The Performance of Urea-Intercalated and Delaminated Kaolinites-Adsorption Kinetics Involving Copper and Lead

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Amostras brasileiras de caulinita foram usadas no estudo dos processos de intercalação com uréia (K_{UR}) e de deslaminação ($K_{UR/DL}$). A amostra intercalada foi deslaminada por tratamento com ultra-som em solução ácida. A distância basal inicial de 0,72 nm passa a 1,08 nm com aumento na área superficial de 20,3 para 90,5 m² g⁻¹ para K_{NAT} e $K_{UR/DL}$. Amostras de argila intercalada e deslaminada adsorveram cátions cobre e chumbo na interface sólido/líquido. Os modelos de isotermas de adsorção de Langmuir, Redlich-Peterson e Toth foram utilizados em processo de regressão linear e não linear para a obtenção de valores de b e K_L para todas as amostras de caulinita. A capacidade de adsorção mais alta, de 12,8 mmol g⁻¹, foi obtida com a caulinita deslaminada. Os parâmetros cinéticos analisados pelos modelos de Lagergren e Elovich forneceram um bom ajuste para uma reação de pseudo-segunda ordem com valores de k₂ variando de 5,0 a 11,0 e de 4,9 a 13,0 mmol⁻¹ min⁻¹ para cátions cobre e chumbo, respectivamente.

Brazilian kaolinite clay was used to study urea intercalation (K_{UR}) and delamination (K_{URDL}) processes. The intercalated sample was delaminated by ultrasonic treatment in acidic solution. The basal distance changed from 0.72 to 1.08 nm with increasing surface area from 20.3 to 90.5 m²g⁻¹ for K_{UR} and $K_{UR/DL}$. Suspended pristine, intercalated and delaminated clays at pH 5.0 adsorbed copper and lead cations at the solid/liquid interface. The Langmuir, Redlich-Peterson and Toth adsorption isotherm models were employed in linear and nonlinear regression processes, to give b and K_L values for all kaolinites. The highest adsorption capacity of 12.8 mmol g⁻¹ was obtained with delaminated kaolinite. The kinetic parameters analyzed by the Lagergren and Elovich models gave a good fit for a pseudo-second order reaction with k_2 in the 5.0 to 11.0 and the 4.9 to 13.0 mmol⁻¹ min⁻¹ ranges for copper and lead cations, respectively.

Keywords: kaolinite, adsorption, heavy metals, kinetics, delamination

Introduction

Some undesirable heavy metals can be removed from aqueous solution through chemical precipitation, ion exchange, electrodeposition, solvent extraction, membrane separation, reverse osmosis and adsorption processes. For this last procedure, the development of low cost adsorbents with easy manipulation and regeneration for possible reuse has been an object of significant attention, mainly when effluents are involved. Various conventional and non-conventional adsorbents have been explored for removal of different metal ions from aqueous solutions.¹⁻⁷ For example, materials containing micro and macropores, such as pillared and intercalated/delaminated clays, are often employed in such operations. In particular, kaolinite and smectite group clay minerals have been successfully explored in many adsorption procedures.⁸⁻¹⁰

From the structural viewpoint, kaolinite is a 1:1 dioctahedral aluminosilicate with two distinct basal cleavage faces.¹¹ One of them consists of a tetrahedral siloxane surface formed by very chemically inert -Si-O-Sibonds, while the other is constituted by an octahedral gibbsite sheet, Al(OH)₃. The neutral layered 1:1 structure can be disrupted and the broken bonds have the ability to accommodate OH groups.

Kaolinite is one of the most important clay minerals and its diverse industrial applications depend on the ability to modify the original properties for different uses. The methodology of surface modification includes, in principle, intercalation of species such as urea, dimethylsulfoxide and potassium acetate, among others. As expected, ion intercalation into the kaolinite structure results in changing

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the amount of reactive acidic and basic sites on the internal and external surfaces.¹²⁻¹⁸

The aim of the present investigation is to study the structural and physical-chemical characteristics of three different kaolinite forms: natural, urea-intercalated and also urea intercalated and delaminated samples. Various parameters, such as synthetic intercalation and delaminated procedures, the influence caused on adsorption capacity due to the intercalation and delamination processes, the kinetics of adsorption and applications for divalent copper and lead from aqueous solution, were explored. The Langmuir, Redlich-Peterson and Toth adsorption isotherm models have been applied to fit the experimental data by using linear and nonlinear regressions. The effect of the variation of parameters on the structure of the modified kaolinite samples was analyzed by different techniques and the kinetics of adsorption was studied through the Lagergren and Elovich models, by applying linear and nonlinear regressions.

Experimental

Raw material

The clay sample used in this investigation was obtained from the Perus area, São Paulo state, Southeast of Brazil. A natural kaolinite sample, named $K_{_{NAT}}$, with less than 2 µm particles, was separated by sedimentation. The cation-exchange capacity (CEC) was measured in order to evaluate the potential use of that clay for intercalation and delamination. The exchange was followed by the ammonium acetate method with a concentration of 2.0 mol dm⁻³ at pH 8.0, giving 0.15 mmol g⁻¹ on an airdried basis.¹¹

Sample preparation

The samples were dried at 333 K to reach humidity in the 12 to 15% range, controlled by thermogravimetry. The solid was ground and passed through a USS sieve No. 200 (0.074 mm). Part of the kaolinite clay was dispersed in doubly deionised water for several hours and was further purified using an established method.¹¹ The sample examined by powder X-ray diffractometry identified the presence of clay similar to those resulting from conventional sample preparation procedures, such as air-dried oriented mounts, ethylene glycol solvated and heated at 573 and 773 K.

