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Ceramic bricks containing Ni ions from contaminated biomass used as an adsorbent

Daiana Simón¹, Nancy Quaranta², Sebastián Gass¹, Raúl Procaccini¹ and Adrián Cristóbal^{1*}

Abstract

This article shows how pine sawdust residues can be used to adsorb nickel ions from synthetic solutions and then to produce porous bricks for civil construction using a mixture of natural clay and biomass containing the adsorbed metals. The adsorption tests were performed by mixing NiCl₂ solutions with pine sawdust during a fixed stirring period of 24 h. The set was filtered and the filtrate was analysed. Highest efficiency adsorbate/adsorbent ratio was 50 mL of 1 M NiCl₂ solution and 20 g L⁻¹ of pine sawdust. This was the contaminated biomass sample used in the manufacture of the bricks. This paper analyses the properties of the bricks achieved and compares them with bricks without added biomass, porous bricks containing zinc and commercial bricks. The obtained values of bulk density, apparent specific weight, apparent porosity, water absorption, apparent volume, weight loss on ignition, compressive strength, flexural modulus of rupture and efficiency retention of metal in the brick, demonstrate that the ceramic pieces obtained are optimal for construction.

Keywords: Toxic metals, Wood waste, Adsorption, Porous bricks, Cooked ceramic immobilization, Sustainability

Introduction

Heavy metals are the main inorganic micro pollutants, and their discharge into aquatic bodies affects the ecosystem and health due to their toxicity and possible carcinogenic effect. The lack of biodegradability of these pollutants results in their bioaccumulation and persistence in the environment, and their danger is given both by the concentrations as well as their chemical behaviour. The anthropogenic sources of heavy metals are very varied, with the metal finishing industry being the most contributor due to the large number of companies that integrate and its geographical dispersion [1–3].

Nickel is a dangerous heavy metal. The World Health Organization states that 0.07 mg L⁻¹ is the allowed limit of this contaminant in water for consumption [4]. This contaminant can cause adverse effects on the blood,

kidneys, bronchi, lung and stomach. The main sources of nickel are the stainless steel and alloy, electroplating, catalysts and Ni/Cd industries. The reduction of nickel concentration in water and wastewater is of great interest due to the current increase in demand of such emitting industries [5, 6].

The most commonly used conventional methods for removing Ni (II) and other toxic metals from wastewater are chemical precipitation, solvent extraction, ion exchange, flotation, adsorption with activated carbon, among others [6, 7]. However, to date, there is no low-cost technology that allows to reach very low levels of these pollutants. In this situation, the removal of heavy metal ions using low-cost adsorbents has recently gained relevance [8–10]. Research has shown that materials of biological origin, such as biomass of algae, fungi and bacteria, as well as residues and by-products from the agro-industrial sector, can retain heavy metal ions [3, 11]. Bioadsorption of pollutants from cellulosic biomass residues has attracted attention because it not only allows the

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removal of heavy metals in industrial effluents, but also treats agricultural waste that is produced today in huge amounts and does not have a specific use [12]. In this sense, they have been studied as adsorbents including rice husk [13, 14], sawdust [15, 16], fruit peels [17, 18] and others [19, 20].

Heavy metals are attracted to the adsorbent through a complex process involving mechanisms such as surface adsorption (by Van Der Waals forces, dipole interactions, or hydrogen bonding), interstitial adsorption (through the pores of the material), ion exchange adsorption (between metal ions and adsorbent ions, facilitated by carboxyl groups and hydroxyl), by electrostatic forces (related to the pH of the solution and the zero charge point of the adsorbent), complexation (through carboxylate groups, amides, phosphate, thiols and hydroxide), precipitation, among others [10, 21]. The availability of the active biomass sites is influenced by the type of biomass, metallic solution, pH, temperature, contact time and stirring speed of the mixture [10, 22].

Currently, there are researches on the production of porous ceramic bricks from the incorporation of agroindustrial biomass residues. These residues act as pore-forming agents because at the sintering temperatures their combustion produces gases and ashes. For this purpose, residue of olive [23], sunflower and wheat [24], fruit pits [25], vine [26] and rice [27] have been investigated.

The aim of this article is to adsorb Ni (II) ions on wood industry residues (pine sawdust) and then use these new contaminated residues as an aggregate in clay to generate optimal porous ceramic matrices for civil construction, which will contain retained metals. The metals will be immobilized in the ceramic pieces, stabilizing them and reducing their release, solubility and toxicity. In this way, the negative effects on the environment are minimized [28, 29].

Materials and methods

Adsorbate

The metal solutions used in this study (0.125 to 1 M) were prepared by dissolving appropriately measured amounts of $\text{NiCl}_2 \times 6 \text{H}_2\text{O}$ salt (analytical grade, Cicarelli) in distilled water.

Bioadsorbent

The adsorbent used was the residue obtained from the processing of *Pinus elliottii* wood in a sawmill from the north east area of Argentina. This residue was sieved to obtain particles smaller than 1 mm. It was then boiled with distilled water for 90 min, filtered and washed repeatedly. Finally, the material was dried in an oven at 50 °C.

Batch adsorption experiments

A qualitative study of the adsorption at different times of Ni (II) ions of 50 mL of 0.25 M NiCl_2 solutions in contact with 10 g L^{-1} of pine sawdust was carried out. Metallic solutions prepared at pH 5–6 (PH/MV/°C HANNA HI 8424), were put in contact with pine sawdust by constant stirring at room temperature of 200 rpm (SK-0330-Pro shaker). Then, each mixture was filtered and the solid residues were dried at 50 °C. The pH was chosen according to what is reported in the literature [10, 30], at higher pH values Ni (II) forms a complex with hydroxide ions, precipitating and decreasing adsorption, and at a pH less than this value, H^+ can compete with ions of heavy metals by the active sites of the adsorbent and also, functional groups of the adsorbent that intervene in the adsorption process can be protonated (general positive charge on the adsorbent), generating an electrostatic repulsion on the positively charged metal ions leading to adsorption. This test was performed with the aim of estimating an optimal time for contact between heavy metal ions and pine sawdust biomass by means of X-ray fluorescence (XRF) measurements on the solids. The tests were done in triplicate.

Subsequently, adsorption tests were performed at different concentrations of NiCl_2 from 0.125 to 1 M and varying from 10 to 40 g L^{-1} the biomass residue content to determine the most efficient adsorbate/adsorbent ratio used in the manufacture of ceramic pieces. For this, the amount of residual metal that was not adsorbed on the biomass and remained in the solution after the filtration process was quantified by UV-Vis Spectrophotometry (UV-Vis) at $\lambda = 722 \text{ nm}$ (this λ was chosen to reduce the signal obtained by the organic matter, in addition to making blank samples). Adsorption tests were performed in triplicate for each system.

