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Organophilic bentonites obtained by microwave heating as adsorbents for anionic dyes

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Abstract

Organobentonites were prepared using three cationic surfactants tetradecyl- (C14), hexadecyl- (C16) and octadecyltrimethylammonium (C18) bromides in the proportions of 100% and 200% of the cationic exchange capacity (CEC) at 323 K under microwave irradiation for 5 min. The XRD patterns indicated the formation of intercalated hybrids with d values in the range 1.83 to 2.00 nm. CHN elemental analysis showed better organofunctionalization of the clay matrix for the organobentonites obtained with 200% of the CEC and longer organic alkyl chains and agreed with thermogravimetric analysis. Infrared spectroscopy suggested the presence of the organic moieties on organobentonites, showing typical absorptions of CH2 and CH3 groups. The solids were applied for Remazol blue RN dye removal from a single and synthetic effluent sample. The influences of the pH, adsorbent dosage, contact time and initial dye concentration were systematically investigated. The data were fitted to different kinetic and equilibrium models. All of the adsorption processes were adjusted to pseudo-second order and Langmuir models. The organobentonites showed good performance in dye removal at a pH of 2 for the solids obtained with 100% of the CEC, and the dye removal was independent of the pH for the other solids, which is in agreement with the zeta potential measurements. The optimal reaction time was 60 min, except for Bent-C16-200%. The best dye removal efficiency (100%-211 mg g⁻¹) for the Bent-C14-200% sample with 500 mg/L of dye concentration.

Keywords Bentonite; organoclay; anionic dyes; adsorption

1. Introduction

Dyestuffs are important pollutants and many of them are non-biodegradable, being widely used in some industries, such as textiles, leather, pharmaceuticals, paper, among others (Tangaraj et al., 2017, Kausar et al., 2018). Reactive anionic dyes, such as Remazol blue RN. disodium 1-amino-9,10-dioxo-4-[3-(2sulfonatooxyethylsulfonyl)anthracene-2-sulfonate, have anthraquinone as a chromophore. They are toxic and recalcitrant dyes, and they represent approximately 30% of the paint production in the world. Their environmental importance is that 10 to 50% of them can remain in solution after the dyeing process (Janaki et al. 2012). The discharge of the effluents containing dyes in aquatic environments damage the ecological balance and living organisms by the influence of the chemical groups present in the dye structure (Torgut et al., 2017). However, the major environmental concern due the presence of dye in aquatic medium is the absorption and reflection of sunlight, which interfere in the growth of the bacteria to levels insufficient to biologically degradation of the impurities in the water resulting in ecological imbalance (Unuabonah et al., 2008).

Several processes for the removal and/or treatment of dyes have been proposed, such as filtration, chemical treatment, advanced oxidative processes, biodegradation, membrane separation, and electrochemical methods (Gupta et al., 2009; Rehman et al., 2014). However, adsorption is the most widely used method, since adsorption is easy to operate and has the possibility of selectivity as well as the reuse of the adsorbent (Huang et al., 2014). Some adsorbents often present high costs of production and regeneration (for example, carbon nanotubes, metal-organic frameworks (MOFs) and ion exchange membranes), which may make them economically unfeasible. Therefore, natural adsorbents with a low cost, a large surface area and/or active sites on the surface

have been extensively explored (Ngulube et al., 2017). Bentonite is a natural mineral composed of at least 50% smectite, and more precisely montmorillonite, a 2:1 phyllosilicate (Brigatti et al., 2013), and meet all of these requirements. Montmorillonite is constituted of an aluminum octahedron sheet sandwiched between two silicon tetrahedron sheets. Isomorphic substitution of Al³⁺ by Mg²⁺ in the octahedral sheet gives an overall negative charge, which is counterbalanced by cations located in the interlayer space (Bergaya et al., 2011; Jaber, et.al., 2014). Furthermore, bentonites are abundant, easy to handle, biocompatible, and non-toxic (Zhu et al. 2015) and presents other properties as thermal stability, nanoscale porous structure, appropriate surface morphology, chemical stability (Fatimah et al., 2011; Hassane et al., 2015; Sahin et al., 2015).

Due to their negatively charged surface, bentonites have a low affinity for anionic species like reactive dyes such as Remazol textile dyes (Silva, et al., 2012; Pereira et al., 2017); therefore the use of organobentonites is an alternative way, because the organic cations inserted in the clay minerals give better affinity for organic species dissolved in solutions (Teppen et al., 2007). The net charge of the solid is changed and made more easily the interaction with negatively charged species (Ma et al., 2011; Tunç et al., 2012). Organobentonites have been applied for dye removal from aqueous solutions (Ma et al., 2011; Fan et al., 2014; Yan et al., 2015; Anirudhan et al., 2015; Tangaraj et al., 2017; Wang et al., 2017).

The synthesis of organobentonites usually involves an ion exchange reaction between organic alkyl quaternary ammonium salts and the interlayer cations (Paiva et al., 2008; Chiu et al., 2014). Generally, ammonium cations with long linear alkyl chains or alkyl phosphonium surfactants could be intercalated into the silicate interlayer gallery to enlarge the basal spacing (Bergaya et al., 2011). Synthesis of organoclays has been well-described in aqueous solutions at room temperature or at moderate temperatures such as 40-50°C (Paiva et al 2008) over a range of times (hours or days).

The hydrophobic nature of the organobentonites results in their use for the sorption of organopollutants. Lagaly et al., (1970, 1971) proposed that organoclays prepared with alkylammonium cations present different arrangements of the organic moieties in the interlayer spacing such as a lateral monolayer, a bilayer parallel to the silicate layers, paraffin-like monomolecular or paraffin-like bimolecular arrangements. More recently, some studies indicated that the properties of the organically-modified bentonites depend on their cation exchange capacity (CEC) and the surfactant structure, that is, the nature of the head group, number of chains per head group and chain length (Heinz et al., 2005; 2007). These studies based on experimental and molecular simulations showed that in low charged montmorillonite, the formation of partially and densely packed alkyl layers can be observed. Thus, when alkylammonium surfactants with C_{10} - C_{22} chains were used, a partial formation of an alkyl bilayer was observed for C_{10} , a low interlayer density of the alkyl bilayer was observed for C_{14} and a densely packed bilayer was observed for a C_{18} chain length. The formation of a partial alkyltrilayer occurred for a chain length of C_{22} (Fu and Heinz, 2010).