Intercalation and delamination procedures

The kaolinite sample was intercalated by mixing about

10 g of the clay with 2.0 g of urea through a grinding procedure, in an agate vibratory mill for a period of 360 min at room temperature. The resulting solid, named K_{UR} , was washed with deionized water and dried for 2 h at 333 K. To perform the delamination, initially 5 g of the intercalated K_{UR} sample was suspended in 5.0 cm³ of 0.10 mmol dm⁻³ hydrochloric acid solution. This suspension was maintained for 1 h in an ultrasonic bath, Bedelin RK 1028 ultrasonic cleaner, operated at 35 kHz. After filtration through sintered glass, the solid named $K_{UR/DL}$ was dried in vacuum for 12 h at room temperature.

Adsorption

The batch adsorption methodology studies were carried out with about 60 mg of the natural, intercalated or delaminated kaolinite samples in a series of flasks containing 20.0 cm³ of aqueous solution containing the metal ion, varying in concentration from 7.0×10^{-3} to 7.0×10^{-4} mol dm⁻³ at 298 ± 1 K. Firstly, the effect of pH on adsorption for all clay samples was evaluated by varying this parameter over the range from 1.0 to 5.0, with addition of 0.10 mol dm⁻³ of nitric acid or sodium hydroxide. For these determinations, the number of moles adsorbed *per* gram (N_f) is calculated by the difference between the number of moles initially present (N_j) and the number of moles of metal remaining in the supernatant (N_s) divided by the mass (m) of the compound,¹⁹⁻²¹ using equation 1.

$$N_{f} = \frac{(N_{i} - N_{s})}{m} \tag{1}$$

The time required to reach equilibrium was obtained through specific isotherms, by considering concentration *versus* time. Equilibrium was reached in less than 10 h; however, a time period of 12 h was chosen to ensure maximum adsorption. Then, after establishing this experimental condition, the same procedure was used to obtain the isotherm with maximum adsorption at chosen pH. Based on these data the final concentration isotherm was obtained using this condition, which enabled obtaining the maxima adsorption capacity. Thus, the number of moles of cation adsorbed (N_f) increased with its concentration in the supernatant (C_s) as a function of pH during 12 h, until a *plateau* was reached, indicating total saturation of the acidic centers in the layered structure.¹⁹⁻²¹

The most commonly used isotherm is that proposed by Langmuir, which was originally derived for gas adsorption on planar surfaces such as glass, mica and platinum. However, this procedure has also been applied to cation adsorption on porous surfaces in the classical Langmuir form. For this adsorption model, the amount of the adsorbate in the equilibrium solution upon reaching the *plateau* of the isotherm enabled determination of the K_L and b parameters, which reflect the activity of the surface for the adsorbing species. The constant b establishes the upper limit for adsorption and represents the maximum value, determined by the number of reactive surface sites in kaolinite structure,¹⁹⁻²¹ as represented by equation 2:

$$N_{f} = \frac{K_{L}bC_{s}}{1+bC_{s}}$$
(2)

Then, the Cs/N_f and the so-called distribution coefficient K_L can be plotted against the concentration of the supernatant in the metal-containing solutions. If the Langmuir equation can be applied, the measured data should fall on a straight line with the slope giving K_L and the intercept K_L b values, since $1/K_L$ b is the angular and 1/b the linear coefficients,^{10,21} as represented by equation 3.

$$\left[\frac{C_{s}}{N_{f}}\right] = \left[\frac{1}{(K_{L}b)}\right] + \left[\frac{C_{s}}{b}\right]$$
(3)

Another fitting of the adsorption process was established by the Redlich-Peterson and Toth procedures as represented by equations 4 and 5, respectively:

$$N_{f} = K_{RP} bC_{S} / (1 + K_{RP} C_{S}^{\beta})$$
(4)

$$N_{f} = K_{T}bC_{S}/(1 + K_{T}C_{S}^{\beta})^{1/\beta}$$
(5)

where K_{RP} and K_T are constants related to adsorption capacity and n is an affinity constant of the adsorbent. Although equation 4 was previously employed empirically, it can be derived with the assumption of a continuous variation in thermal effect during adsorption. There is no assurance that the derivations of the Redlich-Peterson and Toth equations are unique; consequently, if the collected data can fit the equations, it is only probable, but not proven, that the surface is heterogeneous. The Redlich-Peterson and Toth models unfortunately predict both infinite adsorption at infinite concentration and a corresponding thermal effect related to the adsorption at zero coverage.

Non-linear regression was compared to obtain the optimum kinetic sorption and isothermal parameters. A trial and error procedure was employed for non-linear methods using the solver add-in functions of Microsoft Excel software. In the trial and error procedure, isotherm and kinetic parameters were determined by maximizing the coefficients of determination values.²¹ The least square method was used to analyze the linear form of the kinetic and isotherm models.