The adsorption efficiency (R) (%) of Ni (II) was calculated following Eq. (1) and the adsorption capacity (q_e) (mg g^{-1}) was defined by Eq. (2):

$$R (\%) = \frac{(C_i - C) \times 100}{C_i} \quad (1)$$

$$q_e \left(\frac{\text{mg}}{\text{g}} \right) = \frac{(C_i - C) \times V}{M} \quad (2)$$

where C_i and C (mg L^{-1}) are the initial and residual concentration of Ni (II); V (L) is the volume of solution and M (g) is the mass of the adsorbent. The amount C was determined by UV-Vis.

Bricks production and characterization

The bricks used to carry out the mechanical tests were made by mixing 100 g of clay from a natural quarry in eastern Buenos Aires and the biomass containing the adsorbed Ni (II) in a proportion of 20 vol% of clay. The

particle size of the contaminated pine sawdust was less than 1 mm and the particle size of the clay was less than that of the biomass.

The adsorption sample used for the manufacture of bricks was the one representing the highest efficiency adsorbate/adsorbent ratio determined by the adsorption experiments.

For formation, a uniaxial pressure of 25 MPa was used, with the addition of 8 mL of water, in molds of $70 \times 40 \times 18$ mm. After a drying period of 24 h in the environment, the green bricks were calcined at 950°C for 3 h. The heating rate was 1°C min^{-1} and the cooling rate was 5°C min^{-1} .

The sintered ceramic pieces were physically characterized and their mechanical properties were evaluated in triplicate.

Weight loss on ignition (LOI) was determined. Also, apparent porosity (apP), together with bulk density (apD), apparent specific weight (apsW), water absorption (H_2OAbs) and apparent volume (apV), were determined following the Archimedes principle and based on the standard [31].

Flexural modulus of rupture (MOR) and compressive strength (σ_{str}) were evaluated in test specimens in accordance with the provisions of the IRAM 12587 standard and the IRAM 12586:2004 standard, respectively. The MOR was determined on rectified samples of $70 \times 40 \times 18$ mm, submitted to a three-point bending test. The σ_{str} was determined on rectified samples of $35 \times 40 \times 18$ mm.

Leaching tests on 10 g of the bricks were carried out following the procedure described in EPA 1311 Toxicity Characteristic Leaching Procedure (TCLP) and in triplicate. The brick fragments were contacted with 20 times the volume of extraction solvent (leaching solution of $\text{pH } 4.93 \pm 0.05$, prepared from 5.7 mL of 17.4 M glacial acetic acid in 500 mL of distilled water, adding 64.3 mL of 1 N sodium hydroxide and diluting up to 1 L). The mixture was stirred at 50 rpm for 18 h. The system was filtered and the concentration of Ni (II) was determined on the filtered liquid by Atomic Absorption Spectrophotometry (AAS).

Equipment

Thermogravimetric-differential thermal analysis (TGA-DTA): Shimadzu TGA-50 and Shimadzu DTA-50 equipment, with TA-50 WSI analyser. Thermal characterization of pine sawdust residues. Conditions: air flow, heating up to 1000°C at a rate of $10^\circ\text{C min}^{-1}$ and approximately 20 mg of residue.

Scanning electron microscopy (SEM) - energy dispersive analysis of X-ray (EDS): SEM Philips 515 equipment, with energy dispersive analyser (EDAX-Phoenix). Microstructural characterization of pine sawdust residues.

Determination of Ni (II) in the biomass after the adsorption process and in the manufactured bricks.

Fourier-transform infrared spectroscopy (FTIR) in attenuated total reflectance - ATR mode: Nicolet 6700, Thermo Electron equipment. Characterization of the functional groups present in the biomass of pine sawdust before and after the Ni (II) adsorption process.

X-ray diffraction (XRD): PANalytical X'Pert PRO equipment, with $\text{CuK}\alpha$ radiation ($\lambda = 1.5406$ nm). Characterization of pine sawdust before and after Ni (II) adsorption, and characterization of clay and bricks manufactured after the sintering process. Conditions: 40 kV and 40 mA.

XRF: Panalytic Minipal2 PW4024 X-ray spectrometer with copper anode. Qualitative study on sawdust residues from the adsorption of Ni (II) ions as a function of time and qualitative study on the adsorbent from the adsorption of Ni (II) as a function of the initial concentration of metal. Characterization of manufactured bricks. Conditions: helium flux, voltage 20 KV, current 5 mA and time 100 s.

UV-Vis: Shimadzu 3600 Plus UV-VIS-NIR spectrophotometer at $\lambda = 722$ nm. Determination of the concentration of Ni (II) on the filtrates obtained in the adsorption tests.

Mechanical tests: Instron 8501 universal servo hydraulic testing machine. MOR determination. Conditions: room temperature, distance between supports 60 mm, speed of application of the load 0.5 mm min^{-1} in displacement control. σ_{str} determination. Conditions: room temperature, load application speed of 8.25 kN min^{-1} in control per load.

AAS: Shimadzu AA 6800 - Flame mode. Determination of the concentration of Ni (II) on the filtrates obtained in the leaching tests.

