectite (31); Clintonite (30); Kaolinite (22); Quartz (17)
	Specific surface area	$362 \text{ m}^2 \text{ g}^{-1}$
	Cation Exchange Capacity	61.54 cmol/kg
	$pH-(H_2O)$	7.35
	Clay(%) / Silt(%) / Sand (%)	35.5% / 16.0% / 48.5%

3.2 Morphological Characteristics of Clay

According to the electron microscope micrographs, ALVv clay particle morphology was platy and less porous (Figure 1) than that of GARo (Figure 2) which was more spherical, porous and exhibited the tendency to conglomerate as shown in the 500× micrograph. The primary clay minerals contained in ALVv were smectite, kaolinite and quartz whereas GARo contained significant amounts of hematite and quartz (Ngole & Ekosse, 2008). Nonetheless, the diagnostic rod shape entities which are symptomatic to the presence of hematite were visible in the 5000× magnification of Figure 2, and traces of platy entities indicative of the presence of smectite were visible in the 5000× magnification of Figure 1. The nature of clay mineral has profound effects on the adsorptive capacity and heavy metal retention of soils, with smectite and kaolinite (the typical 2:1 swelling clays) having relatively larger surface area and adsorptive capacity than hematite - a distinctive 1:1 clay (see Table 1). The bulk chemical composition of the two clay types was dominated by SiO₂ and Al₂O₃ which made up 66.6% and 31.3%, respectively in ALVv and 69.5% and 11.2%, respectively in GARo (Table 2). However, GARo had a higher Fe₂O₃ content than ALVv corroborating the presence of hematite (an Fe rich mineral). Losses Due to Ignition (%LDI), which was indicative of the presence of volatiles were negligibly low at 0.03% and 2.5% for ALVv and GARo, respectively.



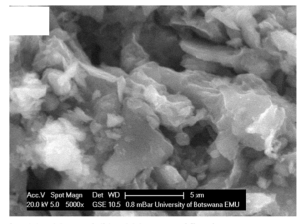


Figure 1. SEM photomicrographs of ALVv under (a) 500× and (b) 5000× magnifications

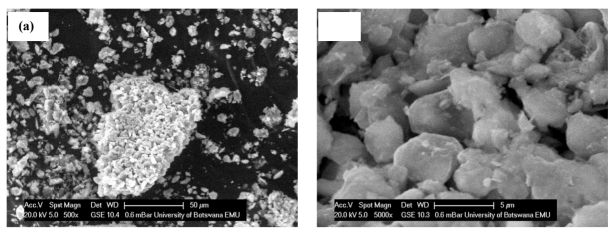


Figure 2. SEM photomicrographs of GARo under (a) 500× and (b) 5000× magnifications

Table 2. Chemical composition of clay extracts

	Mineral Composition (%)										
Clay extract	SiO ₂	Al_2O_3	Fe ₂ O ₃	MnO	MgO	CaO	Na ₂ O	K ₂ O	TiO ₂	P_2O_5	Total
ALVv	66.6	31.3	0.9	0.3	< d.1	< d.1	< d.1	0.3	0.3	< d.1	99.7ª
GARo	69.5	11.2	10.5	< d.1	< d.1	< d.1	< d.1	3.7	2.6	< d.1	97.5 ^b

 $d.1 = detection limit; ^aLDI = 0.03\%; ^bLDI = 2.5\%.$

3.3 Adsorption Isotherms of Single Metal System

The graphical illustrations of the content of metal adsorbed onto solid phases as a function of incremental metal concentrations are shown in Figure 3, where C_e and Q_e are concentrations of dissolved and adsorbed metal forms, respectively. The isotherms of the soil types were identical in shape. They reached a well-defined plateau at high metal ion concentration (± 40 mg/L) and were of type-L, based on the classification described by Giles, McEvan, Nakhawa and Smith (1960). These types of isotherms are characterized by decreasing slopes as equilibrium concentration of metal ions increases. This phenomenon is attributable to high affinity of the sorbent at low concentration of sorbates in solution.