The coefficient of determination values (r^2) were used in order to find the degrees of fit of the isotherm adsorption and kinetic models with the experimental data,²² as defined by equation 6:

$$r^{2} = \frac{\Sigma (N_{fCAL} - N_{fEXP})^{2}}{\Sigma (N_{fCAL} - \bar{N}_{fEXP})^{2} + \Sigma (N_{fCAL} - N_{fEXP})^{2}}$$
(6)

where N_{fEXP} (mmol g⁻¹) is the experimental amount of divalent cation exchanged by the original and modified kaolinite samples and N_{fCAL} (mmol g⁻¹) is the amount of cation obtained by kinetic isotherm models.

Analytical techniques

X-ray powder diffraction (XRD) patterns were recorded with a Philips PW 1050 diffractometer using CuK_{α} (0.154 nm) radiation in the 2 θ region between 2 to 65° at a speed of 2° *per* min and steps of 0.05°.

The samples for infrared spectroscopy were ovendried at 393 K to remove adsorbed water. Each sample of about 1.3 mg was finely ground for 1 min, combined with 100.0 mg of oven-dried spectroscopic grade KBr and pressed with 7.0 ton into a disc under vacuum. The spectrum of each sample was recorded in triplicate between 400 and 4000 cm⁻¹ by accumulating 64 scans at 4 cm⁻¹ resolution, using a Perkin-Elmer 1760X Fourier transform infrared instrument.

BET (Brunauer-Emmett-Teller) surface areas and porosity measurements of the natural, intercalated and delaminated kaolinite samples were determined using a Quantachome/Nova Surface Area-Pore Volume Analyzer, Model 1200/5.01. The mesopore size distribution was obtained by applying the BJH (Barret-Joyner-Halenda) method to the adsorption branch of the isotherm.

The natural kaolinite (K_{NAT}) and the intercalated (K_{UR}) and intercalated-delaminated $(K_{UR/DI})$ forms were analyzed by inductively coupled plasma optical emission spectrometry (ICP OES), using a Perkin Elmer 3000 DV instrument. The oven-dried powdered samples weighing 231.0, 230.0 and 234.0 mg, respectively, were placed separately on weighed glass dishes and transferred quantitatively to pre-cleaned nitric acid digestion bottles. The samples were then digested with 7.0 cm³ of concentrated nitric and hydrochloric acids in 1:3 proportions in volume, with an identical volume of hydrofluoric acid over 5 days. The samples were cooled in an ice-bath and 25.0 cm3 of 0.10 mol dm-3 boric acid was added with stirring, followed by 50 cm³ of deionized water, and the solution was then diluted to 100.0 cm³. For each sample, a blank and a set of elemental standards were run to calibrate the instrument.

Transmission electron microscopy (TEM) images were recorded on a JEM 3010 URP microscope at the LNLS/ Brazil with an accelerating voltage of 300 kV. For this determination the samples were prepared by placing a drop of a suspension of particles dispersed in isopropanol onto a carbon-coated copper grid.

The samples for scanning electron microscopy (SEM) were prepared on orientated blades at 298 ± 1 K and sputtered with gold. The instrument used was a model LEO-ZEISS, 430 Vp, using conditions of analysis for secondary images obtained to 20 kV, with a working distance of 11 mm.

Results and Discussion

Elemental analysis

Elemental analyses from the ICP OES technique for the original clay sample, K_{NAT} , gave results consistent with kaolinite, with aluminum being the major component. The total mineralogical composition is given by 46.46; 39.42; 0.48; 0.27; 0.26; 0.22 and 0.18% of SiO₂, Al₂O₃, Fe₂O₃, MgO, Na₂O, CaO and K₂O, respectively, and 12.63% of mass was lost in the ignition process.

X-ray powder diffraction

The determination of the total mineralogical composition of the pristine natural clay sample was carried out on oriented mounts, exhibiting mainly kaolinite, quartz and goethite, besides the existence of mica in low concentration, as shown in Figure 1a, in agreement with the elemental analysis. These considerations are based on chemical analyses and characteristics X-ray diffraction (XRD) peaks of these minerals.⁴

From diffraction patterns, the d_{001} values were determined and the results are listed in Table 1. After the intercalation process the kaolinite presented, as expected, an increase in the basal distance, changing the d_{001} value from 0.72 nm for natural kaolinite to 1.08 nm for the intercalated form, as shown in Figures 1Aa and 1Ab. It was also observed that the K_{NAT} sample presents a slightly broader and less intense peak, in comparison with K_{UR} , a behavior that could be attributed to differences in the degree of crystallinity, which was calculated by using the Biscaye equation,⁸ as listed in Table 1, due to the effect of urea insertion, to give 0.90 and 0.88 for the natural and intercalated forms. The delaminated clay sample gave characteristics of a collapsed structure and low crystallinity, as shown in Figure 1Ac, which suggested the clear influence of the delamination process on the kaolinite structure. The natural kaolinite and its intercalated form, in expanded scale in part B of Figure 1Bb, presents the new peak with d_{001} at 1.27 nm that is attributed to urea interaction



Figure 1. A: X-ray diffraction patterns for samples of (a) natural kaolinite (C), goethite (G), quartz (Q) and mica (M), (b) K_{UR} and (c) K_{URDL} . B: Detail of patterns for natural kaolinite and K_{UR} from 4 to 11° for (a) and (b).