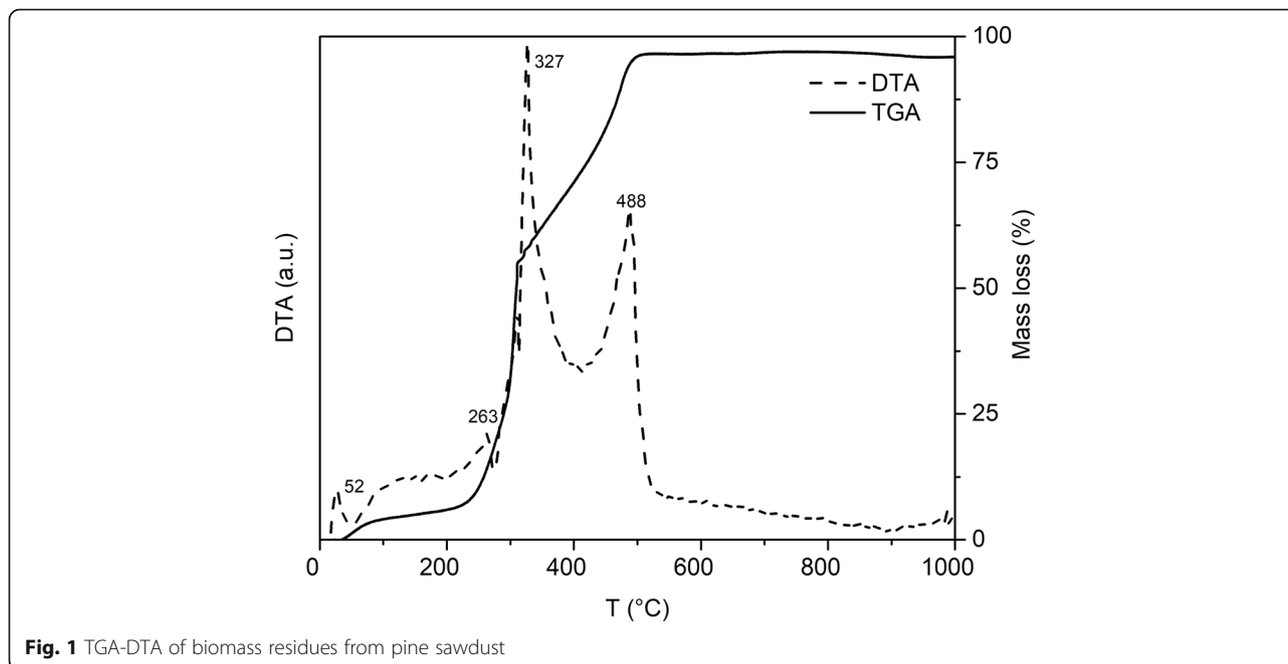
Results and discussion

Bioadsorbent characterization

Thermal analysis by TGA-DTA for pine sawdust is presented in Fig. 1. It is important to know the behaviour of biomass as the temperature rises, since this is closely related to the properties that the brick will reach when sintered.

The DTA diagram shows an endothermic peak at 52°C and a small exothermic peak at 263°C assigned to the loss of water from the sample and the combustion of volatile components, respectively. In addition, in this diagram it is also possible to recognize two exothermic peaks at 327 and 488°C , attributed to the decomposition of hemicellulose (roasted in wood called active pyrolysis), and decomposition of cellulose (active pyrolysis) and lignin (active and passive pyrolysis), respectively [32].

It is possible to observe three major mass losses in the TGA diagram, a first loss up 230°C corresponding to



the loss of moisture and the decomposition of volatile components [33], a second loss up to 311 °C assigned to hemicellulose (to give lower weight compounds molecular, mainly acetic acid), and, finally, a loss of mass up to 502 °C assigned to cellulose and lignin (to finally give CO₂, H₂O and ash). Hemicellulose combustion occurs at lower temperatures due to its linear structure with short side chains. Cellulose and lignin have more complex and stronger structures, with associated dispositions and aromatic compounds, so they have greater resistance to heat.

These results demonstrate that, during the sintering of the ceramic pieces, the biomass added to the clay mix will gradually calcine as the temperature of the oven increases. This will allow the gases formed to diffuse slowly and the fired bricks will not crack. From the data obtained from the TGA analysis, it is possible to obtain an estimated composition for pine sawdust residues. This composition is detailed in Table 1. In particular, the value of inorganic ash was also carried out by the ASTM E 1755–01 standard method and the value obtained, 2.9%, is similar to that obtained by thermogravimetric analysis.

Table 1 Estimated percentage composition of pine sawdust waste from the TGA analysis

Composition	%
Moisture and volatile	7.4
Hemicellulose	47.4
Cellulose and lignin	41.3
Inorganic ash	3.8

Figure 2 shows two SEM images of pine sawdust waste used in adsorption experiments. The images show that pine sawdust has a fibrous and irregular structure with pores smaller than 50 μm and elongated particles [34]. These surface characteristics could make the adsorption of heavy metals possible.

Figure 3 shows the FTIR spectrum of the biomass. FTIR was used for the qualitative analysis of the biomass residues of pine sawdust and to identify the functional groups that could participate in the adsorption of contaminating metals. It is possible to observe the presence of bands corresponding to stretches O-H (3344 cm⁻¹) of hydroxyl groups of phenol, C-H of aromatic methoxide group (2922 cm⁻¹), and vibration of aromatic ring (1508 cm⁻¹), assigned to lignin; stretching C=O (1600 cm⁻¹), and C-OH and C-H (1103 cm⁻¹), assigned to cellulose and lignin; C-H flexion of the methyl group (1327 cm⁻¹), and stretching C-O, C=C and C-C-O (1032 cm⁻¹), assigned to hemicellulose, cellulose and lignin; and asymmetric stretching C-O-C (1158 cm⁻¹) and glycosidic bond (830 cm⁻¹), corresponding cellulose and hemicellulose [35]. The observed peak around 2359 cm⁻¹ of wave-number corresponds to the residual signal of the asymmetric stretch O=C=O of the CO₂ that the target failed to completely eliminate.

Figure 4 shows the XRD diagram of the pine sawdust residue, with the characteristic peaks at 16.2°, 22.4° and 34.8° indicating typical of semi-crystalline cellulose. In addition, a peak at 26.5° corresponding to SiO₂ (quartz) and three other peaks of CaCO₃ (calcite) at 24.5°, 28.2° and 38.2° are observed in the diagram [36].

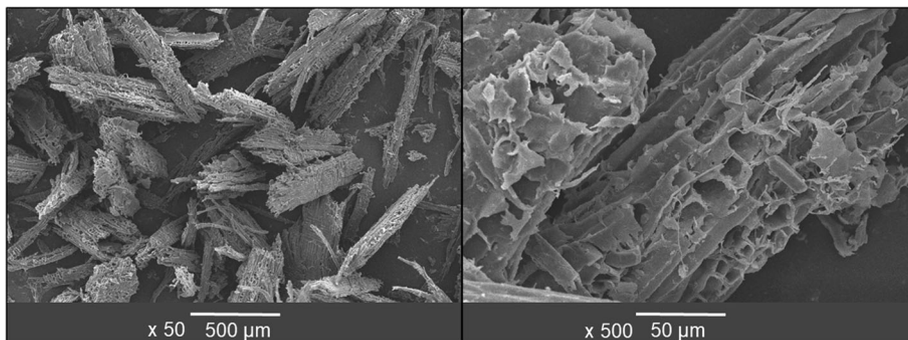


Fig. 2 SEM images (x 50 and x 500) of pine sawdust waste

Qualitative study by XRF of the adsorption of Ni (II) ions as a function of time

Figure 5 shows how the amount of Ni (II) adsorbed increases as the stirring time between the Ni (II) solution and the pine sawdust biomass increases. This process was performed to estimate the optimal contact time for a certain concentration of heavy metal and a certain amount of sawdust. The analysis was performed by XRF on the solids obtained after remaining in contact with the NiCl₂ solution during each of the stirring times studied. Figure 5 shows that Ni (II) adsorption onto pine sawdust increases with contact time because, initially, there is a greater availability of active biomass sites. After a certain time, a plateau is reached which represents a stability in the adsorption of nickel ions. This shows a successive occupation of the active sites of pine

sawdust by the pollutant, which causes, after a time, the adsorption to decrease. According to the results obtained, the stirring time for the adsorption tests was 24 h to ensure greater adsorption.

