

# Adsorption of Cd, Cu, Ni and Zn in tropical soils under competitive and non-competitive systems

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**ABSTRACT:** The adsorption of heavy metals in soils affects their behavior in the environment and their bioavailability to plants. The knowledge of the adsorption mechanisms in competitive systems allows a more realistic evaluation of the metals' behavior in the soil than the single metal adsorption. The objectives of this study were (i) to evaluate Cd, Cu, Ni, and Zn adsorption in 14 surface samples (0–0.2 m) of representative soils of the Brazilian humid-tropical region, in competitive and non-competitive systems, and (ii) to establish metal affinity sequences for each soil, based in the maximum adsorption capacity (MAC) estimated by the Langmuir model. The Rhodic Eutrudox, the Kandiodalf Eutrudox, the Arenic Hapludalf, the Arenic Hapludult and the Typic Argiudoll had the highest metals' adsorption capacity, whereas the Typic Quartzipsamment and the sandy-textured Arenic Hapludult had the lowest values. In general, the MAC values for metals were lower in the competitive than in the non-competitive system. In the non-competitive system, the most common affinity sequence was Cu > Zn > Ni > Cd, whereas the most common sequence was Cu > Cd > Zn > Ni in the competitive system. In general, the Langmuir model fitted well the adsorption data of metals on the studied soils.

Key words: heavy metals, competition, tropical soil

## Adsorção de Cd, Cu, Ni e Zn em solos tropicais em sistemas competitivo e não-competitivo

**RESUMO:** A adsorção de metais pesados em solos afeta seu comportamento e biodisponibilidade às plantas. O conhecimento dos mecanismos de adsorção em sistemas competitivos permite uma avaliação mais realista do comportamento dos metais no solo do que estudos com adsorção de cada metal, isoladamente. Os objetivos desse trabalho foram: (i) avaliar a adsorção de Cd, Cu, Ni e Zn em amostras superficiais (0-0,2 m) de 14 solos representativos da região tropical úmida, em sistema competitivo e não-competitivo, e (ii) estabelecer seqüências de afinidade metálica para cada solo, com base nos valores de capacidade máxima de adsorção (CMA) dos metais estimados por meio do modelo de Langmuir. O Rhodic Eutrudox, o Kandiodalf Eutrudox, o Arenic Hapludalf (Alf2), o Arenic Hapludult (Ult2) e o Typic Argiudoll apresentaram elevadas capacidades de adsorção dos metais, ocorrendo o inverso para o Typic Quartzipsamment e para o Arenic Hapludult textura arenosa. No geral, a CMA dos metais aos solos foi menor no sistema competitivo. A seqüência de afinidade mais comumente encontrada no sistema não-competitivo foi Cu > Zn > Ni > Cd. No sistema competitivo, a seqüência foi Cu > Cd > Zn > Ni. Em geral, o modelo de Langmuir simulou de maneira satisfatória a adsorção dos metais nas amostras de solo.

Palavras-chave: metais pesados, competição, solo tropical

### Introduction

High amount of heavy metals such as Cd, Ni, Zn, and Cu in the soil can decrease crop yield due to toxicity and imply the risk of biomagnification and bioaccumulation in the food chain (Coscione et al., 2009). These metals may accumulate in the soil solid phase in several forms with specific energy levels and variable degree of reversibility of their reactions. Ion exchange, specific adsorption, organic matter complexation, precipitation and dissolution, and redox reactions define the content of bioavailable elements in the soil solution (Chang et al., 2002). Among those processes, adsorption is the most important (Silveira et al., 2002).

Detailing the chemical mechanism is essential to model plant toxicity and groundwater contamination. The interaction between ionic species in solution and the variety of soil components increase the complexity of this mechanism (Casagrande et al., 2004). In addition, these ionic species compete for the adsorption sites on the soil colloidal surfaces. Such complexity justifies the need of studies in tropical soils, because they have contrasting chemical, physical and mineralogical characteristics (Soares et al., 2005; Fontes and Alleoni, 2006).

The objectives of this study were: (i) to evaluate the adsorption of Cd (cadmium), Cu (copper), Ni (nickel), and Zn (zinc), under competitive (CS) and non-competitive system (NCS) conditions, using samples from the

surface layer of representative soils of the Brazilian humid tropical region; and (ii) to establish the sequences of preference of metallic ions for each soil, based on the values of maximum adsorption capacity for the metals, estimated by the Langmuir model.