Table 1. Basal spacing (d_{001}) , surface area (S), micropore area (Ma), crystallinity (Cr), pore volume (P) and pore diameter (PD) for natural (K_{NAT}), intercalated (K_{UR}) and delaminated (K_{URDL}) kaolinite samples

Sample	d ₀₀₁ / nm	$S / (m^2 g^{-1})$	Ma / (m ² g ⁻¹)	Cr	P / (cm ³ g ⁻¹)	PD / nm
K	0.72	20.3	8.21	0.90	3.54	0.12
K _{UR}	1.08	44.5	9.11	0.88	3.90	0.20
K _{UR/DL}	—	90.5	22.98		8.75	0.75

inside the kaolinite layer. When hydrated kaolinite was intercalated with 20% urea and washed with water under ultrasound, a distance of 0.84 nm was obtained.¹⁴ The d_{001} values for the delaminated kaolinite are in agreement with previous results.^{15,16}

Infrared spectroscopy

The Fourier-transform infrared spectra of the natural kaolinite K_{NAT} , intercalated K_{UR} and the delaminated form $K_{UR/DL}$ are shown in Figure 2. The original K_{NAT} sample exhibited all characteristic hydroxyl stretching bands⁵ attributed to the inner-surface hydroxyls oriented towards the interlayer at 3695 cm⁻¹. The bands at 3622 and 3463 cm⁻¹ are assigned to those hydroxyl groups oriented towards the vacant sites in the external layers of the kaolinite structure.⁶ Additional weak, broad bands of kaolinite at 3463 cm⁻¹ and another weak one at 1639 cm⁻¹ can be associated with water adsorbed on the external surface.¹⁴



Figure 2. FTIR spectra of the kaolinite samples, $K_{\rm UR}$ (a), $K_{\rm NAT}$ (b) and $K_{\rm UR/DL}$ (c).

The spectrum of the intercalated material presented a similar set of frequencies for hydroxyl vibrations as does natural K_{NAT} , which is in agreement with other kaoliniteurea complexes.¹⁴ The bands located at 3505, 1683 and 1463 cm⁻¹ were assigned to N-H, C=O and C-N stretching vibrations, respectively. The band at 3505 cm⁻¹ is attributed to the formation of a hydrogen bond between the NH₂ urea group and oxygen atoms from the basal tetrahedral sheet.¹⁷ The intercalated K_{UR} displayed characteristic hydroxyl stretching bands at 3670 and 3650 cm⁻¹, but with reduced intensity and strongly broadened. This behavior may be attributed to the loss of hydrogen bonding of the outer hydroxyls between the layers. In addition, there are two new sets of weak bands: (*i*) the formation of hydrogen bonds between NH₂ urea groups and O-Si-O groups of kaolinite in the 3381 to 3505 cm⁻¹ interval and (*ii*) the symmetric and asymmetric vibrational modes of NH₂ urea groups that interact with oxygen atoms of the Si₂O₅ layers in the 3204 to 3413 cm⁻¹ region.¹⁷ The C=O stretching vibration of the urea molecule is also observed at 1683 cm⁻¹ for the intercalated urea-kaolinite, as observed before.¹⁸

Textural analysis

The gaseous nitrogen adsorption values for natural, intercalated and delaminated kaolinite samples are listed in Table 1. The nitrogen BET data was used to calculate the specific surface areas for comparative purposes. The BET surface areas of the kaolinite samples demonstrated that intercalation and delamination processes caused the formation of mesopores in the solid particles, resulting in a higher surface area, 90.5 m² g⁻¹, relative to the natural sample, K_{NAT} , with 20.3 m² g⁻¹. The pore size distribution in the mesopores region was obtained by applying the BJH method from the nitrogen isotherms at 77 K. The pore size distribution was compared by considering natural and intercalated samples, as shown in Figure 3. The natural kaolinite sample exhibited a pore volume of 3.54 cm³ g⁻¹, while the intercalated and delaminated kaolinites gave values of 3.90 cm³ g⁻¹ and 8.75 cm³ g⁻¹ respectively. The K_{UR} and $K_{UR/DL}$ samples presented a unimodal distribution of pore sizes, while $\mathbf{K}_{_{\rm NAT}}$ has a bimodal distribution. The present values of surface areas are in agreement with those reported before in the 5 to 25 m²g⁻¹ interval for kaolinite,²⁴ which depends on particle size distribution, particle shape, and distribution of cracks and pores in the material, and therefore, cannot be represented as a general characteristic of a particular type of material.25



Figure 3. Pore size distribution curves of natural and modified kaolinite samples: $K_{NAT}(\blacksquare)$, $K_{UR}(\bigcirc)$ and $K_{URDL}(\blacktriangle)$.

Microscopy

The TEM and SEM images were obtained in parallel position related to the layers, in 010 or 100 planes, as shown in Figures 4 and 5, respectively. The images for natural and delaminated kaolinites in the 001 plane are shown in Figures 4a and 4b, respectively. The individual crystals presented themselves as flakes in a pseudo-hexagonal form, as confirmed by TEM images, and the delaminated kaolinite structure is shown in Figure 4b. The reduction of particle size after the delamination process is shown in the TEM image, where the separation and disintegration into small stacks of flakes is clear, in comparison with Figure 4a, which could be attributed to the use of ultrasonic procedure.