Effect of the initial concentration metal and dose of bioadsorbent

Ni (II) adsorption tests were carried out by modifying the initial concentration of the metallic solutions, but maintaining the volume of the solution, and studying adsorption on 3 different amounts of biomass, 10, 20 and 40 g L⁻¹. After a 24 h stirring time, the mixtures were filtered and the remaining solutions were analysed to determine their residual nickel concentration. Figure 6a shows the relationship between adsorbent dose, initial adsorbate concentration, and Ni (II) adsorption efficiency.

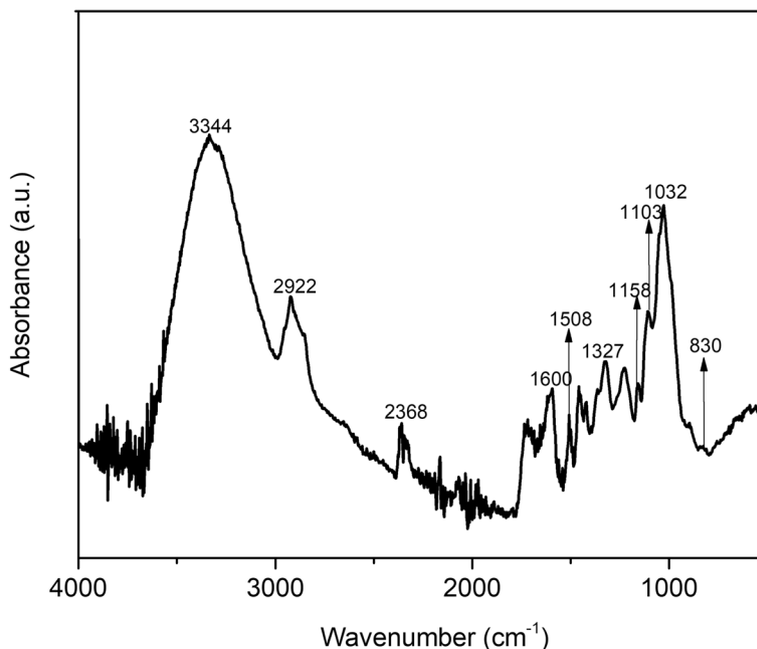


Fig. 3 FTIR of pine sawdust waste

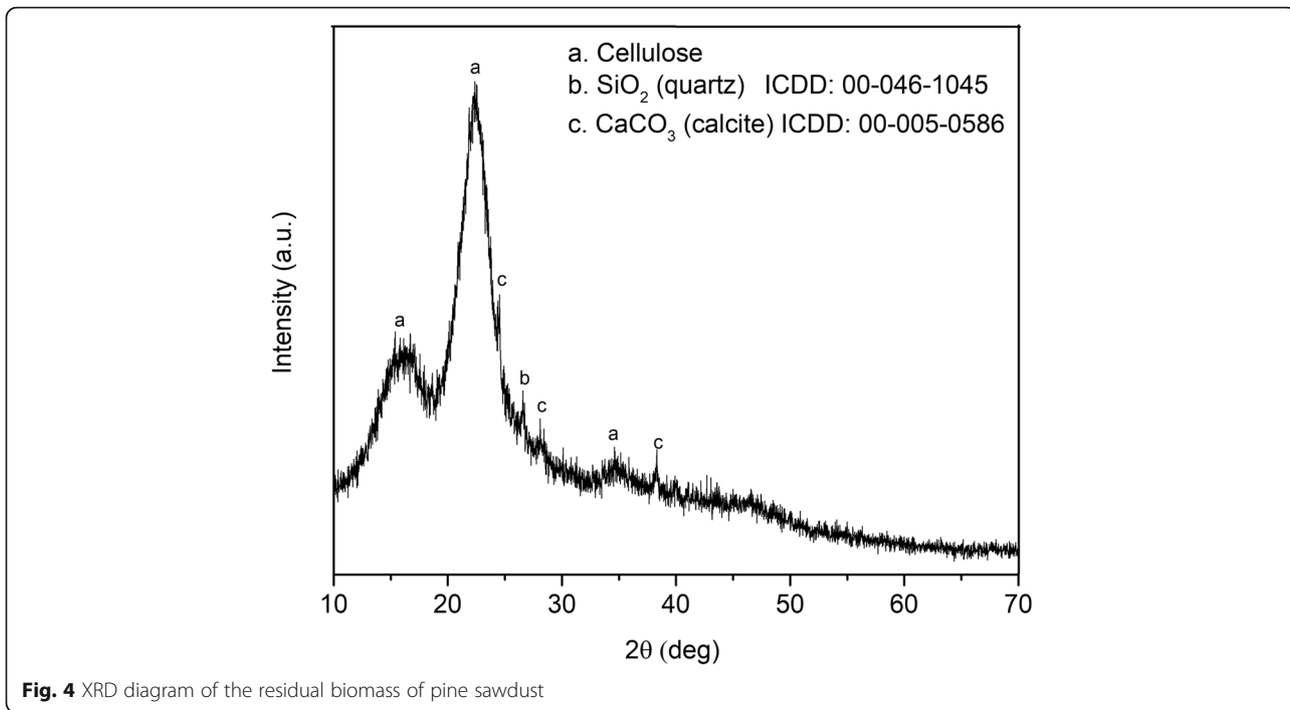


Fig. 4 XRD diagram of the residual biomass of pine sawdust

R values are higher for low NiCl₂ concentrations because there are many binding sites available in pine sawdust biomass but then decrease for all residue doses studied. In Fig. 6b, it is possible to observe that higher initial concentrations of NiCl₂ yield greater adsorption capacity. As the concentration of metal ions increased, the collisions between these and the adsorbent increased. Finally, it seems that at high metal concentrations there is a tendency to

reach a plateau. According to Fig. 6b, in the concentration range studied, at 50 mL of 1 M NiCl₂ ($5.87 \times 10^4 \text{ mg L}^{-1} \text{ Ni}^{+2}$), the adsorption capacity is higher. This value increases when the adsorbent dose of 20 g L^{-1} is used. Furthermore, it is observed in Fig. 6a that when the amount of biomass doubles from 10 to 20 g L^{-1} , the *R* increases 2.5 times, but when the sawdust dose is doubled again, the *R* increases only 1.5 times. Therefore, the adsorbate/