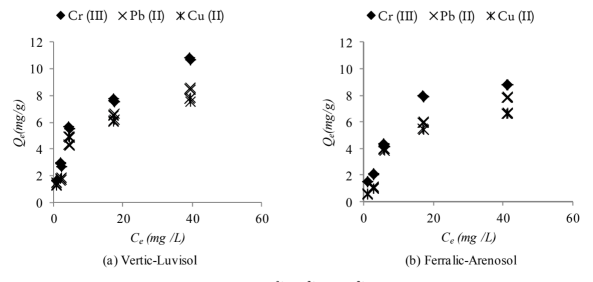


Figure 3. Equlibrium adsorption isotherms of Cr^{3+} , Cu^{2+} and Pb^{2+} ions onto ALVv and GARo clays in a mono-metal system. C_e is the concentration of metalic ions in equilibrium, and Q_e is the adsorption

Augmentation of metal ions in contact with solid phases resulted in significant acidification of the systems as shown in Figure 4. In both soils, the initial pH was circum-neutral then dropped to ± 4.5 when the solid phases approached saturation.

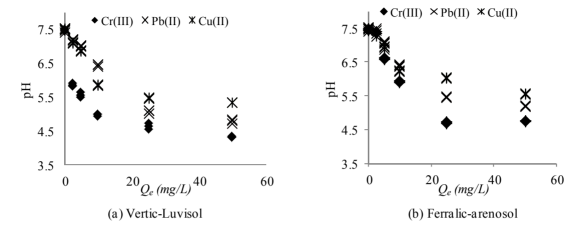


Figure 4. Resultant equlibrium solution's pH as a function of incremental metal ion concentration interacting with the solid phases

3.4 Isothermal Data Fitting

3.4.1 Monometal System

The experimental data for the uptake of Cr³⁺, Pb²⁺ and Cu²⁺ were processed in accordance with two most widely used Langmuir and Freundlich classical adsorption isotherms. The results of regression analysis for calculating the parameters of Langmuir and Freundlich models are presented in Table 3.

Table 3. Langmuir and Freundlich isothermal parameters in mono-metal systems

	The Langmuir isotherm constants								
Clay type	Metal ion $(n)^a$	<i>b</i> (L mg ⁻¹)	$Q_{max} (\text{mg g}^{-1})$	r^2					
ALVv	$Cr^{3+}(10)$	3.1±0.9	32.2±3.1	0.9824					
	$Cu^{2+}(10)$	1.2±1.1	12.2±2.1	0.9679					
	$Pb^{2+}(9)$	2.1±3.1	28.6±2.5	0.9477					
GARo	$Cr^{3+}(8)$	2.8 ± 2.1	10.4±1.9	0.9863					
	$Cu^{2+}(10)$	0.8 ± 2.4	6.85 ± 2.1	0.9563					
	$Pb^{2+}(10)$	1.1 ± 0.7	9.37 ± 2.4	0.9645					
		The Freundlich isotherm constants							
Clay type	Metal ion (n)	$k_F (\mathrm{mg}\;\mathrm{g}^{\text{-}1})$	n	r^2					
ALVv	$Cr^{3+}(9)$	42.0±0.4	0.67±0.01	0.9442					
	$Cu^{2+}(10)$	29.4 ± 0.9	0.45 ± 0.01	0.8619					
	$Pb^{2+}(9)$	16.4 ± 0.3	0.31 ± 0.01	0.9114					
GARo	$Cr^{3+}(10)$	30.5 ± 0.7	0.59 ± 0.01	0.9472					
	$Cu^{2+}(10)$	18.9±1.1	0.51 ± 0.01	0.9210					
	Pb ²⁺ (9)	11.7±0.9	0.62 ± 0.01	0.9324					

 $⁽n)^*$ denotes the number of experimental points in the isotherm.

3.4.2 Binary-Metal System

The mutual interference effects of binary-metal ions (Cr³⁺-Pb²⁺; Cr³⁺-Cu²⁺ and Cu²⁺-Pb²⁺) on sorption as well as single and binary-metal isothermal parameters are presented in Table 4. The results indicate that overall adsorption capacity was significantly decreased by mutual coexistence of competing species (Antoniadis, Tsadilas, & Ashworth, 2007).