### Material and Methods

Surface samples (0-0.2 m) from 14 non-cultivated soils were collected in areas under native vegetation or ancient reforestation in the State of São Paulo, Brazil. The natural vegetation is the evergreen Atlantic Forest, some fragments of the semideciduous forest, and remnants of the Cerrado biome (Soares and Alleoni, 2008).

Chemical characterization was performed in air-dried soil samples passed through a 10 mesh (2 mm) sieve (air-dry fine earth, ADFE). The pH was potentially determined in H<sub>2</sub>O and M KCl solution (soil:solution ratio 1:2.5), and the difference  $\Delta\text{pH} = \text{pH}_{\text{M KCl}} - \text{pH}_{\text{H}_2\text{O}}$  was used to estimate the sign of the net charge (Mekaru and Uehara, 1972). Organic carbon was quantified by oxidation with potassium dichromate in the presence of sulfuric acid, followed by titration with ammonium Fe<sup>II</sup> sulfate (Nelson and Sommers, 1996).

Cation exchange capacity (CEC) was calculated as the sum of exchangeable basic cations (Ca + Mg + K + Na), extracted by an ionic exchange resin (van Raij et al., 1986), plus the total acidity (H+Al), extracted with a 1 M calcium acetate (pH 7 buffered) solution, with further determination by titration with ammonium hydroxide (0.025 M). Ca and Mg were determined by atomic absorption spectrometry (AAS), and Na and K by flame photometry. Mineralogical analyses of the clay fraction were performed in an X-ray diffractometer, operating at 40kV and 40 mA, using Cu-K $\alpha$  radiation and scanning from 2 to 35° after their treatment with H<sub>2</sub>O<sub>2</sub> and Na-citrate-dithionite-bicarbonate (Na-cdb) to remove the organic matter and the Fe oxides, respectively.

Iron, aluminum was extracted using a 9 M H<sub>2</sub>SO<sub>4</sub> solution, and silicon was removed with NaOH from the residue of the acid attack. Contents of Fe and Al were determined using AAS, and Si was quantified by gravimetry. The Ki weathering index was calculated by  $\text{Ki} =$

$(\% \text{SiO}_2/60)/(\% \text{Al}_2\text{O}_3/102)$ . Na-cdb solution was used to extract the 'free' iron oxides (Fed) according to the method of Mehra and Jackson (1960). The hydrometer method was used for particle-size analysis after dispersion in 0.015 M (NaPO<sub>3</sub>)<sub>6</sub>.Na<sub>2</sub>O/M NaOH and overnight shaking.

Initial concentrations of Cu as Cu(NO<sub>3</sub>)<sub>2</sub>.2H<sub>2</sub>O, Ni as Ni(NO<sub>3</sub>)<sub>2</sub>.6H<sub>2</sub>O, Cd as Cd(NO<sub>3</sub>)<sub>2</sub>, and Zn as Zn(NO<sub>3</sub>)<sub>2</sub> were prepared in 20 mL of a 0.01 mol L<sup>-1</sup> NaNO<sub>3</sub> solution (1:10 ratio). Both non-competitive and competitive solutions were prepared and mixed with 2 g of the ADFE. Nitrate was chosen as the supporting electrolyte because it has a small capacity to complex metallic cations (Msaky and Calvet, 1990; Silveira et al., 2003). Concentrations were standardized by molar concentrations, suppressing the mass effect difference among the metal ions (Table 1) (Echeverría et al., 1998).

Each batch was shaken for 1 h in a horizontal shaker at 100 cycles per minute (Fontes and Gomes, 2003). After shaken, samples were centrifuged at 1,100 x g for 10 minutes and filtered. Concentration of Cd, Cu, Ni and Zn were determined by AAS.

Metal concentrations of adsorbed heavy metals were estimated by  $C_{\text{ads}} = (C_0 - C_{\text{eq}}) \text{DF}$ ; where  $C_{\text{ads}}$  is the amount of metal adsorbed onto the solid phase,  $C_0$  is the concentration of metal in the initial solution,  $C_{\text{eq}}$  is the concentration of metal in the solution after interaction with soils, and DF is the dilution factor (in the present case, 20 mL of solution to 2 g of ADFE, DF=10).

Adsorption isotherms were adjusted using the algorithm Fitfunc.bas (Barrow 1987), which adjusts non-linear equations to the results using the "minimum deviation" method (Mead, 1981; Shani et al., 1992) to calculate the maximum adsorption for each metal. In soils with high Cu adsorption, the isotherm could not be fitted to the experimental data even in the cases of small concentration in the solution (Ox1, Ox5, Oll and Alf1). In such cases, the data did not fit the model, because errors in the estimation of the maximum adsorption may reach 50% (Harter, 1984).