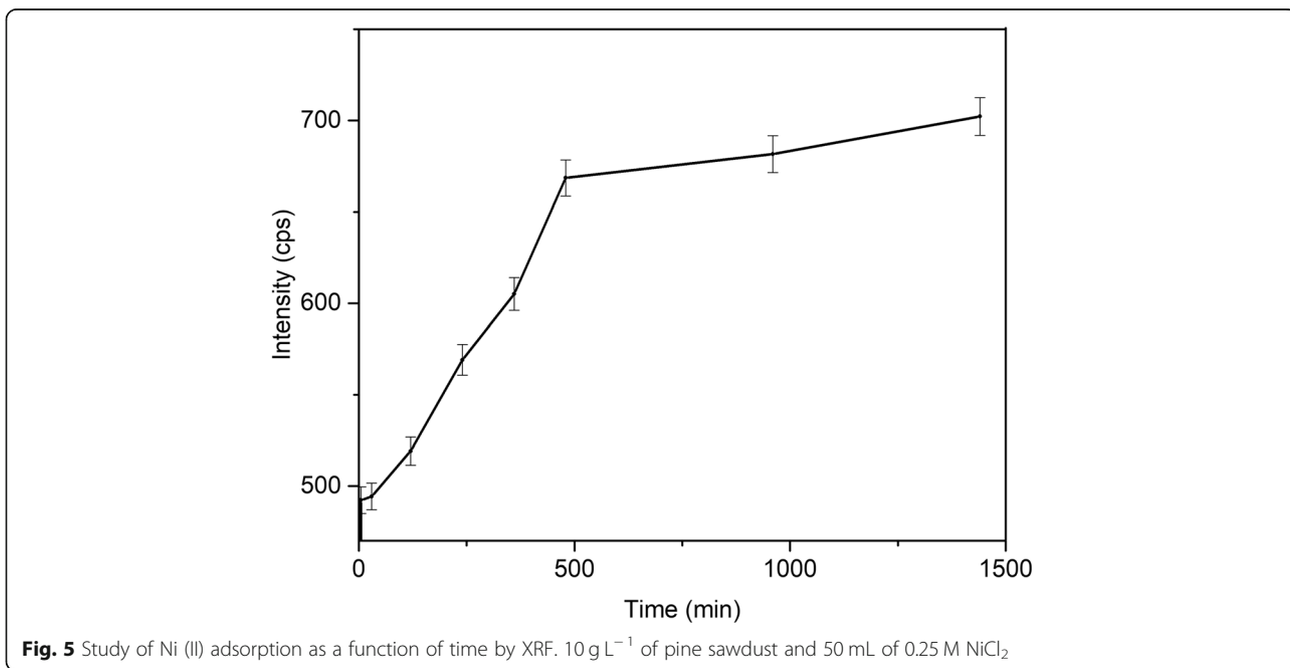


Fig. 5 Study of Ni (II) adsorption as a function of time by XRF. 10 g L^{-1} of pine sawdust and 50 mL of 0.25 M NiCl₂

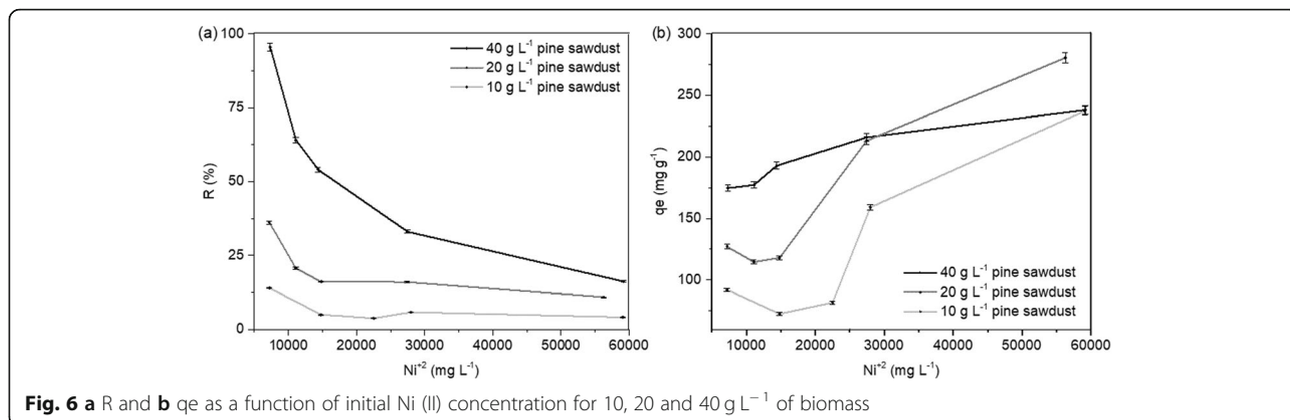


Fig. 6 a R and b q_e as a function of initial Ni (II) concentration for 10, 20 and 40 g L⁻¹ of biomass

adsorbent ratio of 50 mL of 1 M NiCl₂ and 20 g L⁻¹ sawdust was considered the most efficient and was then used in the construction of clay bricks.

Figure 7 shows qualitative XRF analysis of the amount of Ni (II) adsorbed in the solids obtained after contact with 50 mL of NiCl₂ solutions in a concentration range between 0.125 and 1 M NiCl₂ using a dose of 20 g L⁻¹ of adsorbent. As the initial concentration of NiCl₂ increases, the nickel adsorbed on the pine sawdust increases and there is a tendency to reach a plateau that marks the saturation of the biomass. This is observed for 50 mL of 1 M NiCl₂.

In Fig. 8, the FTIR spectrum of pine sawdust is compared with that measured after 24 h of stirring with 50 mL of 1 M NiCl₂ solution. A slight change in the position and intensity of the bands in the biomass residue

can be observed after the metal adsorption, due to the probable complexation or electrostatic interaction reactions and the Van der Waals forces. It is important to highlight the evidence of new bands at 3487, 1606, 717 cm⁻¹ (indicated with * in Fig. 8), visible from the contact of the biomass with 50 mL of 1 M NiCl₂ and belonging to this compound, giving an indication of its adsorption as such in the biomass.

Figure 9 shows the XRD of sawdust after the sorption process with 50 mL of 1 M NiCl₂. In the diffraction diagram it is possible to identify the presence of NiCl₂(H₂O)₂ crystalline phase (indicated with * in Fig. 9).