Table 4. Mono-metal and binary-metal systems isothermal parameters

Clay	Metal	System	k_F \/ k_F	Q_e'/Q_e	Isotherms					
					Langmuir			Freundlich		
ALV v	_				b	Q_{max}	r^2	k_F	n	r^2
		Cr ³⁺	_	_	3.1±0.9	32.2±3.1	0.982	42.0±0.4	0.67	0.944
	_	Pb^{2+}	_	_	2.1±3.1	28.6±2.5	0.948	16.4±0.3	0.31	0.911
	_	Cu^{2+}	_	-	1.2±1.1	12.2±1.1	0.968	29.4±0.9	0.45	0.862
	Cr ³⁺	Cr ³⁺ +Cu ²⁺	0.28	0.36	5.8±2.1	11.5±2.8	0.977	11.8±2.4	0.68	0.985
	Cr ³⁺	$Cr^{3+} + Pb^{2+}$	0.25	0.35	4.9±0.9	11.2±2.4	0.982	10.5±2.7	0.79	0.945
	Pb^{2+}	$Pb^{2+}+Cu^{2+}$	0.45	0.35	2.3±2.5	10.1±1.9	0.978	7.98±0.9	0.61	0.930
	Pb^{2+}	$Pb^{2+}+Cr^{3+}$	0.50	0.27	2.0±1.6	7.69±2.1	0.974	8.23±1.4	0.84	0.951
	Cu^{2+}	$Cu^{2+} + Pb^{2+}$	0.23	0.92	2.1±3.4	11.2±3.7	0.977	6.75±2.4	0.49	0.933
	Cu^{2+}	$Cu^{2+} + Cr^{3+}$	0.11	0.44	2.2±2.4	5.32±2.1	0.981	3.25±1.1	0.53	0.962
						Langmuir Freundlich				h
GAR	=				b	Q_{max}	r^2	k_F	n	r^2
0	=	2±								
	-	Cr ³⁺	-	-	2.8±2.1	10.4±1.9	0.986	30.5±0.7	0.59	0.947
	-	Pb^{2+}	-	-	1.1 ± 0.7	9.37 ± 2.4	0.965	11.7±0.9	0.62	0.932
	-	Cu ²⁺	-	-	0.8 ± 2.4	6.85 ± 2.1	0.956	18.9 ± 1.1	0.51	0.921
	Cr^{3+}	$Cr^{3+}+Cu^{2+}$	0.39	0.96	4.1±0.9	9.98 ± 0.9	0.910	12.1 ± 2.2	0.91	0.942
	Cr^{3+}	$Cr^{3+} + Pb^{2+}$	0.33	0.97	3.9 ± 2.1	10.1±3.9	0.936	10.0 ± 3.1	0.47	0.961
	$Pb^{2^{+}}$	$Pb^{2+}Cu^{2+}$	0.61	0.94	2.2±1.6	8.85 ± 2.5	0.961	7.13±4.0	0.45	0.935
	$Pb^{2^{+}}$	$Pb^{2+}+Cr^{3+}$	0.74	0.39	2.8 ± 3.1	3.65 ± 4.4	0.971	8.62 ± 3.1	0.81	0.956
	Cu^{2+}	$Cu^{2+}+Pb^{2+}$	0.49	0.69	3.9±1.7	4.65±3.6	0.969	9.32±3.5	0.79	0.962
	Cu^{2+}	$Cu^{2+}+Cr^{3+}$	0.47	0.33	2.4 ± 3.2	2.29 ± 4.3	0.983	8.91±2.6	0.81	0.978

3.5 Sorption Kinetics

Adsorption capacities of metal ions were measured as a function of time (0-400 min) while keeping the initial concentration, adsorbent dosage and/or sorbent-solution ratio, temperature and pH constant at 100 mg/L, 1 g/L, $25\pm2^{\circ}\text{C}$ and pH = 6.5 ± 0.02 , respectively. The results showed two contrasting adsorption processes among the two soil types. In GARo, sorption was two phased, in which the initial phase was characterized by a fast sorption process in the first 200 minutes followed by a slow sorption process. This suggests that availability of sorption sites on solid phases significantly influenced the sorption process. Conversely, the sorption process in ALVv was linear, in which sorption increased with time duration, suggesting that time was a limiting factor in the sorption process.