Values of maximum adsorption capacity (MAC) were obtained after the linearization of experimental data of each metal according with the equation  $C_{\text{eq}} / C_{\text{ads}} =$

Table 1 – Amount of Cd, Cu, Zn and Ni used for the obtaining the adsorption isotherms and its molar equivalency.

Cd	Cu	Zn	Ni	
mg L <sup>-1</sup>				mmol L <sup>-1</sup>
1.9	1.1	1.1	1.0	0.017
3.8	2.2	2.2	2.0	0.034
9.6	5.4	5.6	5.0	0.085
19.1	10.8	11.1	10.0	0.170
28.7	16.2	16.7	15.0	0.255
57.3	32.5	33.3	30.0	0.510
95.5	54.1	55.6	50.0	0.850
143.3	81.2	83.3	75.00	1.275

$(1/K \times MAC) + C_{eq} / MAC$ ; where  $C_{eq}$  is the ion concentration in the solution at equilibrium with the solid phase,  $C_{ads}$  is the amount of ion adsorbed in the solid phase,  $K$  is the affinity constant and  $MAC$  is the maximum adsorption capacity. Considering the large number of isotherms obtained in the present study, this paper presents those with contrasting adsorptive behavior.

The experiment was designed to be completely randomized. Comparison among soil samples and the preference sequences were made based on the maximum adsorption values.

## Results and Discussion

### Soil characterization

Most of the soil samples were acidic ( $pH$  in  $H_2O \leq 5.5$ ) (Table 2). All samples had a negative balance of charge ( $\Delta pH < 0$ ) probably due to the contribution of organic matter. Organic carbon (OC) content was higher than  $15 \text{ g kg}^{-1}$  for the majority of samples, except for the Arenic Hapludult (Ult1), Typic Quartzipsamment and Arenic Hapludalf. The Typic Eutraquox and the Typic Medifibrst had the highest contents of OC ( $130.7$  and  $119.1 \text{ g kg}^{-1}$ , respectively). Rhodic Eutradox, Rhodic Acrudox and Kandiudalfic Eutradox were very clayey ( $> 600 \text{ g kg}^{-1}$  clay), while Rhodic Hapludox, Anionic Acrudox, Arenic Hapludalf (Alf2), Typic Eutraquox and Typic Argiudoll were clayey ( $350\text{-}600 \text{ g kg}^{-1}$ ). On the other hand, the Arenic Hapludult (Ult1) and the Typic Quartzipsamment were sandy ( $< 150 \text{ g kg}^{-1}$  clay). The remaining samples were medium-textured ( $150\text{-}350 \text{ g kg}^{-1}$  clay) (Table 2).

### Adsorption isotherms for Cd, Cu, Ni and Zn

The adsorption curves fitted well to a Langmuir-type isotherm in both the competitive (CS) and non-competitive (NCS) systems and were divided into three sub-groups: L1 (maximum adsorption up to  $1,200 \text{ mg kg}^{-1}$  and equilibrium concentration up to  $15 \text{ mg L}^{-1}$  for the NCS; maximum adsorption up to  $1,200 \text{ mg kg}^{-1}$  and equilibrium concentration up to  $60 \text{ mg L}^{-1}$  for the CS); L2 (maximum adsorption up to  $700 \text{ mg kg}^{-1}$  and equilibrium concentration up to  $60 \text{ mg L}^{-1}$  for the NCS; maximum adsorption up to  $700 \text{ mg kg}^{-1}$  and equilibrium concentration up to  $120 \text{ mg L}^{-1}$  for the CS); and L3 (maximum adsorption up to  $400 \text{ mg kg}^{-1}$  and equilibrium concentration up to  $120 \text{ mg L}^{-1}$  for both systems (Table 3)).

### Non-competitive system

It was not possible to obtain the isotherm for Ox1 in the non-competitive system because the adsorption was exceedingly higher than the other soils. This behavior halted the linearization procedure and the determination of the maximum adsorption capacity. These findings were in agreement with the high adsorption capacity of this soil, as reported by Silveira et al. (2002) for Cu. Isotherms that had similar equilibrium concentration for different metal concentration may have had metal precipitation during the experiments (Brümmer et al., 1983). Therefore, the high  $pH$  of sample Ox1 (7.2) may have induced metal precipitation. For the remaining soils, a more contrasting behavior among metals was observed. The smaller steepness of the isotherms of these soils implies that most of the metals were adsorbed even at the smaller concentrations, decreasing as the metal concentration increased.