The biomass electron micrographs after adsorption are shown in Fig. 10. Agglomerates of pine sawdust particles exceed 25 μm. From EDS, the homogeneous presence of

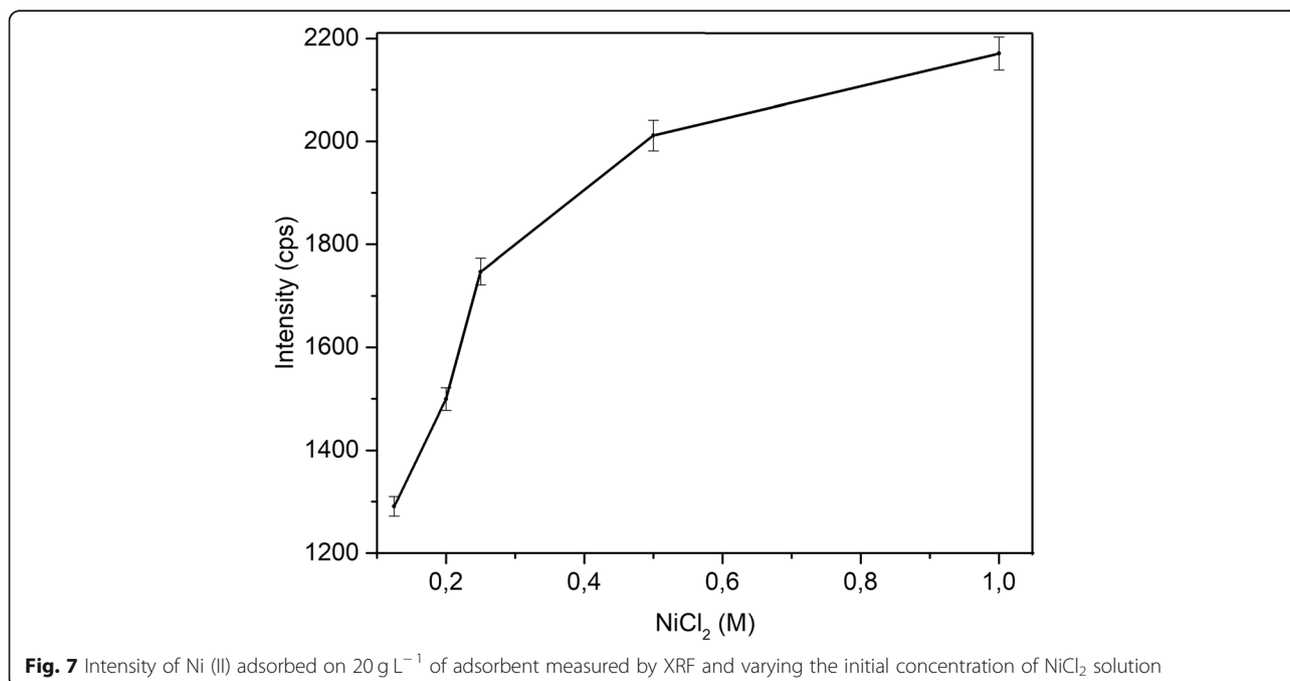
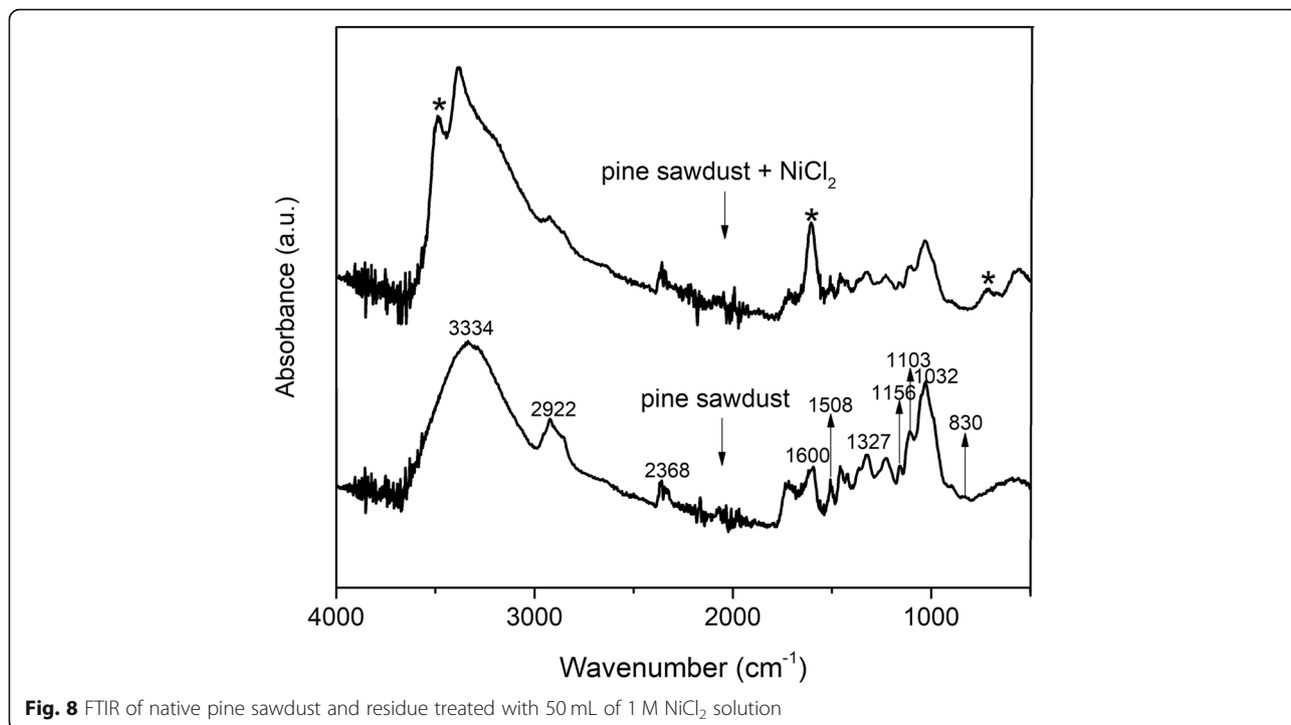


Fig. 7 Intensity of Ni (II) adsorbed on 20 g L⁻¹ of adsorbent measured by XRF and varying the initial concentration of NiCl₂ solution



adsorbed Ni can be identified. Semi-quantitative chemical analysis by EDS for the pine sawdust residues after the adsorption process shows that the percentage composition by mass, without considering the carbon content in the sample, is 22.9% oxygen, 0.2% silicon, 36.7% chlorine and 40.2% nickel.

Characterization and evaluation of the bricks

The clay was provided by a local brickyard and its composition obtained by XRD is shown in Fig. 11.