SEM images were used to probe the changes in the morphological features of K_{NAT} and $K_{UR/DL}$ samples, as shown in Figures 5a and 5b, respectively. The surface morphology of a natural sample presents corn-flake like crystals with a cottony appearance, revealing its extremely fine plate-like structure, Figure 5a. However, after intercalation, the clay becomes more porous and



Figure 4. TEM of natural- K_{NAT} (a) and modified kaolinite- $K_{UR/DL}$ (b) samples.

fluffy, Figure 5b. This appearance is probably due to the change in the surface charge of the particle, as a result of the intercalation process, and the reduction in amorphous phase originally associated with the natural clay sample. However, both samples present thick stacks with thin single flakes, as are observed for other chemically modified kaolinites. This texture is not consistent with the propensity of $K_{UR/DL}$ to convert spontaneously into individual flakes or small stacks of pseudo-hexagonal flakes as shown in Figure 5b.

Effect of pH

The pH of the aqueous solution, which increased almost linearly up to pH 5.0, is an important controlling parameter in the adsorption process. Both the extent of adsorption and the amount adsorbed (N_f) showed positive changes. The influence of the pH on the concentration adsorbed for natural and modified kaolinites is shown in Figure 6, demonstrating that the cation exchanges strongly as the acidity of the solution decreases. The adsorption experiments at pH values higher that 5.0 were not carried out due to the risks of cation hydrolysis.²¹





Figure 5. SEM of natural- $K_{_{NAT}}$ (a) and modified kaolinite- $K_{_{UR/DL}}$ (b) samples.





Figure 6. Effect of pH on cation adsorption from aqueous solution for 2.0 mg dm⁻³ clay, with copper(II) at 12.08 mg dm⁻³ and lead(II) at 14.62 mg dm⁻³, at 298 ± 1 K for: $K_{UR/DL}$ (a) and K_{NAT} (b) with Cu (\bigcirc) and Pb (\blacktriangle).

For a strongly acidic medium, the proton concentration outnumbers the lead ions in the adsorptive solution. Thus, and it is expected that, at low pH, the adsorptive sites will be covered by protons. By increasing the pH, the surface sites become free for adsorption of the various lead species. Similar observations have already been reported for lead in natural and chemically modified clay adsorption processes.¹⁹

The number of available hydrogen ions is high at low pH values and copper cations must compete with them to exchange all adsorption sites on the inorganic surface. These active sites on the adsorbent surface are gradually deprotonated with the increase in pH, favoring copper uptake, with an adsorption mechanism similar to that previously reported.¹⁻³

Cation adsorption

Adsorption processes involve an array of phenomena that can alter the distribution of undesirable agents on the constituent phases and interfaces of a given system.²⁰ Thus,

Table 2. The maximum adsorption capacity, N_f^{max} , for each metal on natural, K_{NAT} and modified kaolinites, K_{IIR} , $K_{IIR/DL}$

M ²⁺	K _{NAT}	K _{UR}	K _{ur/dl}
Cu ²⁺	5.0 ± 0.2	7.5 ± 0.1	12.3 ± 0.2
Pb ²⁺	5.6 ± 0.2	11.3 ± 0.2	12.8 ± 0.2

Clay, 3.0 g dm⁻³; initial Cu²⁺, 12.08 g dm⁻³; initial Pb²⁺, 14.62 g dm⁻³; controlled temperature, 298 \pm 1 K; pH = 5.0.

adsorption at a surface or interface is mainly a result of binding forces between the individual atoms, molecules or ions of the adsorbate to the surface, all of these forces having some electrostatic effects to promote interaction.²¹ Divalent cations (M^{2+}) can be adsorbed on the available surface sites, denoted by \equiv SOH, where S represents silicon or aluminum atoms in the inorganic structure, these atoms being main elements in the clay composition, as listed in Table 2. For this process, the reactive edge sites (OH and NH₂ groups) on the surface can interact with cations in solution, as given by equations 7 to 9:

$$\equiv SOH + M^{2+} \iff \equiv SOM^{+} + H^{+}, \tag{7}$$

$$\equiv SOH + M^{2+} + H_2O \implies \equiv SOMOH + 2H^+, \tag{8}$$

$$2 \equiv \text{SOH} + \text{M}^{2+} \iff \equiv \text{SO}_2\text{M} + 2\text{H}^+.$$
(9)

On the surface sites the original \equiv SOH group can exchange with cations (M²⁺), as represented in equations 10 and 11:

$$2\equiv SOH + M^{2+} \iff \equiv SO_2 M + 2H^+, \tag{10}$$

$$\equiv SOH + M^{2+} + H_2O \iff \equiv SOMOH + 2H^+.$$
(11)

During the course of the adsorption the inner-sphere interactive process is predominant over that of the outer-sphere for divalent metal cations adsorbed by edge sites.²⁰ Both cations presented similar isotherms with kaolinite and its intercalated and delaminated forms, as exemplified for copper in Figure 7 and lead in Figure 8. As observed, for the matrix containing urea, the adsorption amounts are higher at the saturation isotherm *plateau*. The maximum adsorption capacities, N_f^{max} , for each metal on natural and modified kaolinites are listed in Table 2. The adsorptions are higher for modified matrices.