Table 2 – Selected chemical, physical, and mineralogical properties of 14 Brazilian superficial samples from different soil orders, collected under native vegetation.

US Taxonomy <sup>1</sup>	Symbol	Geographical coordinates	$pH_{H_2O}$	$\Delta pH^2$	CEC <sup>3</sup>	mmol $kg^{-1}$ ----- g $kg^{-1}$ -----				Qualitative mineralogy <sup>7</sup>
						OC <sup>4</sup>	Clay	Fe <sub>dep</sub> <sup>5</sup>	Ki <sup>6</sup>	
Rhodic Eutradox	Ox1	21°05'S, 47°08'W	7.3	-0.6	64.1	64.9	684	185	1.1	Kt, Gb
Rhodic Hapludox	Ox2	22°43'S, 47°38'W	4.6	-1.2	66.7	23.9	530	87	1.2	Kt
Rhodic Acrudox	Ox3	21°10'S, 47°48'W	4.7	-0.9	66.6	35.5	716	208	1.0	Kt, Gb
Anionic Acrudox	Ox4	20°10'S, 48°02'W	4.7	-1.0	53.4	25.5	470	113	0.9	Kt, Gb
Kandiudalfic Eutradox	Ox5	21°10'S, 47°48'W	5.5	-0.8	97.1	35.7	658	192	1.6	Kt, Gb
Typic Eutraquox	Ox6	22°43'S, 47°38'W	4.9	-1.1	144.0	130.7	476	5	2.1	Kt
Arenic Hapludult	Ult1	22°32'S, 47°54'W	5.3	-1.0	10.0	4.2	60	3	2.3	Kt
Arenic Hapludult	Ult2	22°38'S, 47°11'W	5.4	-0.8	61.6	19.1	247	33	2.8	Kt
Arenic Hapludalf	Alf1	22°12'S, 49°56'W	5.4	-1.1	85.9	20.7	100	5	3.0	Kt, Il
Arenic Hapludalf	Alf2	22°12'S, 49°39'W	5.8	-2.0	13.5	6.4	40	8	4.5	Kt, Il, HIV
Typic Quartzipsamment	Ent	22°32'S, 47°54'W	4.4	-0.6	14.6	4.9	80	3	2.4	Kt
Typic Medifibrst	Ist	24°43'S, 47°52'W	4.0	-0.6	124.7	119.1	275	9	2.4	Kt
Typic Dystrochept	Ept	22°22'S, 46°56'W	4.3	-0.7	39.6	17.6	243	19	1.9	Kt
Typic Argiudoll	Oll	22°07'S, 47°39'W	5.8	-1.7	153.6	28.7	543	113	2.6	Kt

<sup>1</sup>USDA Soil Taxonomy (Soil Survey Staff, 2010); <sup>2</sup> $\Delta pH = pH_{M_{KCl}} - pH_{H_2O}$ ; <sup>3</sup>CEC = cation exchange capacity; <sup>4</sup>OC = organic carbon; <sup>5</sup>Fe<sub>dep</sub> = iron content extracted by Na citrate-dithionite-bicarbonate solution; <sup>6</sup>Ki = weathering index; <sup>7</sup>Kt = kaolinite; Gb = gibbsite; Il = illite; HIV = hydroxy-interlayered vermiculite.

For all soils, the amount of adsorbed Cd, Cu, Ni and Zn increased as the concentration of the metals increased, although the rate of the increase became smaller as lesser adsorption sites were available (Petruzzelli et al. 1985; Ross 1994). Almost the whole amount of metals added were adsorbed in soils Oll, Ult2, Alf1 and Ox5, as can be seen by the steep isotherms of the group L1 (Figure 1). This behavior demonstrated the high affinity of metals by the adsorption sites of these soils.

Cu adsorption was higher than the adsorption of other metals (Figure 1, Groups L1 and L2), except for the soils of Group L3 - Arenic Hapludult (Ult1) and Typic Quartzipsamment (Ent) (Figure 1). Similar results were reported by Matos et al., (1996); Fontes et al., (2000); Gomes et al., (2001); Alumaa et al., (2001) and Silveira and Alleoni (2003) for highly-weathered tropical soils.

The maximum adsorption capacity (*MAC*) of metals was high for Oll, Ult2, Alf1, Ox5 and Ox1 (Table 4). Conversely, Ent and Ult1 had smaller *MAC*, which may be explained by the small OC contents (4.9 and 4.2 g kg<sup>-1</sup>,

respectively) and the sandy texture of these soils (Table 1). The Cd isotherm for Ent was not shown because the linearization of the isotherm was impossible. The remaining soils had intermediate values of *MAC*.