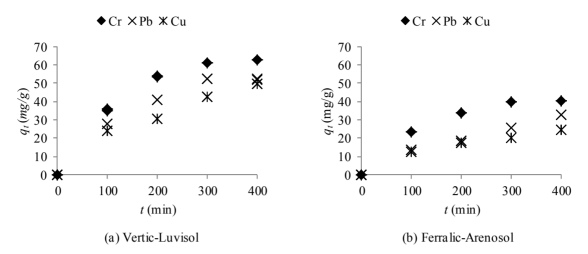


Figure 5. Effect of equlibration time on adsorption of Cr^{3+} , Pb^{2+} and Cu^{2+} on (a) ALVv and (b) GARol. The pH was held constant at pH = 6.05 ± 0.2 , metal ions concentrations held constant at 100 mg L⁻¹ and working temperature at 25 °C

The kinetic parameters for modeling adsorption of Cr^{3+} , Pb^{2+} and Cu^{2+} ions by the soil types of interest are shown in Table 5. In both soil types, the sorption kinetics of the metal ions followed pseudo-second order as shown by $r^2 > 994$.

4. Discussion

4.1 Comparison of Sorption Processes Based on Isothermal Plots

4.1.1 Monometal System

The graphical illustrations of sorption isotherms for single metal systems showed that an increase in metal ion concentration resulted in an increase in overall adsorption (Figure 3). Moreover, metal-wise comparison of isothermal plots for the soil types showed a general preference of sorption for Cr^{3+} over Pb^{2+} and Cu^{2+} . This is in agreement of Irha, Steinnes, Kirso, and Petrsell (2009) along with de Matos, Fontes, da Costa and Martinez, (2001) who found that adsorption of metallic ions to solid phases favoured trivalent ions over divalent ions. This preferential trend of sorption described above was similar for both soil types. However, higher sorption amounts per metallic ion (Q_e) were achieved in ALVv (Cr^{3+} , 11 mg/g; Pb^{2+} , 8 mg/g & Cu^{2+} , 7 mg/g) than GARo (Cr^{3+} , 8 mg/g; Pb^{2+} , 7 mg/g & Cu^{2+} , 6 mg/g). Notably, a series of salient points were observed on isothermal plots when the concentration of equilibrated metal ions increased in the ranges 0-5, 10-20, and 25-40 mg/L for ALVv and 0-2.5, 5-10, and 10-20 mg/L for GARo, respectively. As expected, the results revealed an important underlying role of soil type on sorption *i.e* soil properties such as specific surface area of clay particles as well as cation exchange capacity were influential in determining sorption process of metal ions. de Matos et al. (2001) demonstrated the influence of soil properties on sorption processes of pollutants. Thus, ALVv with superior soil properties (Table 2) had higher metal loading capacity than its counterpart.

Furthermore, augmentation of metal ions in contact with the solid phases resulted in a significant acidification of the system (Figure 4). Soil-wise comparison showed that adsorption-induced acidity was more pronounced in ALVv than GARo. The acidification of the equilibrium solutions observed during the sorption of metallic ions by the solid phases is usually explained by the displacement of H^+ in favour of multivalent $M^{2^+/3^+}$ ions (Bradl, 2004). Thus, the adsorbent surface was dominated by negative charges that attracted positively charged metallic ions. It is widely reported that the surface charges on most clay crystals emanate from hydroxyl (OH) groups (Adebowale et al., 2006; Bradl, 2004), however this study did not measure any Infra-Red (IR)-related parameters to identify functional groups.

4.2 Isothermal Data Fitting

4.2.1 Langmuir Model

Experimental data showed that the Langmuir model had a slightly higher correlation coefficient for linearity and conclusively suggested that it was reliable and valid in modelling sorption data of Cr^{3+} , Pb^{2+} and Cu^{2+} in both soils. An isothermal soil-wise comparison based on Langmuir parameter, Q_{max} (maximum adsorption capacity

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per unit mass of adsorbent) showed higher cumulative sorption of metal ions by ALVv than GARo (Table 3). It seems reasonable to link higher retention capacities of metal ions to cation exchange capacity (CEC) and particle specific surface area, values of which were greater in ALVv than GARo. Studies by Appel et al. (2008), Bradl (2004) and de Matos et al. (2001) demonstrated that heavy metal loading capacity is correlated to high specific surface area of clay particles. Metal-wise comparison based on Langmuir parameter Q_{max} established a higher sorption capacity for Cr^{3+} than Pb^{2+} and Cu^{2+} . The preferential-sorption of metal ions based on Q_{max} can be ordered into $Cr^{3+} > Pb^{2+} > Cu^{2+}$ for the two soils. This adsorption sequence was similar for the two soils and did not reveal any underlying soil based variability amongst the soil types. This decisively suggested that the sorption process was driven by factors other than the expected soil properties.