Figure 12 shows the physical appearance of the manufactured sintered ceramic pieces, (a) without added residue (ARC), and (b) with added 20 vol% of sawdust

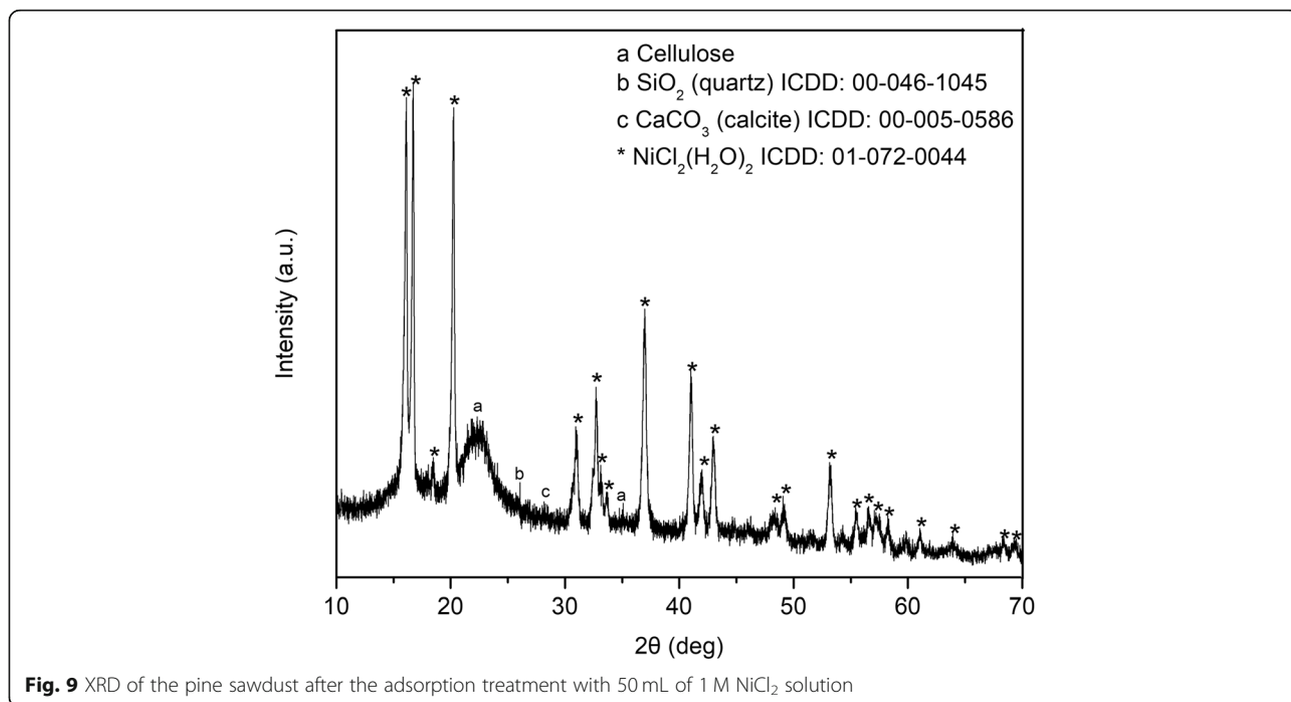
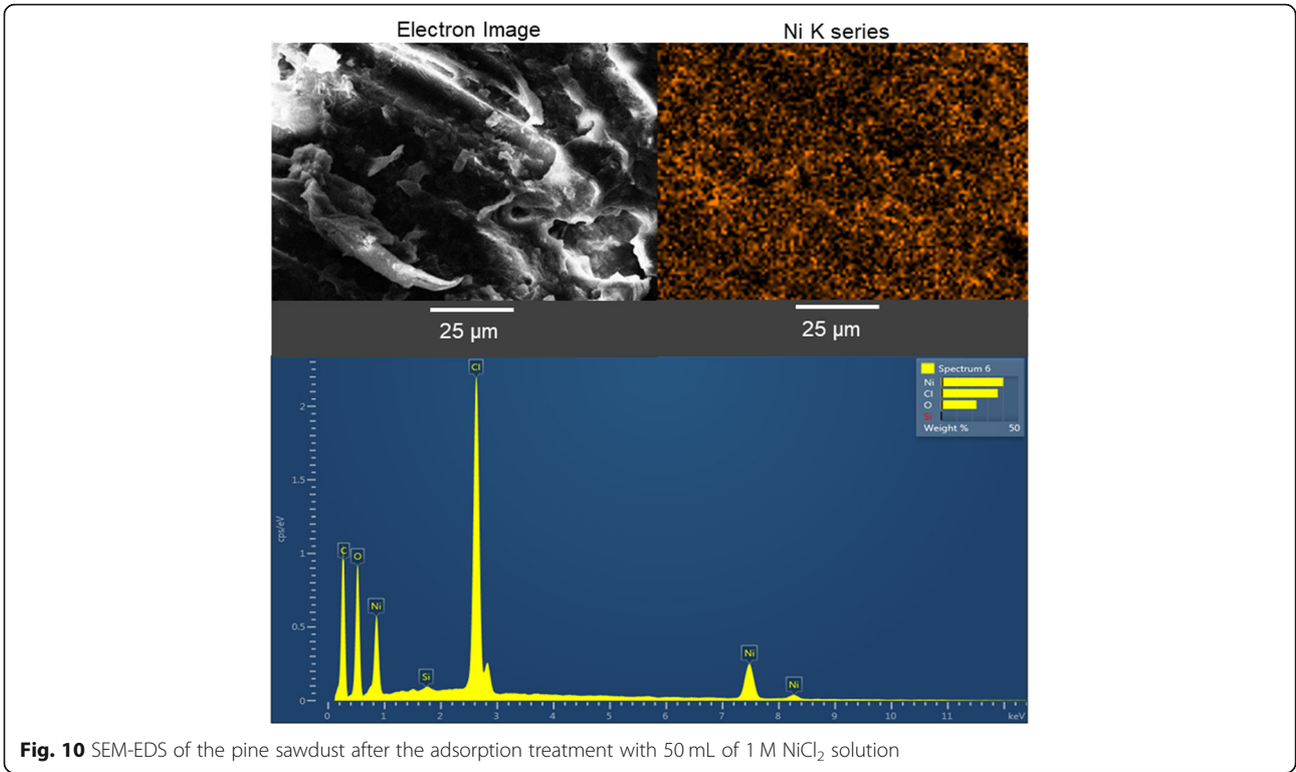