In the present study, organobentonites were prepared by the reaction between ammonium salts and bentonites under microwave heating. Currently, microwave chemistry has received remarkable attention, owing to its powerful thermal effect that helps in the completion of the reactions in the shortest time by means of rapid and uniform heating, because the use of microwave radiation targets mainly to carbon-based material with good microwave absorbency (Kar et al., 2016). When the carbon material is exposed to microwave radiation, dipole rotation in atomic scale is occurred up to million times per second within the material. As a result, heat energy is rapidly generated from the frictions between the atoms and molecules within the material which subsequently leads to fast heating of the material (Lam et. al., 2016; Liew et al., 2018). Some studies described the use of the microwave assisted reactions for synthesis of modified bentonites (Farias et al., 2015; Kar et al., 2016; Fabryanty et al., 2017; Mishra et al., 2017).

Therefore, the aim of the present work was to prepare organobentonites by using three surfactants (tetradecyltrimethylammonium bromide (C14), hexadecyltrimethylammonium bromide (C16) and octadecyltrimethylammonium bromide (C18)) by microwave heating. The prepared bentonites were applied to Remazol blue RN removal, while evaluating different experimental conditions. The influence of the organic chain size in the chemical structure of the adsorbents was investigated. The efficiencies of the solids were also tested for a synthetic effluent, which was composed of three different anionic dyes (Remazol blue RN, Remazol red RB and Remazol yellow GR) in the presence of inorganic salts.

2. Experimental

2.1. Materials

A sodium bentonite sample was obtained from the Bentonise Bentonita Company, Brazil. The sample was ground and sieved to a particle size below 74 μ m. The CEC was 88 cmol(+)/kg, as measured by the ammonium-exchange method (Dohrmann, 2006). The chemical composition of the bentonite after reaction with a HF/HNO₃ mixture was obtained by atomic absorption spectroscopy. Bentonite has the following constituents SiO₂ (52.98%), Al₂O₃ (18.35%), Fe₂O₃ (3.96%), Na₂O (2.56%), MgO (2.47%), TiO₂ (0.18%), K₂O (0.22%) and CaO (0.01%). Heat loss at 1223 K was 18.6%.

Three cationic surfactants C14, C16 and C18 (99% of purity) were supplied from Sigma-Aldrich and used as received.

Remazol Blue RN ($C_{22}H_{16}N_2Na_2O_{11}S_3$), remazol red GR (tetrasodium 4-[[4chloro-6-(4-sulfonatoanilino)-1,3,5-triazin-2-yl]amino]-5-oxido7-sulfo-6-[[4-(2sulfonatoethylsulfonyl) phenyl]diazenyl] naphthalene-2-sulfonate, $C_{24}H_{18}ClN_7Na_4O_{15}S_5$) and remazol yellow GR (disodium 4-(4-((2-methoxy-5-methyl-4-((2-sulfonatoethyl) sulfonyl)phenyl) diazenyl) -3-methyl-5-oxo-4,5-dihydro-1Hpyrazol-1-yl)benzenesulfonate) were supplied by DyStar Company, Brazil.

For synthetic effluent the following salts were used as received. Na₃PO₄ (Synth, 99%), Na₂CO₃ (Vetec, 99%), NaCH₃COO (Vetec, 99%), NH₄Cl (Fmaia, 99.5%), NaCl (Vetec, 99%), KNO₃ (Sigma-Aldrich, 99%) and Na₂SO₄ (Moderna, 99%).

2.2. Preparation of the organobentonites

Organobentonites were prepared as follows: 4.0 g of the clay was transferred to a Teflon vessel reactor and suspended in 100 mL of the surfactant solution. The amount of the surfactant in solution was equivalent to 100 and 200% of the CEC of the clay. The amount of each surfactant was above the critical micelle concentration, which had values of 0.0033 (C14), 0.0009 (C16) (Acisli et al., 2017) and 0.00032 (C18) (Mellott et al., 2004). The mixture was heated in a microwave reactor for 5 min at 323 K in an IS-TEC MW reactor model RMW-1, Brazil, with potency of 1100 W and 2.45 GHz. The final suspension was centrifuged, and the solid was washed until no bromide ions were detected by 0.1 mol L^{-1} silver nitrate solution. The solid was dried at 333 K for 48 h, pulverized in an agate mortar and sieved to 200 mesh. All experiments were conducted in triplicate, and the initial and final pH were monitored; however, pH variations were not observed.

2.3. Characterization

X-ray diffraction (XRD) patterns were obtained using a Shimadzu XD3A model diffractometer, with CuK α radiation ($\lambda = 0.15406$ nm), operating at 40 kV and 30 mA. The diffractions patterns were performed between 1.5-70° (2 θ) with scanning rate of 0.05 deg min⁻¹ at room temperature. Fourier transform infrared spectroscopy (FTIR) was performed in a Bomem MB-series FT spectrometer. The samples were prepared at KBr pellets with a concentration of 2%. The FTIR spectra (4000-400 cm⁻¹) were recorded with resolution of 4 cm⁻¹ and 32 accumulations. TG measurements was carried out by a TG I/1600 HT simultaneous thermal analyzer in a argon flow of 50 mL min⁻¹ with 10 mg of samples and a heating rate of 283 K min⁻¹ until 1273 K. The degree of organofunctionalization was determined based on CHN elemental analysis, which was obtained on a Perkin-Elmer PE-2400 microelemental analyzer. Zeta potential (ζ) was monitored on a Zetasizer Nano Zs -Malvern Instruments for isoelectric titration through pH titration. The pH of solutions was adjusted with 0.100 mol L⁻¹ NaOH or 0.500 mol L⁻¹ HNO₃.

2.4. Adsorption experiments

Batch experiments were employed to determine the adsorption capacities of the dyes on the modified bentonites. Adsorbent (0.05 g) was added to 20 mL of 500 mg L^{-1}

dye solution in an Erlenmeyer flask, and the samples were then shaken at 303 K for 24 h. The initial solution pH was adjusted to be in a range of 2 to 7 with 0.1 mol L^{-1} HNO₃ or NaOH. After reaching adsorption equilibrium, the mixtures were centrifuged at 5000 rpm, and the solution was collected in a syringe. The dye was quantified by a UV–VIS spectrophotometer (Shimadzu UV-2550) at 592 nm.

The amount of dye removed on the adsorbents was calculated from Equation 1:

$$q = \frac{(C_i - C_e).V}{m} \tag{1}$$

where q is the adsorption capacity (mg g^{-1}); C_i and C_e are the initial and equilibrium dye concentrations (mg L^{-1}), respectively; m (g) is the mass of adsorbent; and V (L) is the dye solution volume.