These results were adjusted to fit the Langmuir, Redlich-Peterson and Toth models, as listed in Table 3, and the linear and nonlinear correlation adjustments from these equations are shown in Figure 9. All three adsorption models can be used to explain the significant adsorptive capacity of the intercalated matrix and to quantify copper and lead



Figure 7. Experimental data for copper adsorption by kaolinite samples, using 2.0 g dm⁻³ clay; copper(II) concentration 12.08 mg dm⁻³; 298 ±1K at pH 5.0 for K_{NAT} (\blacksquare), K_{UR} (\bigcirc) and $K_{UR/DL}$ (\blacktriangle).



Figure 8. Experimental data for lead adsorption by kaolinite samples, using 2.0 g dm³ clay; lead(II) concentration 12.08 mg dm³ at pH 5.0; 298 ± 1 K for K_{NAT} (\blacksquare), K_{IIR} (\bigcirc) and $K_{IIR/DL}$ (\blacktriangle).

interactions with these inorganic matrices, although the Langmuir model presents a significant advantage in comparison with the others. It allows the quantification of the capacity of cations in the structure of the matrix and the evaluation of the constant related to the binding energy.

The interaction of these cations with the intercalated surfaces of the kaolinite structure is governed by the microenvironment around each arrangement, which is usually composed of basic centers, mainly in the internal structure of hydration. Hydroxyl groups act as bridges that result in the kaolinite matrix intercalation process.

Kinetic processes

Pseudo-first order kinetics using the Lagergren equation²⁶⁻³⁸ is generally expressed by equation 12:

$$\frac{\partial N_{t}}{\partial t} = k_{1} (N_{fEQ} - N_{f})$$
(12)

After integration and applying the boundary conditions, $N_t = 0$ for t = 0 and $N_t = N_t$ at t = t, the integrated form of equation 12 becomes:

$$\ln \left(N_{fEQ} - N_{f} \right) = \ln N_{fEQ} - k_{1}t$$
(13)

where N_{fEQ} and N_f are the amounts of metal adsorbed at equilibrium and at a given time t (mmol g⁻¹), respectively, and k_1 is the rate constant for pseudo-first order adsorption (min⁻¹).

When the rate of reaction of an adsorption reaction is controlled by chemical exchange, then a pseudo-second order model can be better adjusted to the experimental kinetic data,²⁶⁻³⁹ as expressed by equation 14.

$$\frac{\partial N_{f}}{\partial t} = k_{2} (N_{fEQ} - N_{f})^{2}$$
(14)

After integration and applying the boundary conditions, $N_t = 0$ for t = 0 and $N_t = N_t$ at t = t, the integrated form of equation 14 becomes:

$$\frac{\mathbf{t}}{\mathbf{N}_{\rm f}} = \left(\frac{1}{\mathbf{k}_2 \mathbf{N}_{\rm fEQ}^2}\right) + \left(\frac{1}{\mathbf{N}_{\rm fEQ}}\right) \mathbf{t} \tag{15}$$

where k_2 is the pseudo-second order rate (mmol⁻¹ min⁻¹). The values of k_2 can be obtained from the *y*-intercept of the linear plot of t / N_f versus t.

Carrying out a set of experiments at constant temperature and monitoring the amount adsorbed with time, the kinetics of the adsorption process should be known. The adsorption using the Lagergren model can also be explored,²⁶⁻³⁸ as proposed in equation 12.

The useful Elovich equation for energetically heterogeneous solid surfaces is represented by equations 16 and 17 in nonlinear and linear¹ forms, respectively:

$$N_{f} = \ln(\alpha\beta t)^{\beta} \tag{16}$$

$$N_{f} = \beta \ln(\alpha\beta) + \beta \ln t \tag{17}$$

where α and β , the Elovich coefficients, represent the initial adsorption rate (mmol g⁻¹ min⁻¹) and the desorption coefficient (g mmol⁻¹), respectively.

The modified and natural kaolinites showed identical behavior towards copper and lead uptake with increasing interaction time at pH 5.0. Adsorption increased rapidly up to 1 h and slowly increased as equilibrium was reached, as shown in Figure 10. The lead uptake became almost