The Langmuir parameter b, characterizing the binding strength between metal ions and functional centres on the sorption surface showed higher values for ALVv than GARo (Table 3). This could be ascribed to a series of cationic properties notably, the differences in the first hydrolysis constant (pK_h) amongst metal ions. According to Bradl (2004), ion exchange prefers cations with low a pK_h, thus Cr^{3+} with the lowest pK_h value sorbed more than other metal ions. The same sorption trend was reported by Merdy, Gharbi, and Lucas (2009) and Fonseca, Figueiredo, Rodrigues, Queiroz, and Tavares (2011) with the former attributing the trend to the conventional hard-soft-acid-base (HSAB) principle. The hard ions are those with low polarizability and small ionic size and prefer hard ligands and form weak complexes with them. On the other hand, soft ions have greater affinity for soft sites and are more strongly bounded (Sposito, 1984).

4.2.2 Freundlich Model

Based on the Freundlich coefficient k_F , related to the metal ion's affinity to the solid phase, sorption of metals were ordered into the following sequence: $Cr^{3+} > Pb^{2+} > Cu^{2+}$ for the both soils. Thus, k_F values did not unearth any sorption variabilities between the two soils. However, the values were higher in ALVv than GARo and in each instance Cr^{3+} was twice higher than Pb^{2+} which in turn was twice higher than Cu^{2+} . Likewise, the Freundlich adsorption intensity constant, n for the metal ions and soils under study did not show any variability, except that all values were less than 1.

4.3 Effect of Cationic Competition on Sorption

4.3.1 Binary-Metal System

To investigate the overall competitive effect on sorption by metal ions, CACRs were developed based on the sorption model that had the best fit of experimental data which turned out to be the Langmuir equation (Table 4). The Langmuir-derived CACRs showed an overall suppressive effect of competition on sorption (Zhu et al., 2012). For instance, in ALVv, the total and/or cumulative sorption amount in non-competitive system was ΣQ_{max} $(Cr^{3+}, Pb^{2+}, Cu^{2+}) = 73.0 \text{ mg/kg}$ which was significantly higher than the total sorbed amount in the binary-metal (competitive) system, ΣQ_{max} (Cr^{3+} - Cu^{2+} , Cr^{3+} - Pb^{2+} , Pb^{2+} - Cu^{2+} , Pb^{2+} - Cr^{3+} , Cu^{2+} - Pb^{2+} , Cu^{2+} - Cr^{3+}) = 57.0 mg/kg. In contrast, the total sorbed amount in GARo was higher in binary-metal system than in single-metal (non-competitive) system, where ΣQ_{max} (Cr³⁺, Pb²⁺, Cu²⁺) = 26.2 mg/kg for monometal system and ΣQ_{max} $(Cr^{3+}-Cu^{2+}, Cr^{3+}-Pb^{2+}, Pb^{2+}-Cu^{2+}, Pb^{2+}-Cr^{3+}, Cu^{2+}-Pb^{2+}, Cu^{2+}-Cr^{3+}) = 39.5 \text{ mg/kg for binary metal system.}$ Metal-wise comparison within a soil type showed that in all cases where Cr³⁺ co-existed with the other metal in solution, it exerted suppressive effect on the other competing metal. This suppressive effect was substantiated by the CACR of less than 1 in all such cases (antagonistic effect of coexisting species). For instance in the Cr^{3+} - Cu^{2+} and Cr^{3+} - Pb^{2+} combinations in ALVv, the Q_{max} for Cr^{3+} as the target analyte was 11.5 and 11.2 mg/kg in the former and latter combinations, compared to 5.32 and 7.69 mg/kg for Cu²⁺ and Pb²⁺, respectively. Furthermore, the suppressive effect of Cr³⁺ over other competing metal ions was manifested in GARo. In similar binary-metal combination systems (Cr^{3+} - Cu^{2+} & Cr^{3+} - Pb^{2+}), the Q_{max} for Cr^{3+} as the target analyte were 9.98 and 10.1 mg/kg for the former and latter combinations, respectively. While the Q_{max} for Pb²⁺ and Cu²⁺ under similar ionic combinations were 3.65 and 2.29 mg/kg, respectively. Ionic charge and/or metal ion's oxidation state seemed have been the main driver of the competitive adsorption process in a binary-metal system (Irha et al., 2009; de Matos et al., 2001). The ionic charge favoured Cr³⁺ over Pb²⁺ and Cu²⁺, hence the observed suppressive effect (Antoniadis et al., 2007). Summarily, the suppressive effect of competition for adsorption was more emphatic in ALVv than GARo.