The influence of the adsorbent dosage was monitored at a fixed pH using masses of 25, 50, 75 and 100 mg, which reacted with 20 mL of 500 mg L^{-1} dye solution for 24 h. Afterwards, the same procedure was followed to separate the solid and to quantify the dye in aqueous solution.

Adsorption kinetics analysis was conducted with a 500 mg L^{-1} initial dye concentration at 303 K at the optimum pH between 10-240 min.

Equilibrium isotherms were obtained for initial dye concentrations varying from 100 to 1400 mg L^{-1} at the optimum pH, adsorbent dosage and contact time.

2.5. Kinetic and equilibrium models

Kinetic equations pseudo-first order (Lagergren and Svenska, 1898), pseudosecond order (Ho and Mckay, 1999) and Elovich model (Elovich and Larinov, 1962) were applied to study the adsorption process as described the following equations 2-4: pseudo-first order: $\ln(q_{e,exp} - q_t) = \ln q_{e,cal} - k_1 t$ (2)

pseudo-second order:
$$\frac{t}{q_t} = \frac{1}{k_2 q_{e,cal}}^2 + \frac{1}{q_{e,cal}}t$$
(3)

Elovich:
$$q_t = \frac{1}{\beta} \ln(\alpha\beta) + \frac{1}{\beta} \ln t$$
 (4)

Where q_e and q_t mg g⁻¹ are the adsorption capacities at the equilibrium and in a time t (min) respectively.

In the Eq. 4 α (mg g⁻¹ min⁻¹) and β (g mg⁻¹) are the initial adsorption rates and Elovich constant related to the extent of surface coverage and also to the activation energy involved in chemisorption, respectively.

The data of the equilibrium isotherms were fitted to Langmuir (Langmuir, 1916), Freundlich (Freundlich, 1906) and Temkin models (Temkin and Pyzhev, 1940), following the equations 5-7:

Langmuir:
$$\frac{C_e}{q_e} = \frac{1}{q_{máx}K_L} + \frac{C_e}{q_{máx}} \qquad R_L = \frac{1}{1 + K_L C_e}$$
 (5)

Freundlich: $\ln q_e = \ln K_f + \frac{1}{n_f} \ln C_e$ (6)

Temkin:
$$q_e = \frac{RT}{b_T} \ln A_T + \frac{RT}{b_T} \ln C_e$$
 (7)

Where $C_e (mg L^{-1})$ is the dye equilibrium concentration, $q_e (mg g^{-1})$ is the dye adsorbed amount on solid/liquid interface, $q_{max} (mg g^{-1})$ is the maximum removal to form a monolayer of the dye on surface, $k_L (L mg^{-1})$ is the Langmuir constant.

In the Freundlich model K_F (mg.g⁻¹) (mg.L⁻¹)^{-1/n} and n are the Freundlich constant and a factor which are related to capacity and intensity of the adsorption, respectively. For Temkin model, b_T is the constant related to adsorption energy (J mol⁻¹), At (L mg⁻¹) is the Temkin isotherm constant, R is the gas constant (8.314 J mol⁻¹ K) and T is the temperature (K).

2.6. Test with a simulated effluent solution

A synthetic dye solution consisting of three anionic dyes (Figure SM1) at 33.3 mg L^{-1} of each dye in the presence of each inorganic salt at 15 mg L^{-1} was prepared and used as the synthetic effluent (Thue et al. 2018).

The adsorption test was performed using 20 mL of the effluent solution, which reacted with 75 mg of solid and the samples were then shaken at 303 K for 3 h at pH values of 2 and 6.4. The adsorption was monitored by UV-VIS spectrometry in the range of 190 to 900 nm. The dye mixture showed maximum absorptions at 412, 540 and 609 nm associated with Remazol yellow GR, Remazol red RB and Remazol blue RN, respectively.

2.7. Stability test

The desorption experiments were conducted by suspending the dye-loaded adsorbent (50 mg of the solid previously exposed to 1000 mgL⁻¹ initial dye concentration) with 20 mL of 0.1 mol L⁻¹ NaOH or 0.1 mol L⁻¹ HCl, ethanol (95%),

separately. The mixtures were shaken at 200 rpm and 298 K for at least 12 h. After desorption, the supernatant was centrifuged and the dye in solution was quantified as described in the subsection 2.2.

3. Results and discussion

3.1. XRD

The XRD patterns of the pristine and organobentonites are depicted in Figure 1. The pure bentonite presented the characteristic peak of dioctahedral montmorillonite with reflections at 20 values of 7° (001) and 62° (060) associated with a basal spacing of 1.2 nm and interplanar distance of 0.149 nm (Duane and Robert, 1997).



Figure 1. XRD patterns for (a) pristine bentonite and organo bentonites, (b) Bent-C14-100%, (c) Bent-C16-100%, (d) Bent-C18-100%, (e) Bent-C14-200%, (f) Bent-C16-200% and (g) Bent-C18-200%.

For the organobentonites, the reflections in the *001* plan were changed for lower 20 values (Figure 1i), reflecting an increase in the basal spacing values and the occurrence of the intercalation of the surfactants in the Mt. The values were 1.83, 1.89 and 1.98 nm for Bent-C14-100%, Bent-C16-100% and Bent-C18-100%, respectively. For the samples Bent-C14-200%, Bent-C16-200% and Bent-C18-200%, the distances were 1.83, 1.90 and 2.00 nm, respectively (Figure 1ii). The increase in the basal spacing suggested the intercalation of the ammonium ions into the interlayer space of the montmorillonite. Higher values were observed for longer organic chain lengths, as reported in the literature for samples obtained by using conventional heating (Ma et al., 2011; Schampera et. al., 2016; Acisli et al., 2017). Increasing the amount of organic modifier (100 and 200% of the CEC) did not drastically alter the interlayer spacing. Although the basal spacing depends on the interlayer cations and their hydration (Lepoitevin et al., 2014), the obtained values were compared with ammonium-modified montmorillonites obtained by conventional heating (Table SM1).