### Competitive system

For the competitive system, the amount of adsorbed metal also increased as the concentration increased (Figure 2). As the concentration increased, the competition among the metals influenced their adsorption in different degrees. Such results were similar to those reported by Basta and Tabatabai (1992) and Echeverría et al. (1998) in Haplaquolls, Hapludols, Xerochrept, Xerorthent and Haplumbrept. Fontes and Gomes (2003) evaluated the competitive adsorption of metals in samples of tropical soils (Oxisols, Ultisols and one Alfisol) and observed that Cu, Cr and Pb kept the high affinity for soil solid surfaces, while Ni, Zn and Cd were displaced from the adsorbing surfaces. Ni, Zn and Cd are more affected by the electrostatic interactions with the surface exchange

Table 3 – Soil Classification, grouped according to the adsorption behavior of the adjusted Langmuir isotherm for the competitive (CS) and non competitive (NCS) systems.

Isotherm type (group)	NCS	CS
L1		Kandiudalfic Eutrudox (Ox5) Arenic Hapludult-2 (Ult2) Arenic Hapludalf-1 (Alf1) Typic Argiudoll (Oll)
L2	Anionic Acrudox (Ox4) Typic Eutraquox (Ox6) Rhodic Hapludox (Ox2) Typic Medifibril (Ist) Arenic Hapludalf-2 (Alf2) Typic Dystrochept (Ept) Rhodic Acrudox (Ox3)	Anionic Acrudox (Ox4) Typic Eutraquox (Ox6) Rhodic Hapludox (Ox2) Typic Medifibril (Ist) Rhodic Acrudox (Ox3)
L3	Arenic Hapludult-1 (Ult1) Typic Quartzipsamment (Ent)	Arenic Hapludult-1 (Ult1) Typic Quartzipsamment (Ent) Typic Dystrochept (Ept) Arenic Hapludalf-2 (Alf2)

L1: Maximum adsorption up to 1,200 mg kg<sup>-1</sup> and equilibrium concentration up to 15 mg L<sup>-1</sup> for the NCS; maximum adsorption up to 1,200 mg kg<sup>-1</sup> and equilibrium concentration up to 60 mg L<sup>-1</sup> for the CS; L2: Maximum adsorption up to 700 mg kg<sup>-1</sup> and equilibrium concentration up to 60 mg L<sup>-1</sup> for the NCS; Maximum adsorption up to 700 mg kg<sup>-1</sup> and equilibrium concentration up to 120 mg L<sup>-1</sup> for the CS; L3: Maximum adsorption up to 400 mg kg<sup>-1</sup> and equilibrium concentration up to 120 mg L<sup>-1</sup> for both system.

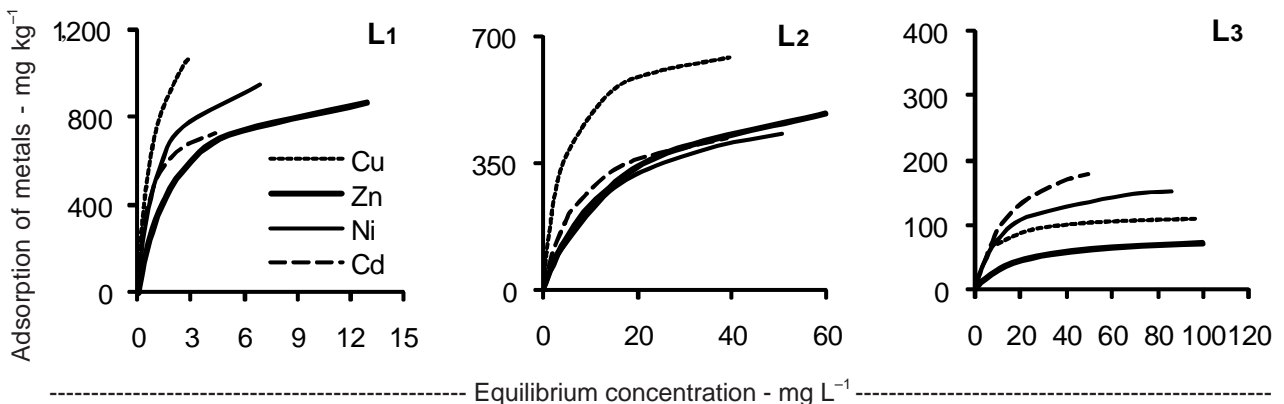


Figure 1 – Adsorption isotherms of Cd, Cu, Ni and Zn for L1, L2 and L3 groups in non competitive system (NCS).

sites, while Cu is more affected by covalent binding to mineral surfaces (McBride, 1994). Such ideas are in agreement with the findings of Matos et al. (1996), Fontes et al. (2000), Gomes et al. (2001) and Fontes and Gomes (2003).