4.4 Sorption Kinetics

The time dependence of metal ion's sorption on solid phases of the two soil types is shown in Figure 5. In ALVv, the sorption process was biphasic, a phenomenon attributable to sorption to sites of variable reactive and/or diffusion limited sorption (Covelo et al., 2004). The shape of the curve was linear, suggesting that the sorption process increased with time. This observation is in line with existing information that bioaccumulation

of heavy metals in terrestrial ecosystems is time dependent (Usman, 2008). The longer the period of exposure of a body of soil to heavy metals, the higher the chances of heavy metal loading beyond toxic threshold levels (Appel et al., 2008). In GARo, the fast sorption process was terminated at the 200th minute, followed by a rather progressively slow sorption as shown by the shape of plot which reached a well-defined plateau. Soil-wise comparison based on amount of metal ions adsorbed per unit time showed that ALVv attained higher metal ion adsorption capacities than GARo. The environmental implications derived from the results indicate that the sorption process in ALVv was infinite and progressive with time rendering this soil type prone to pollution. Conversely, the sorption process in GARo was terminal and significantly influenced by availability of sorption sites on solid phases. However, metal-wise comparison showed a similar adsorption preference for metal ions in both soils, thus both soils had significant affinity for Cr³⁺, Pb²⁺ and Cu²⁺ in that order.

The kinetic parameters for sorption of metal ions by the soil types of interest are shown in Table 6. Based on superiority of correlation coefficient for linearity, r^2 the pseudo-second-order rate equation was more suited to illustrate the kinetic processes of the three heavy metal ions onto the two soil types. The pseudo-second order model is based on the assumption that the rate-determining step is a chemical sorption involving valence force through sharing of electrons between adsorbent and adsorbate (Sposito, 1984; Bradl, 2004; Mohan, Singh, & Singh, 2006; Wu, Zhang, Guo, & Huang, 2008). As mentioned earlier, the kinetic sorption of the metallic ions onto solid phases was characteristically biphasic - a fast initial sorption followed by a much slower and gradual sorption. This two-step kinetic process is characteristic to the heterogeneity of the surface binding sites on sorbent, explaining that different binding sites had different binding affinities to metal ions and resulted in different binding rates (Wu et al., 2008). This assertion was corroborated by the Environmental Scanning Electron Microscope/EDAX micrographs (Figures 2 and 3) which highlighted the differences in morphology of the clay particles of the two soil types.

5. Conclusion

Sorption characteristics of selected heavy metals on soil were used as a yardstick to uncover inherent soil type variabilities amongst two semi-arid soils. The equilibrium sorption data of both soils was fittable to the Langmuir model and therefore fell short of revealing any variabilities amongst the two soil types. Nonetheless, the model was efficacious in identifying ALVv as a soil type having a higher heavy metal loading capacity potential (Q_{max}) than its counterpart (GARo). Furthermore, metal ion's affinity for the solid phases were similar for both soils and assumed the order; $Cr^{3+}>Pb^{2+}>Cu^{2+}$ and therefore inconvenient in exposing any sorption discrepancies amongst the soil types. However, preferential sorption of heavy metal onto soils was found to be driven by factors other than the expected soil type *i.e* cationic properties such as first hydrolysis constant (pK_h), oxidation state and/or cationic charge etc. Metal-wise competition for sorption sites was confirmed using CACRs and showed Cr^{3+} to have a tendency to suppress sorption of other competing entities, a feature attributable to cationic properties. When in coexistence with another competing species, Cr^{3+} suppressed sorption of Pb^{2+} and Cu^{2+} by up to half. The adsorption kinetics of metallic species onto the two soil types followed the pseudo-second order.

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