The values of the basal spacings for the prepared organobentonites with 100% of the CEC were close to those obtained by Fu and Heinz (2010) using a montmorrillonite with 91 cmol(+)/kg of CEC, which was organo-modified with the same ammonium quaternary salts (C_{14} , C_{16} and C_{18}). Based on more recent studies cited by Heinz (2011), the arrangements of the organic moieties were suggested based on the difference between the d (*001*) values and the layer thickness of Mt (~0.97-1.0 nm) (Bergaya et al., 2011). For example, for the solids modified with C_{18} , the difference was 0.98 nm, which corresponds to the arrangement of the surfactant ions in a bilayer parallel to plane of the layers (Figure 2). The obtained thicknesses of the intercalated surfactants were, therefore, 0.98 nm, and this value is approximately double the thickness of the surfactant, which is known to be 0.51-0.67 nm for the hydrophilic head group from other studies, thus confirming a bilayer arrangement (Daoudi et al., 2013; Park et al., 2017; Taleb et al., 2018). Therefore, the obtained values were higher than 0.95 nm for all of the prepared organobentonites, indicating a bilayer arrangement of the alkyl ammonium cations with charged surfactant head groups disposed alternatively in the interlayer spacing of the montmorillonite. These present results are in agreement with other studies (Park et al., 2011; Daoudi et al., 2013; Sun et al., 2013; Tangaraj et al., 2017; Acisli et al., 2017).



Figure 2. Illustration of the surfactant molecules in the interlayer spacing of montmorillonite in (a) monolayer and (b) bilayer arrangements.

3.2. CHN elemental analysis

The CHN elemental analysis results for the organobentonites obtained by microwave heating are presented in Table 1. High values for the organic content were observed for solids obtained with longer organic chain lengths for both groups of modified bentonites, as indicated by the carbon percentages. These results agree with those of other works (Mizutani et al., 1995; Teppen and Aggarwal, 2007), where it was observed that the smectites are more selective for organic cations with higher chain length and with more ramified chains, where the variation in the hydration energy is a factor for the selectivity of the reaction.

Hybrid	С		Н	Ν	
	%	mmol g ⁻¹	%	%	mmol g⁻¹
Bent	0.11	0.09	1.94	-	-
Bent-C14-100%	15.64	13.03	3.57	1.08	0.77
Bent-C16-100%	17.82	14.85	3.78	1.10	0.79
Bent-C18-100%	19.15	15.96	4.12	1.15	0.82
Bent-C14-200%	18.87	15.72	4.15	1.31	0.94
Bent-C16-200%	24.74	20.62	5.20	1.50	1.07
Bent-C18-200%	28.42	23.68	5.79	1.54	1.10

Table 1. CHN elemental analysis for the organobentonites

The incorporation was better when the amount of surfactant used was 200% of the CEC. This fact can be indicative that the surfactant:bromide ion pair can be intercalated together, and therefore, ion exchange is not the only mechanism of interaction between

the montmorillonite and ammonium salts. Similar results were observed for interaction with C16 (Sun et al., 2017; Ma et al., 2011) and C18 (Sahnoun et al., 2018).

Another aspect to be considered is that basal spacings were close for both groups of solids; this indicates that the solids obtained with 200% of the CEC have higher packaging than those prepared with 100% of the CEC.

3.3. Infrared spectroscopy

Infrared spectroscopy is useful to indicate the presence of organic moieties on the solids. Bentonite was mainly composed of sodium montmorillonite and typical absorptions bands of the inorganic sheets were observed. Infrared spectrum of pristine bentonite (Figure 3) presents a band at 3630 cm⁻¹ assigned to OH structural stretching (M-OH, $M^{n+} = Al^{3+}$, Mg^{2+}) and a broad band at 3433 cm⁻¹ associated with the OH stretching of the adsorbed and interlayer water as well as SiOH (Madejová et al., 1998; Petit and Madejová, 2013). The band associated with water deformation is present at 1638 cm⁻¹. The absorption related to silicate anions is observed at 1035 cm⁻¹ due to Si-O stretching. The bands observed at 914 and 840 cm⁻¹ were assigned to Al-Al-OH and Al-Mg-OH, respectively and suggested the partial Al^{3+} substitution by Mg^{2+} in the octahedral layer (Madejová et al., 1998). Other absorptions were observed at 631 cm⁻¹ (M-O-Si; M = Al, Mg), 522 cm⁻¹ and 460 cm⁻¹ (Si-O-Al and Si–O-Mg, respectively) (Madejová et al., 1998; Petit and Madejová, 2013).



Figure 3. FTIR spectra for (a) pristine and organobentonites obtained with (i) 100% and (ii) 200% of CEC using the surfactants (b) C14, (c) C16 and (d) C18.

For the organobentonites, a displacement in the wavenumbers of the OH absorptions compared with the spectrum of the pristine solid was not observed. This indicated that interaction is preponderantly in the interlayer region (Xi et al., 2005). New, sharp and strong bands were observed for all hybrids at 2926 and 2848 cm⁻¹ which were assigned to the asymmetric and symmetric C-H stretchings of methylene groups (Ouellet-Plamondon et al., 2014; Acisli et al., 2017; Martinez-Costa et al., 2017; Santos et al., 2018) and suggest the presence of the ammonium moieties on the bentonite (Acisli et al., 2017).

For all of the hybrids, both C-H stretching bands were sharp and intense without alterations in the wavenumbers. Other new bands were observed at 1475 cm⁻¹ and 725 cm⁻¹ and were assigned to the scissoring and rocking bending of CH_2 , respectively (Ouellet-Plamondon et al., 2014; Martinez-Costa et al., 2017; Lazorenko et al., 2018; Santos et al., 2018). The CH₂ scissoring band normally overlaps with the CH₃ antisymmetrical bending band, which occurs in the same region (Lin-Vien et al. 1995).

3.4. Thermogravimetry (TG)

Thermogravimetry analysis was useful to quantify the percentages of organic matter in the solids. The TG results are presented in Figure 4 and Table SM2.

The first mass loss (2.3%) for pristine bentonite between 298-383 K was attributed to the desorption of external and interlayer water (Park et al., 2011; Ma et al., 2015; Santos et al., 2018) and the second loss of 5% between 560 – 1126 K was associated with the dehydroxylation of bentonite. The percentages of total mass losses between 298-1200 K were higher for organobentonites than pristine bentonite. The first mass loss associated with the desorption of interlayer water was lower for the

organobentonites (<1%) than the pristine sample, which was attributed to the hydrophilic character of the samples after the organofunctionalization (Moslemizadeh et al., 2016; Santos et al., 2018). The thermal elimination of alkyl surfactants proceeded at temperatures above 473 K. The second, third, fourth and fifth events in the TG curves of the hybrids were attributed to loss of the surfactant moieties and are in agreement with the literature (Park et al., 2011; Zhu et al., 2012; Taleb et al., 2018).