Adsorbed	Langmuir			Redlich-Peterson			Toth		
M ²⁺	b	K	r	K _{RP}	β	r	Ν	K _T	r
				Linear					
				K _{NAT}					
Cu	6.2 ± 0.1	0.8 ± 0.3	0.988	2.2 ± 0.2	2.3 ± 0.1	0.992	1.9 ± 0.1	3.7 ± 0.2	0.997
Pb	6.3 ± 0.2	0.9 ± 0.1	0.999	2.0 ± 0.3	2.3 ± 0.3	0.998	1.8 ± 0.3	3.3 ± 0.3	0.996
				K _{ur}					
Cu	8.5 ± 0.2	1.0 ± 0.2	0.999	3.169	3.0 ± 0.5	0.985	2.1 ± 0.2	4.3 ± 0.1	0.997
Pb	9.6 ± 0.2	1.1 ± 0.3	0.994	3.651	3.9 ± 0.3	0.987	2.0 ± 0.2	4.2 ± 0.2	0.996
				$K_{UR/DL}$					
Cu	9.6 ± 0.2	1.0 ± 0.3	0.999	3.6 ± 0.1	3.1 ± 0.5	0.987	3.7 ± 0.1	5.6 ± 0.3	0.998
Pb	9.9 ± 0.2	1.2 ± 0.2	0.998	4.0 ± 0.2	4.1 ± 0.2	0.998	4.2 ± 0.3	5.9 ± 0.2	0.998
				Nonlinear					
				K _{NAT}					
Cu	5.9 ± 0.3	0.9 ± 0.1	0.988	3.0 ± 0.2	2.4 ± 0.2	0.992	1.9 ± 0.1	3.7 ± 0.2	0.998
Pb	6.6 ± 0.1	0.8 ± 0.7	0.999	3.4 ± 0.4	2.3 ± 0.1	0.998	1.8 ± 0.5	3.3 ± 0.5	0.997
				K _{ur}					
Cu	9.3 ± 0.3	0.9 ± 0.1	0.999	3.8 ± 0.2	2.0 ± 0.2	0.986	2.1 ± 0.1	4.3 ± 0.3	0.999
Pb	1.0 ± 0.1	1.2 ± 0.2	0.997	3.7 ± 0.1	3.9 ± 0.1	0.989	2.1 ± 0.3	4.3 ± 0.2	0.998
				$K_{UR/DL}$					
Cu	9.7 ± 0.3	0.942	0.999	3.7 ± 0.2	3.2 ± 0.3	0.998	3.6 ± 0.2	5.5 ± 0.1	0.999
Pb	1.1 ± 0.1	1.2 ± 0.3	0.999	4.1 ± 0.2	4.1 ± 0.1	0.999	4.0 ± 0.1	6.0 ± 0.2	0.998

Table 3. Parameters from Langmuir, Redlich-Peterson and Toth equations for metal adsorption on natural, intercalated and delaminated kaolinite samples; linear and nonlinear forms

Clay, 3.0 g dm⁻³; initial Cu²⁺, 12.08 g dm⁻³; initial Pb²⁺, 14.62 g dm⁻³; controlled temperature, 298 \pm 1 K; pH = 5.0.



Figure 9. Experimental data fitting of lead adsorption in K_{UR} applied to linear (a) and nonlinear of copper adsorption in $K_{UR/DL}$ (b) models: Langmuir (\blacktriangle), Redlich-Peterson ($\textcircled{\bullet}$), Toth (\blacksquare) models and experimental (V) for 2.0 g dm⁻³ clay. Lead and copper concentrations 14.62 and 12.08 mg dm⁻³, respectively, at pH 5.0.

constant after 200 min for natural and after 150 min for intercalated kaolinite samples, which can be considered reaching the equilibrium condition. The initial high rate of adsorption can be attributed to the existence the exposed



Figure 10. Adsorption isotherms of K_{UR} and K_{NAT} with divalent cations for 2.0 g dm⁻³ clay, copper at 12.08 and lead at 14.62 mg dm⁻³, 298 ± 1 K, pH 5.0, for K_{NAT} . Cu (\Box), Pb (\blacktriangle) and K_{UR} : Cu (\bigcirc), Pb (\blacktriangledown).

basic centers for interaction on the readily accessible surfaces. However, as the coverage increased, the number of available surface sites for adsorption decreases until it reaches equilibrium, when the uptake is controlled by the rate at which the adsorbate is transported from the exterior to the interior sites of the adsorbent particles.

Second order kinetic plots presented better linearity with R > 0.99, and the second order rate constant, k_2 , varied from 4.8 to 10.9 and from 4.9 to 12.9 g mg⁻¹ min⁻¹ with the linear method and from 5.0 to 11.0 and 4.9 to 13.0 g mg⁻¹ min⁻¹ with the nonlinear method for copper and lead cations, respectively, as listed in Table 4. A comparison of N_f experimental values and those obtained from the slopes of the second order plots showed values in the 11.40 to 2.5 mmol g⁻¹ range for copper with kaolinite and lead for intercalated urea matrices with linear and nonlinear analyses, respectively, as shown in Figures 11 and 12. Considering all the above results, the kinetics of second order for cation adsorption on natural and modified kaolinites is better adjusted for both metals.¹

Second order kinetics suggests that the number of adsorption sites on the clay surface and the number of