The majority of soils revealed a decrease in *MAC* of metals in the non-competitive system, except for Cu adsorption in Ox4 and Ist, and for Cd in Ult2, Alf1, Ox5 and Oll and for (Table 4). For these soils, the competition among metals did not change the adsorption behavior but somehow increased the specific retention. The *MAC* values reported the dominance of Cu in the adsorption complex, particularly for Ox1, Ult2, Oll, Ox5, Alf1 and Ist. The high adsorption capacity for Cu into these soils implied that, besides the covalent, specific binding, Cu retention was also related to ionic binding (Silveira et al., 1999). Similar results were reported for

Cu in Oxisols, Ultisols and Alfisol (Fontes and Gomes, 2003) and other soil orders (Haplaquolls, Hapludols, Xerochrept, Xerorthent, Haplumbrept) (Basta and Tabatabai, 1992; Echeverría et al., 1998). Fontes and Gomes (2003) observed that the presence of Cu in the system decreased the adsorption of Cd, Zn and Ni; the maximum adsorption and the distribution coefficient of the metals were greater in the non competitive system as compared to the competitive system.

For the majority of soils, Cu adsorption was the one with less contrast between the two systems, indicating that the ionic competition did not affect its preferential binding to specific adsorption sites. Although Ent, Ox4, Ult1 and Ox3 had small CEC, these soils had the smallest decrease in the maximum adsorption for Cu in the competitive system. This behavior may be related to the preferential adsorption of Cu, particularly when there

Table 4 – Maximum adsorption of Zn, Ni, Cu, and Cd, estimated by fitting metal adsorption data to the Langmuir model in the non competitive (NCS) and competitive (CS) systems.

Soil	NCS				CS			
	Ni	Cu	Zn	Cd	Ni	Cu	Zn	Cd
	mg kg <sup>-1</sup> (mmol <sub>c</sub> kg <sup>-1</sup> )*							
Ox2	288.2 (4.9)	584.5 (9.2)	385.2 (5.9)	438.2 (3.9)	88.2 (1.5)	412.9 (6.5)	143.6 (2.2)	247.2 (2.2)
Ox1	-	-	-	-	452.9 (7.7)	997.4 (15.7)	1423.4 (21.8)	1999.9 (17.8)
Ox3	417.6 (7.1)	622.6 (9.8)	555.0 (8.5)	505.6 (4.5)	300.0 (5.1)	476.5 (7.5)	182.8 (2.8)	303.4 (2.7)
Ox4	347.1 (5.9)	584.5 (9.2)	417.9 (6.4)	449.4 (4.0)	111.8 (1.9)	584.5 (9.2)	156.7 (2.4)	258.4 (2.3)
Ult1	105.9 (1.8)	165.2 (2.6)	352.6 (5.4)	370.8 (3.3)	64.7 (1.1)	146.1 (2.3)	84.9 (1.3)	101.1 (0.9)
Ult2	835.3 (14.2)	2,001.2 (31.5)	999.0 (15.3)	662.9 (5.9)	400.0 (6.8)	908.5 (14.3)	398.3 (6.1)	831.4 (7.4)
Alf1	911.8 (15.5)	1,664.5 (26.2)	907.6 (13.9)	764.0 (6.8)	347.1 (5.9)	768.7 (12.1)	476.6 (7.3)	910.1 (8.1)
Ent	76.5 (1.3)	101.6 (1.6)	84.9 (1.3)	-	41.2 (0.7)	95.3 (1.5)	19.6 (0.3)	-
Ox5	835.3 (14.2)	1,245.2 (19.6)	829.2 (12.7)	764.0 (6.8)	335.3 (5.7)	832.2 (13.1)	476.6 (7.3)	910.1 (8.1)
Ox6	417.6 (7.1)	622.6 (9.8)	430.9 (6.6)	471.9 (4.2)	111.8 (1.9)	590.8 (9.3)	182.8 (2.8)	370.8 (3.3)
Ist	382.4 (6.5)	584.5 (9.2)	476.6 (7.3)	449.4 (4.0)	105.9 (1.8)	768.7 (12.1)	189.4 (2.9)	337.1 (3.0)
Ept	205.9 (3.5)	400.2 (6.3)	587.6 (9.0)	393.2 (3.5)	58.8 (1.0)	343.1 (5.4)	137.1 (2.1)	179.8 (1.6)
Alf2	258.8 (4.4)	343.1 (5.4)	666.0 (10.2)	415.7 (3.7)	88.2 (1.5)	190.6 (3.0)	117.5 (1.8)	179.8 (1.6)
Oll	711.8 (12.1)	1,111.8 (17.5)	999.0 (15.3)	764.0 (6.8)	452.9 (7.7)	832.2 (13.1)	620.3 (9.5)	1,101.1 (9.8)