Ma et al (2015) observed a single mass loss with a maximum at 579 K in the DTG curve and attributed the mass loss to the desorption of the surfactant. In fact, in the present case, an intense loss mass at 579 K was observed for all hybrids, but multiple mass losses were observed. This fact was associated with the different interactions between the surfactants and bentonite as the concentration of organic moieties increased (Teppen and Aggarwal, 2007; Park et al., 2011; Taleb et al., 2018). The last mass loss for all hybrids (~2.0%, above 799 K) was attributed to the dehydroxylation of the solid (Ma et al., 2015; Santos et al., 2018).



Figure 4. (i) TG and (ii) DTG curves for (a) pristine and organobentonites (b) Bent-C14-100%, (c), Bent-C16-100%, (d) Bent-C18-100%, (e) Bent-C14-200%, (f) Bent-C16-200% and (g) Bent-C18-200%.

3.5. Zeta potential

The zeta potential measurement is an important method to define the electrokinetic properties of a surface and determine the point of zero charge (pH_{PZC}) since the anionic dye adsorption is favorable in pH values lower than the point of zero charge (Yagub et al., 2014). At the point of zero charge, the quantity of positive and negative charges on surface is equal and the net charge is zero (McCafferty, 2010).

Point of zero charge (PZC) is the pH where the surface charge of a system double-layer reversible is zero, being determined by a particular ion activity, (Parks and Bruyn, 1962). The pH_{IEP} (IEP for isoelectric point) and pH_{PZC} correspond to the pH of the aqueous solution, in which amphoteric substances have a neutral charge, however, the pH_{IEP} determines the charge of the external surface of the adsorbent particles in solution, and the pH_{PZC} results in the total value, that is, the charge of both the internal and external surfaces (Menéndez et al., 1995). The point of zero charge is the pH value where negative and positive surface concentrations are equal, i.e., surface charge charge is 0. The surface charge is negative at pH > pHpzc and positive at pH < pHpzc. The isoelectric point is the pH at which zeta potential of a molecule or surface is equal to zero, ξ = 0. In order to get a sharp isoelectric point, the surface must be amphoteric, meaning it must have both acidic and basic functional groups. In the simplest case, if the specific sorption of ions and dissociation of counter ions are neglected, then pHpzc coincides with pHiep (Gulicovski et. al., 2008).

Therefore, at pH values below the point of zero charge (pH_{PZC}), the edges sites of the clay mineral surface are positively charged and the electrostatic interaction is enhanced. At pH values higher than the PZC, the edges the clay mineral surface was deprotonated, the negatively charged dye was barely able to reach the surface adsorption sites.

Bentonite has a net negative charges in aqueous medium independent of the pH (Swartzen-allen and Matijevic, 1974). The organofunctionalization of the bentonite can modify the surface charge and change it to partially or entirely positive. In this present study, the hybrids prepared with 100% of the CEC presented pH_{PZC} at 4.1 ± 0.1; 10.5 and 10.5 ± 0.1 for Bent-C14-100%, Bent-C16-100% and Bent-C18-100%, respectively. This result indicated that at pH lower than the point of zero charge, the hybrids are positively charged (Figure 5), favoring in these values the adsorption of the anionic dye. For the hybrids prepared with 200% of the CEC, a positive surface charge was observed for all pH values, without a point of zero charge. Therefore, Bent-C14-200%, Bent-C16-200% and Bent-C18-200% presented higher dye removal in all pH. The same behavior was observed for other studies in the literature (Ma et al., 2011; Tunç et al., 2012).



Figure 5. Zeta potential (ζ) measurements for (a) Bent-C14-100%, (b) Bent-C16-100%,
(c) Bent-C18-100%, (d) Bent-C14-200%, (e) Bent-C16-200% and (f) Bent-C18-200%.

3.6. Microscopy

3.6.1 Scanning electronic microscopy (SEM)

SEM measurement was used to investigate alterations in the morphology of the solids after reactions with surfactants. SEM images (Figure SM2) showed typical flaky morphology for the pristine bentonite. After organofunctionalization, the solids presented a more irregular surface and dispersed flaky. The same behavior was observed for organobentonites (Santos et al., 2018; Sun et al., 2017).

3.6.2 Transmission electronic microscopy (TEM)

TEM micrographs (Figure 6) show that organobentonites maintained the layered structure of the sodium bentonite without the formation of exfoliated phases. The measured interlayer spacings were 1.27, 1.82, 1.92, 2.05, 1.82, 1.94 and 2.17 nm for Bent, Bent-C14-100%, Bent-C16-100%, Bent-C18-100%, Bent-C14-200%, Bent-C16-200% and Bent-C18-200%, respectively and were in concordance with the values of the basal spacing obtained in the XRD analysis.



Figure 6. TEM for (a) pristine and organobentonites (b) Bent-C14-100%, (c), Bent-C16-100%, (d) Bent-C18-100%, (e) Bent-C14-200%, (f) Bent-C16-200% and (g) Bent-C18-200%.

3.7. Adsorption

3.7.1. Influence of the pH

The influence of the pH on adsorption is well-documented due to the charge on the surface and the speciation of the pollutant species in solution (Yagub et al., 2014; Ngulube et al., 2017; Kausar et al., 2018). In acidic conditions, some surfaces are positively charged, and the removal of anionic dye is improved (Ngulube et al., 2017).

The effect of the pH on RN blue dye removal is shown in Figure 7 for the test at 303 K with 50 mg and a 500 mg·L⁻¹ dye concentration.

The dye removal by the three hybrids obtained with 100% of the CEC was pH dependent, and the highest value of retained dye was at a pH of 2.0. This behavior agrees with the positive charge (20 - 36 mV) (pH < 4.1) based on zeta potential measurements. The Bent-C14-100% sample showed a dye removal of 160 mg g⁻¹ at a pH of 2.0, while the value was 173 mg·g⁻¹ for both Bent-C16-100% and Bent-C18-100% under the same conditions. Early studies reported that lower pH values contribute to higher adsorption (Ozcan et al., 2007; Ma et al., 2011).