Sample	Pseudo-first order			Pseudo-second order			Elovich			
	k ₁ x 10 ² / min ⁻¹	${f N_{fcal}}/{(mmol \ g^{-1})}$	r ²	k ₂ x 10 ³ / (mmol g ⁻¹ min ⁻¹)	N _{fcal} / (mmol g ⁻¹)	r ²	$\propto x \ 10^3$ / (g mmol ⁻¹ mim ²)	β/ (mmol g ⁻¹ mim ⁻¹)	r ²	
				Line	ear					
				copper	r (II)					
K _{NAT}	1.0 ± 0.1	4.2 ± 0.3	0.978	4.8 ± 0.1	6.1 ± 0.2	0.998	9.7 ± 0.2	1.5 ± 0.2	0.981	
K _{ur}	1.1 ± 0.2	6.2 ± 0.2	0.977	8.9 ± 0.3	8.7 ± 0.1	0.999	17.4 ± 0.2	1.8 ± 0.1	0.975	
K _{UR/DL}	1.8 ± 0.2	7.2 ± 0.3	0.999	10.9 ± 0.2	9.1 ± 0.1	0.998	23.0 ± 0.1	2.2 ± 0.2	0.998	
				lead	(II)					
K _{NAT}	1.2 ± 0.5	4.7 ± 0.5	0.976	4.9 ± 0.2	6.3 ± 0.2	0.997	$5,3 \pm 0.2$	1.2 ± 0.1	0.986	
K _{ur}	1.2 ± 0.1	7.8 ± 0.1	0.977	11.8 ± 0.2	9.9 ± 0.1	0.998	18.2 ± 0.2	1.6 ± 0.2	0.976	
K _{UR/DL}	2.0 ± 0.5	9.8 ± 0.1	0.998	12.9 ± 0.2	10.0 ± 0.1	0.998	25.3 ± 0.1	2.8 ± 0.1	0.998	
				Nonli	near					
copper(II)										
K _{NAT}	1.2 ± 0.2	4.1 ± 0.2	0.999	5.0 ± 0.2	6.1 ± 0.1	0.999	9.8 ± 0.2	1.5 ± 0.2	0.998	
K _{ur}	1.1 ± 0.2	6.2 ± 0.2	0.998	8.8 ± 0.2	8.3 ± 0.2	0.999	18.0 ± 0.2	1.9 ± 0.2	0.998	
K _{UR/DL}	2.1 ± 0.2	8.9 ± 0.5	0.999	11.0 ± 0.1	9.1 ± 0.1	0.998	23.1 ± 0.3	2.3 ± 0.2	0.999	
				lead	(II)					
K _{NAT}	1.2 ± 0.1	4.7 ± 0.2	0.976	4.9 ± 0.2	6.3 ± 0.4	0.999	$5,3 \pm 0.2$	1.2 ± 0.1	0.986	
K _{UR}	1.4 ± 0.2	7.8 ± 0.3	0.997	11.8 ± 0.2	10.0 ± 0.2	0.998	18.2 ± 0.1	1.6 ± 0.3	0.996	
K _{UR/DL}	2.1 ± 0.5	10.2 ± 0.4	0.989	13.0 ± 0.2	11.0 ± 0.2	0.999	25.5 ± 0.3	2.7 ± 0.1	0.999	

Table 4. Kinetics for pseudo-first order and pseudo-second order Lagergren and Elovich models calculated for metal adsorptions onto natural, intercalated and delaminated kaolinite samples

Clay, 3.0 g dm⁻³; initial Cu²⁺, 12.08 g dm⁻³; initial Pb²⁺, 14.62 g dm⁻³; controlled temperature, 298 \pm 1 K; pH = 5.0; time, 360 min.

heavy metal cations in the liquid phase together determine the kinetics. Depending on pH, different cationic species may be held to the clay surface at appropriate ion-exchange sites.²⁷



Figure 11. Linearization of adsorption isotherms obtained by kinetic models: (a) pseudo-second order; (b) pseudo-first order and (c) Elovich equation. Clay: 2.0 g dm⁻³; divalent copper at 12.08 and lead at 14.62 mg dm⁻³; 298 ± 1 K, pH 5.0, time 360 min. K_{NAT}: Cu (□), Pb (▲); K_{UR}: Cu (○), Pb (▼).



Figure 12. The experimental fitting of lead adsorption data applied to nonlinear models: Elovich (\bullet), Lagergren pseudo-first order (\blacktriangle), pseudo-second order models (\blacksquare) and experimental (—) for 2.0 g dm⁻³ clay. Lead concentration 14.62 mg dm⁻³ at pH 5.0.

Conclusions

Synthesized urea-intercalated kaolinite, originally from the Perus region, Brazil, has high stability, into which active basic sites can be introduced, improving the physicalchemical properties of the precursor clay. For example, the specific area of the original kaolinite, 20.3 m² g⁻¹, changed after intercalation and delamination to 44.5 and 90.5 m² g⁻¹, respectively. Identically, the modified clays, when applied to the metal adsorption process, increased their capacities for divalent lead and copper removal, a favorable condition attributed to the available edge sites on the inorganic backbone, to give the order Pb > Cu.

Adsorption either on natural or intercalated clays increased continuously with pH, by limiting the hydrolysis of both cations and NH_2 anchored in the kaolinite surface. The most appropriate condition was chosen to be pH 5.0 at room temperature, presenting a *plateau* at 200 and 150 min for natural and modified kaolinites, respectively. The copper and lead cations showed very rapid initial uptakes on both inorganic supports before reaching the *plateau*. Langmuir, Redlich-Peterson and Toth isotherms yielded good fits with the adsorption data for cation/clay interactions.

The kinetics of adsorption of both heavy metals used in this investigation on natural, intercalated and delaminated kaolinites were complex, but it seems that this interactive effect is better expressed by a pseudosecond order process. Linear and nonlinear regression methods were compared to evaluate the experimental data with a pseudo-second order kinetic model. Non-linear regression exhibited a higher coefficient of determination for the isotherm and the kinetic analyses than a linear regression.

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