(-)not linearized isotherm. \* values in parenthesis are expressed in mmol kg<sup>-1</sup>.

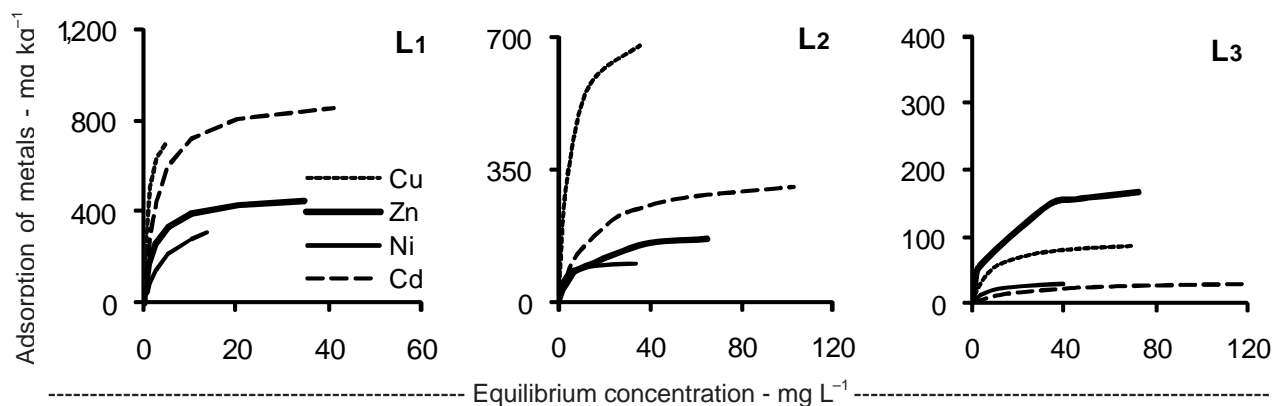


Figure 2 – Adsorption isotherms of Cd, Cu, Ni and Zn for H, L1, L2 and L3 groups in competitive system (CS).

was a small number of adsorption sites. In the soils with high CEC, the ionic competition was greater, but still the high affinity of Cu by the specific adsorption sites seemed not to be affected appreciably. As it occurred in the non-competitive system, the MAC for Cd was not estimated for the Ent soil due to the impossibility of the isotherm linearization (Table 4).

### Metal affinity sequences

The most common sequence of metal affinity in the NCS was Cu > Zn > Ni > Cd (Table 5). For the CS, the most common sequence was Cu > Cd > Zn > Ni, resulting from the preference for Cd over Zn and Ni. The sequence for the NCS was in agreement with the sequence of the negative log of the first hydrolysis dissociation constant for these metals [Cu (8.0); Zn (9.0); Ni (9.9) and Cd (10.1)]. In this case, the covalent binding was the main mechanism of metal retention, as a result of metallic hydrolysis. Similar affinity sequences, based on the hydrolysis constants, were reported for soils with high clay and iron oxide contents (Tyler and McBride, 1982; Tiller et al., 1984). The Oxisols used in this study (except Ox1) had a high affinity for all the four metals tested. Such results reinforce the role of oxides in the metal retention by highly-weathered soils. In addition to the Oxisols, the Arenic Hapludult (Ult2), the Typic Medifibril (Ist) and the Typic Argiudoll (Oll) also presented the same sequence Cu > Zn > Ni > Cd, in accordance with the hydrolysis constant of these metals.