For the hybrids obtained using 200% of the CEC (Figura 7ii), the behavior was different. Indeed, the performances of the solids were very close, and these results are in opposition to the zeta potential measurements since the hybrids synthesized with 200% of the CEC showed a positive surface throughout the entire pH range. A similar result was observed by Yan et al., (2015). It is possible that the electrostatic force is the principal factor in relation to the affinity between the dye and the hybrids, but the amount of the organocations in the hybrid influenced the adsorption (Zhang et al., 2016). Therefore, the maximum removal was close to 206 mg·g⁻¹ for Bent-C14-200% at pH values in the range of 3.0 to 7.0, and the minimum value was 188.3 mg·g⁻¹ at a pH of 2.0. The Bent-C16-200% and Bent-C18-200% hybrids presented almost the same behavior and maximum adsorption values of 169 mg·g⁻¹. For Bent-C14-200%, the maximum adsorption was higher than 99% at all pH values, while the maximum values were 82% and minimum 62% for Bent-C16-200% and maximum 78% and minimum 66% for Bent-C18-200%.

Considering that many residual waters have pH values in the range of 6.0 to 9.0 (Yan et al., 2015), the hybrids prepared with 200% of the CEC have the advantage with regards to practical applications because pH adjustment of the solution is not necessary.



Figure 7. Effect of the pH on adsorption of remazol blue RN (inserted figure) by (a) pristine and organobentonites obtained with (i) 100% and (ii) 200% of CEC using the

surfactants (b) C14, (c) C16 and (d) C18 at 303 K. Conditions: 50 mg of solid, 20 mL of the 500 mg L^{-1} dye solution, contact time of 24 h, pH 2.0 to 7.0 at 303 K.

3.7.2. Dosage of the adsorbent

The influence of the adsorbent dosage (Figure SM3) was monitored by contacting different amounts of the solid (25-100 mg) with 20 mL of a 500 mg \cdot L⁻¹ dye concentration at 303 K and a pH 2.0 for the maximum removal for 24 h.

The adsorption efficiency of solids initially increases with the adsorbent dosage, and it finally reaches a plateau. The optimum dosage for dye removal was found to be 75 mg of solid for all hybrids synthesized by using 100% and 200% of the CEC, except for Bent-C16-200%, for which the better dosage was 100 mg. Generally, the increase in adsorption efficiency with increasing the amount of the adsorbent in the system is associated with the availability of more adsorption sites resulting from the increase in adsorbent particles (Ngulube et al., 2017). The optimum adsorbent dose is mainly related to availability of active site but also is directly linked with the concentration of the adsorbate (Iftekhar et al., 2018). The effect of the adsorbent dosage gives an idea of the capacity of an adsorbent in the adsorption of a determined dye, so as to use the smaller amount of adsorbent needed for determined experimental condition and concentration of adsorbent needed for determined experimental condition and concentration of adsorbent needed is quick because of the readily available active sites and when the adsorbent dosage is high, the dye ions cannot readily access the adsorption sites easily until the attainment of equilibrium (Sarma et al., 2016).

3.7.3. Effect of time on the adsorption

The adsorption kinetics were evaluated by contacting 75 or 100 mg of the solid with 20 mL of the 500 mg·L⁻¹ dye solutions at the optimum pH (Figure 8). Corroborating with the literature, the adsorption was fast during the first few minutes of the interaction, and after 20 min, all of the solids obtained with 100% of the CEC adsorbed 100 mg·g⁻¹, reaching a dye removal higher than >95%. The behavior was similar and the equilibrium was reached after 60 min with an adsorption capacity of close to 125 mg·g⁻¹ for all hybrids.

For the hybrids obtained with 200% of the CEC, the equilibrium time was achieved after 60 min with maximum removals close 140 mg \cdot g⁻¹, except for Bent-C16-200%, for which the equilibrium time was 240 min.

In order to interpret the adsorption kinetics, the pseudo-first-order, pseudosecond-order and Elovich models were applied (Figure SM4). The kinetic parameters of the fittings for the cited models are presented in Table 2. The higher R^2 regression and lower difference between the $q_{e,exp}$ and $q_{e,theor}$ show that the experimental data were well-fitted to the pseudo-second-order model. The fit of this model indicates that adsorption rate is controlled by chemisorption process, which involves valence forces from the sharing of electrons between dye molecules and material surface (Foo et al., 2012; Silva 2016). Other studies related to adsorption of the same dye on different adsorbents demonstrated the same behavior (Pereira et al., 2017; Silva et al., 2012; Malakootian et al., 2015).



Figure 8. Time isotherms for the adsorption of remazol blue RN by (a) pristine and organobentonites obtained with (i) 100% and (ii) 200% of the CEC using the surfactants (b) C14, (c) C16 and (d) C18. Conditions: 75 or 100 mg of solid, 20 mL of the 500 mg L^{-1} dye solution, contact times of 60-240 min, pH 2.0 or 6.5 at 303 K.

Pseudo-first order

Table 2. Kinetic parameters obtained by fitting the adsorption data of Remazol blue RN

 on organobentonites to pseudo-first order, pseudo-second order and Elovich models.

Hybrid	q _{e,exp}	q _e ,theor	k ₁			
	$(mg g^{-1})$	$(mg g^{-1})$	10^{-2} (min ⁻¹)	\mathbf{R}^2		
Bent-C14-100%	123.0	12.6	9.82	0.9619		
Bent-C16-100%	125.4	3.3	5.22	0.5684		
Bent-C18-100%	125.4	28.2	9.69	0.9461		
Bent-C14-200%	142.8	49.9	3.00	0.7148		
Bent-C16-200%	75.1	29.2	0.20	0.3908		
Bent-C18-200%	140.7	16.6	4.40	0.7092		
Pseudo-second order						
Hybrid	q _e ,exp	q _e ,theor	k ₂			
	$(mg g^{-1})$	$(mg g^{-1})$	$10^{-2} (g mg^{-1} min^{-1})$	R^2		
Bent-C14-100%	123.0	124.2	1.68	0.9999		
Bent-C16-100%	125.4	125.9	2.87	0.9999		
Bent-C18-100%	125.4	126.8	0.91	0.9998		
Bent-C14-200%	142.8	166.7	0.07	0.9998		
Bent-C16-200%	75.1	71.4	0.07	0.9345		
Bent-C18-200%	140.7	142.8	0.54	0.9992		
Elovich						
Hybrid	$q_{e,exp}$	α	β	R^2		
	(mg g)	(mg g^{-1})	$(g mg^{-1})$			
Bent-C14-100%	123.0	4.76 10 ¹⁹	0.3917	0.9785		
Bent-C16-100%	125.4	4.11 10 ²⁷	0.5331	0.7447		
Bent-C18-100%	125.4	3.11 10 ⁷	0.1555	0.5649		
Bent-C14-200%	142.8	532.03	0.0594	0.8248		
Bent-C16-200%	75.1	146.46	0.1331	0.5999		
Bent-C18-200%	140.7	1.31 10 ¹²	0.2184	0.8129		

3.7.4. Effect of the initial dye concentration

Equilibrium isotherms for the adsorption of Remazol Blue RN on organobentonites are shown in Figure 8. Studies reported that dye removal increased with the increase of the initial dye concentration until saturation is reached (Makhoukhi et al., 2015; Ngulube et al., 2017).

The initial dye concentration of Remazol blue RN modified the quantity of adsorbed dye on the different solids until equilibrium (Figure 9), where saturation occurred at the initial concentration of 1000 mg L^{-1} for solids obtained with 100% of the CEC and 1200 mg L^{-1} for those prepared with 200% of the CEC. The maximum retentions were 195, 199 and 209 mg g⁻¹ for Bent-C14-100%, Bent-C16-100% and Bent-C18-100% as well as 211, 163 and 287 mg g⁻¹ for Bent-C14-200%, Bent-C16-200% and Bent-C18-200%, respectively.

The equilibrium data were adjusted to Langmuir, Freundlich and Temkin models (Figure SM5), and the resulting parameters are summarized in Table 3.

Taking into account the coefficients of correlation (R^2) , the better adjustments were observed for the Langmuir model for all of the investigated systems, indicating that chemisorption can be the preponderant mechanism involved in the Remazol blue RN dye removal.



Figure 9. Equilibrium isotherms for the adsorption of remazol blue RN by (a) pristine and organobentonites obtained with (i) 100% and (ii) 200% of the CEC using the surfactants (b) C14, (c) C16 and (d) C18. Conditions: 75 or 100 mg of solid, 20 mL of the 100-1400 mg L^{-1} dye solution, 60 or 240 min, pH 2.0 or 6.5 at 303 K.

Langmuir									
Hybrid	q_{exp}	q _e ,theor	R^2						
	$(mg g^{-1})$	$(mg g^{-1})$							
Bent-C14-100%	192.9	200.0	0.9992						
Bent-C16-100%	194.5	200.0	0.9984						
Bent-C18-100%	205.4	250.0	0.9996						
Bent-C14-200%	211.9	250.0	0.9961						
Bent-C16-200%	163.7	169.4	0.9985						
Bent-C18-200%	287.4	361.1	0.9933						
Freundlich									
Hybrid	n _F	K _F	R^2						
		$(\text{mg g}^{-1})(\text{mg L}^{-1})^{-1/n}$							
Bent-C14-100%	5.9524	70.74	0.6445						
Bent-C16-100%	6.3694	72.82	0.4085						
Bent-C18-100%	6.2112	82.35	0.5763						
Bent-C14-200%	6.6445	84.02	0.1652						
Bent-C16-200%	1.3928	3.06	0.1420						
Bent-C18-200%	1.1429	2.73	-0.0060						
Temkin									
Hybrid	b _T	A _T	R^2						
Bent-C14-100%	126.00	43.07	0.8182						
Bent-C16-100%	127.64	48.77	0.6310						
Bent-C18-100%	123.24	82.99	0.7474						
Bent-C14-200%	126.26	107.89	0.3130						
Bent-C16-200%	40.38	0.05	0.3974						
Bent-C18-200%	17.13	0.03	0.1165						

Table 3. Parameters obtained by fitting the adsorption data of Remazol blue RN on

 organobentonites to Langmuir, Freundlich and Temkin models.

The same behavior was observed in other studies for bentonites modified by surfactants applied in dye adsorption, which were adjusted to the pseudo second order model kinetic (Anirudhan et al., 2015; Yan et al., 2015; Sahnoun et al., 2018) and to the Langmuir model equilibrium (Ren et al., 2018; Sahnoun et al., 2018;).

Table SM3 shows the adsorption capacity for remazol blue RN dye by different adsorbents including alginate gel beads (Ergene et al., 2009), modified montmorillonites (Silva et al., 2012; Pereira et al., 2017), activated carbon (Isah et al., 2015; Silva et al. 2016); alumina-coated multi-walled carbon nanotubes (Malakootian et al., 2015) and porous nano-crystalline CoFe₂O₄ composite (Khan et al., 2015).

The XRD patterns and infrared spectra of the hybrids after the adsorption process of the Remazol blue RN did not have significant variations attributed to the overlapping of the absorptions since the bands of the dye occur in the same region of the organobentonite. Solid state UV-VIS spectra showed a new band at approximately 592 nm attributed to Remazol blue RN dye (Figure SM6).

Based on zeta potential measurements and the pH isotherms and kinetic and equilibrium models, the preponderant mechanism of adsorption could be electrostatic interaction between the anionic dye and the alkyl ammonium moieties disposed in the interlayer spacing and external surface of the montmorillonite and also the chemisorption process (Figure SM7) (Zhang et al., 2016). However, the present set of techniques are not conclusive about the chemical forces between the dye and the surface. Another possible mechanism is by anion exchange between the bromides intercalated as counter ions in the ionic pair with the cations. Some works have proposed the interaction by anion exchange involving the dye and Br⁻ previously intercalated as the counter ion of the ammonium salt (Ma et al., 2011; Sun et al., 2017; Sahnoun et al., 2018).

In order to check this hypothesis, qualitative tests were carried out with silver nitrate solution in the supernatant obtained after the adsorption process. The formation of a white precipitate was associated with the presence of bromide ions in the supernatant, indicating that anion exchange between the Br⁻ in the montmorillonite and the anionic dye in solution contributed to the removal. As a control, the procedure was repeated by suspending the organobentonites in distilled water, and no precipitate was formed, indicating the absence of bromides on the solid due insufficient washing. Therefore, the presence of bromide ions in the supernatant after dye adsorption was associated with the anion exchange mechanism.

3.7.5 Desorption test

This study was performed to evaluate the regeneration capacity of the adsorbent for reuse. The desorption test defines whether the potential of the solid for reuse or stability of the dye/solid complex for other purposes, such as pigment. The reusability is one of the important characteristics of any kind of adsorbents and is a critical point on the economy of the process (Salehi et al., 2016). However, in recent years great importance was given to the preparation of new stable pigments based on clay minerals (Fournier et al., 2016; Zhang et al., 2017; Guillermin et al., 2018; Trigueiro et al., 2018). The study of new materials includes the search for new pigments by immobilization of the dyes in resistant substrates such clays minerals to get new color fonts and materials with high stability (Fournier et al., 2016; Trigueiro et al., 2016).