Except for Cd, the Arenic Hapludalf (Alf2), Typic Quartzipsamment (Ent), Kandiudalfic Eutradox (Ox5) and Typic Eutraquox (Ox6) had affinity sequences fol-

lowing the “hardness” degree of metals [Cd (3.04); Cu (2.89); Ni (2.82) and Zn (2.34)]. This concept is derived from the Pearson’s hard/soft theory for acids (Pearson, 1963) and is an estimate of the degree of covalence of the chemical bonds between the metals and soil. Except for Zn, there was a relationship between electronegativity parameters [Cu (2.0); Ni (1.91); Cd (1.69) and Zn (1.65)] of metals and the sequence obtained in the NCS, which emphasizes the importance of covalent binding in metal adsorption by soils. The electronegativity indicates the tendency of an atom to attract electrons. The ionic potential (ratio between the charge and radius of an ion) is a measure of the ability of an ion to bind through electrostatic bond [Ni (5.71); Cu (5.48); Zn (5.33) and Cd (4.21)] and was also observed in the NCS in the majority of cases, except for Ni.

In the CS, there was a small variability in sequences among soils, as a consequence of metal competition. The sequence of metal affinity in the CS is in accordance with Basta and Tabatabai (1992); Fontes et al. (2000); Gomes et al. (2001); Fontes and Gomes (2003) and Agbenin and Olojo (2004).

The dominance of Cu in the majority of affinity sequences is in agreement with its hydrolysis constant, reinforcing the role of specific binding mechanisms with high stability and binding energy (James and Healy, 1972) in the behavior of this metal in soil. The sequence Zn > Ni found in most soils is also in agreement with the hydrolysis of metals.

Although soils had a greater affinity for Cu and Cd than for Zn and Ni, the sequences resulted from the CS for the four metals were in agreement with the “hardness” of these elements. Regarding the electronegativity, there was agreement only for the sequence Cu > Cd > Zn in the Rhodic Hapludox (Ox2), Arenic Hapludult (Ult2), Kandiudalfic Eutradox (Ox5), Typic Medifibril (Ist) and Typic Argiudoll (Oll). Except for Ni, these results are in agreement with those reported by Basta and Tabatabai (1992), obtained in Haplaquolls and Hapludols (Pb > Cu > Ni ≥ Cd ~ Zn).

The affinity sequences did not correlate with the ionic potential for the four metals in any soil in the CS. The poor agreement between the experimental results and those expected based on the ionic potential imply that the retention of these metals by soils is not of electrostatic nature in scenarios of high competition.

### Conclusions

Competition has affected the adsorption of all the metals, in general, it diminished the maximum adsorption. The most common sequence of affinity in the non competitive system was Cu > Zn > Ni > Cd. In the competitive system, the most common sequence was Cu > Cd > Zn > Ni. In the competitive system, Rhodic Eutradox (Ox1), Kandiudalfic Eutradox (Ox5), Arenic Hapludult (Ult2), Arenic Hapludalf (Alf2) and Typic Argiudoll (Oll) had the largest metal adsorption capacity, and the opposite was observed for Typic

Table 5 – Affinity sequences of Cd, Cu, Ni, and Zn, based on the maximum metal adsorption data for the non competitive (NCS) and competitive (CS) systems.

Soil	NCS	CS
Ox2	Cu>Zn>Ni>Cd	Cu>Cd>Zn>Ni
Ox1	Cu ≅ Zn ≅ Cd ≅ Ni*	Zn>Cd>Cu>Ni
Ox3	Cu>Zn>Ni>Cd	Cu>Ni>Zn>Cd
Ox4	Cu>Zn>Ni>Cd	Cu>Zn>Cd>Ni
Ult1	Zn>Cd>Cu>Ni	Cu>Zn>Ni>Cd
Ult2	Cu>Zn>Ni>Cd	Cu>Cd>Ni>Zn
Alf1	Cu>Ni>Zn>Cd	Cu>Cd>Zn>Ni
Ent	Cd>Cu>Ni>Zn	Cu>Zn ≅ Cd>Ni*
Ox5	Cu>Ni>Zn>Cd	Cu>Cd>Zn>Ni
Ox6	Cu>Ni>Zn>Cd	Cu>Cd>Zn>Ni
Ist	Cu>Zn>Ni>Cd	Cu>Cd>Zn>Ni
Ept	Zn>Cu>Cd ≅ Ni	Cu>Zn>Cd>Ni
Alf2	Zn>Cu>Ni>Cd	Cu>Zn>Cd>Ni
Oll	Cu>Zn>Ni>Cd	Cu>Cd>Zn>Ni

\*Affinity sequences based on the visual observation of the adsorption isotherms.

Quartzipsamment (Ent) and Arenic Hapludult (Ult1). The Langmuir model satisfactorily simulated the metal adsorption by soils in the majority of cases.

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