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Some studies show that acid and basic solutions are favorable to desorption (Sathishkumar et al., 2012; Anirudhan et al., 2015; Salehi et al., 2016). In the present work, the dye/bentonite hybrid presented high stability and did not show any desorption in any of the acid and basic solutions.

3.8. Adsorption with simulated dye effluent

UV-VIS spectra (Figure SM8) of the synthetic effluent before and after adsorption were recorded in the range of 190-900 nm. The tests were conducted at natural pH of the dye solution 6.4, and a controlled pH of 2.0 for 3 h at room temperature. The absorbance of the mixture was monitored and used to calculate the dye concentration in solutions before and after the interaction with the organobentonites.

For all of the hybrids, the absorbance in the monitored wavenumbers ($\lambda_{max} = 412$ nm, Remazol yellow GR dye; $\lambda_{max} = 540$ nm, Remazol red RB dye; and $\lambda_{max} = 609$ nm, Remazol blue RN dye) decreased significantly after adsorption (Figure 10 and Figure SM9). For Bent-C14-100% and Bent-C14-200%, the dye removals were very close to 100% at pH 6.4. The Bent-C16-100% and Bent-C18-100% showed higher decay in the absorptions than Bent-C16-200% and Bent-C18-200%. Therefore, for the dye mixtures, the high amounts of the organic moieties were not a preponderant factor due the competition between the anionic dyes and the higher packing density of the solids with long organic moieties.



Figure 10. Results for the removal of the dyes at pH value of 6.4 in a synthetic effluent. The macroscopic picture is the solution after centrifugation using Bent-C14-200%, Bent-C16-200%, Bent-C18-200%.

The best results were for Bent-C14-100% and Bent-C14-200%, with removals close 98% and 99%, respectively. Bent-C16-200% and Bent-C18-200% hybrids presented less efficiency to remove all of the dye in the mixture; however both hybrids were selective for Remazol yellow GR, which has a maximum absorbance at 412 nm. The insert image in Figure 9 shows the macroscopic picture of the systems with the synthetic effluent, and while the solution is clear after removal using Bent-C14-200%, there is color in the suspension after the interaction with Bent-C16-200% and Bent-C18-200%. However, both solids were yellow after adsorption in the synthetic effluent.

It was observed that Bent-C14 showed the best performance for Remazol blue RN and for the dyes in the synthetic effluent. The behavior of Bent-C14 was associated with the lower density packing of the organic moieties compared with Bent-C18, as suggested in the literature (Heinz, 2011).

4. CONCLUSIONS

The use of microwave heating for the preparation of organobentonites at 323 K is a rapid and efficient methodology (5 min). The prepared solids showed the intercalation of surfactant molecules into the interlayer spacing with basal spacings varying in the range of 1.83 to 2.00 nm.

The organosolids have good performance for dye removal from aqueous solution, although increasing the amount and the length of organic alkyl chain for hybrids prepared with 100% and 200% of the CEC did not result in a higher efficiency of the solids. The change in the surface charge of organobentonites contributed to high Remazol blue RN removal as well as good removal of the dyes in the synthetic effluent. The best performance in both tests was for Bent-C14-100% and Bent-C14-200%. Furthermore, the Bent-C14-200% was efficient in acidic conditions and at the pH of the dye solution. The Bent-C16-200% and Bent-C18-200% hybrids were selective for Remazol yellow GR.

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Supplementary material

Figure SM1. Molecular structure of the dyes a) Remazol yellow GR, b) Remazol red RB and c) Remazol blue RN.

Figure SM2. SEM for (a) pristine and organobentonites (b) Bent-C14-100%, (c), Bent-C16-100%, (d) Bent-C18-100%, (e) Bent-C14-200%, (f) Bent-C16-200% and (g) Bent-C18-200%.

Figure SM3. Effect of the dosage of the adsorbent on the adsorption of Remazol blue RN by (a) pristine and organobentonites obtained with (i) 100% and (ii) 200% of the CEC using the surfactants (b) C14, (c) C16 and (d) C18 at 303 K. Experimental conditions: 25 to 100 mg of solid, 20 mL of the 500 mg g-1 dye solution, contact time of 24 h, pH 2.0 at 303 K.

Figure SM4. Fitting of the adsorption data of Remazol blue RN on organobentonites (a) Bent-C14, (b) Bent-C16 and (c) Bent-C18 prepared with (i) 100 and (ii) 200% of the CEC for pseudo-first order; (iii) 100 and (iv) 200% of the CEC for pseudo-second order and (v) 100 and (vi) 200% of the CEC for Elovich models.

Figure SM5. Fitting of the adsorption data of Remazol blue RN on organobentonites (a) Bent-C14, (b) Bent-C16 and (c) Bent-C18 prepared with (i) 100 and (ii) 200% of the CEC for Langmuir; (iii) 100 and (iv) 200% of the CEC for Freundlich and (v) 100 and (vi) 200% of the CEC for Temkin models.

Figure SM6. i) XRD patterns; ii) UV-VIS solid stated spectra for Bent-C14-200% a) before and b) after dye removal and iii) infrared spectra for a) Bent-C14-200%, b) Remazol blue dye and c) Bent-C14-200% after dye removal.

Figure SM7. Proposed mechanism for the interaction between the dye and the organobentonites on the external and interlayer regions.

Figure SM8. UV-VIS absorption spectra of the synthetic dye effluent before and after adsorption for pure bentonite and hybrids prepared with (i) 100% and (ii) 200% of the CEC in a pH of 2.0 and (iii) 100% and (iv) 200% of the CEC in a pH of 6.4 for (a) initial dye effluent solution and the solutions after adsorption using solids modified with (b) Bent, (c) C14, (d) C16 and (e) C18. Experimental conditions: 3 h, 298 K, 75 mg of solid and 20 mL of effluent solution.

Figure SM9. Results for the removal of the dyes at pH value of 2.0 in a synthetic effluent by monitoring using UV-Vis spectrometry at the maximum absorptions of 412, 540 and 609 nm for organobentonite.

Table SM1. Values of the basal spacing for organobentonites prepared with the surfactants C14, C16 and C18 for solids obtained by conventional heating accordingly to the literature.

Table SM2. Data of mass losses and temperature (T) obtained from thermogravimetry (TG/DTG) for the pristine and organo bentonites.

Table SM3. Adsorption capacity and time for remazol blue RN dye by using different adsorbents. R^2 is the correlation coefficient obtained by fitting experimental data to pseudo-second order model.