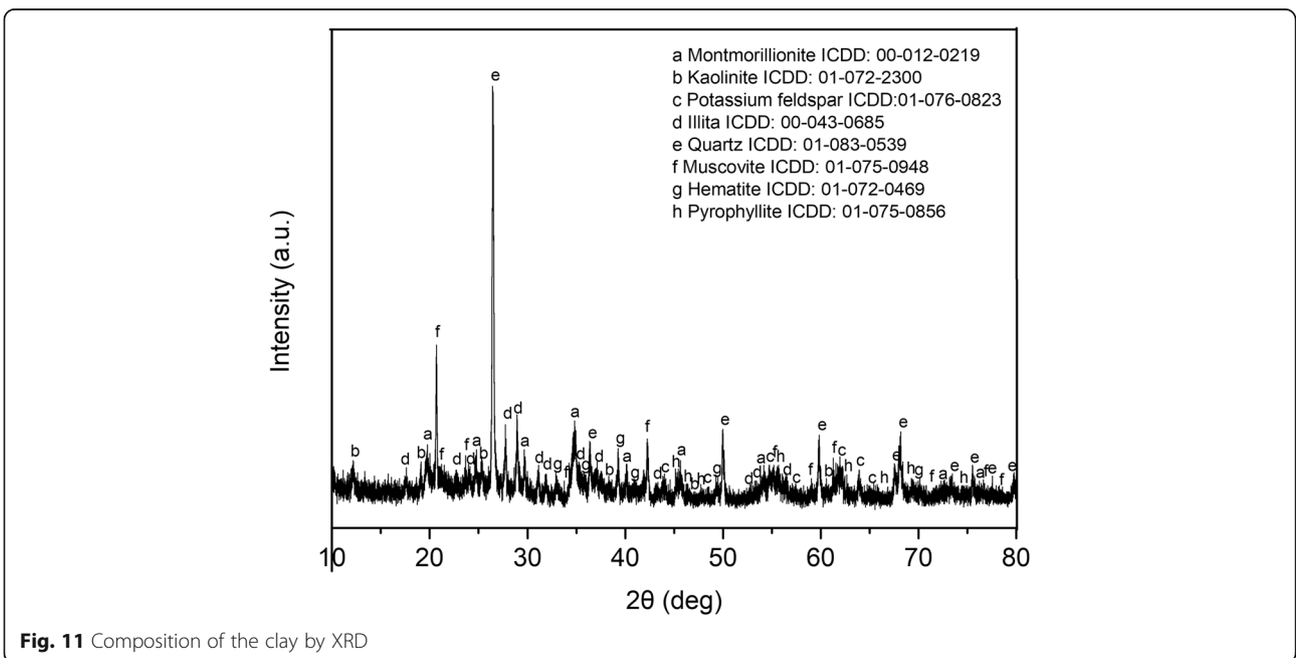
Fig. 9 XRD of the pine sawdust after the adsorption treatment with 50 mL of 1 M NiCl₂ solution



containing nickel adsorbed (AN20). Each AN20 brick weighs approximately 90 g and an estimate of its nickel content from Fig. 6 is 0.8 g. Both pieces are compact, but ARC has a reddish color due to the presence of Fe in the clay, while AN20 has a darker color that could be contributed by high levels Ni (II)

retained on the biomass and then immobilized in the cooked brick.

Figure 13 shows the XRD obtained for AN20 after cooking. It is possible to observe the presence of SiO₂ together with peaks corresponding to numerous secondary phases because the clay used was obtained from



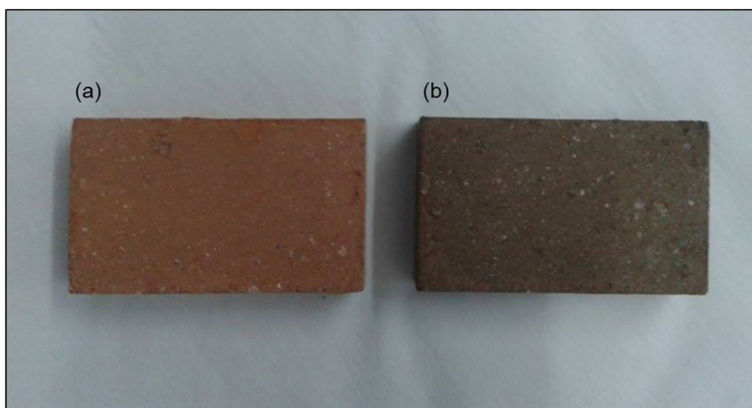


Fig. 12 Macroscopic appearance (a) brick without added residue (ARC), and (b) brick with added 20 vol% of sawdust containing nickel adsorbed (AN20)

natural quarries. Furthermore, it is possible to observe peaks at 37°, 43°, 53° and 63° corresponding to NiO.

Figure 14 shows the XRF diagrams of the AN20 and ARC bricks together with a commercial brick (COM). The presence of lines at 7 and 8 keV corresponding to nickel (NiKa and NiKb respectively) corroborates that the metal remains in the clay matrix after the sintering process.

The EDS images of powder AN20 are shown in Fig. 15. These images allow to corroborate the presence and immobilization of Ni after the sintering process. Semi-quantitative chemical analysis by EDS for AN20 shows that the percentage composition by mass, without considering the carbon content in the sample, is 42.0%

oxygen, 0.8% sodium, 1.1% magnesium, 11.5% aluminum, 27.9% silicon, 2.3% potassium, 1.8% calcium, 0.6% titanium, 8.3% iron and 3.8% nickel.

Table 2 shows the average values obtained from LOI and the parameters determined from the apparent porosity test of the manufactured bricks. The LOI was greater for AN20 than for ARC due to the combustion, during sintering, of the aggregate biomass. This combustion of sawdust waste generates a greater apP observed in ceramic pieces made from clay and the addition of 20% biomass. Higher values of apP and H₂OAbs determine lower values of apD and apsW for AN20. In addition, the mechanical properties of the AN20 and ARC ceramic matrices were evaluated and the average values obtained are presented

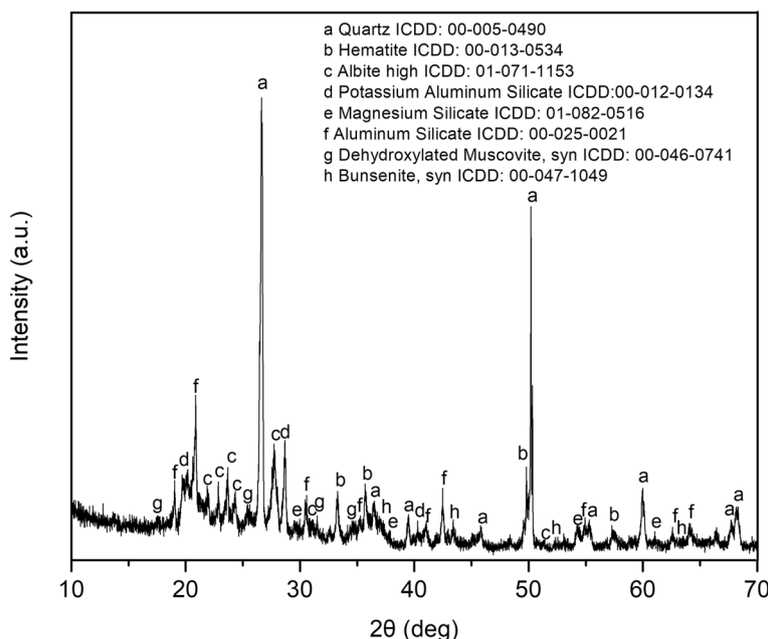


Fig. 13 XRD of AN20 after calcination at 950 °C

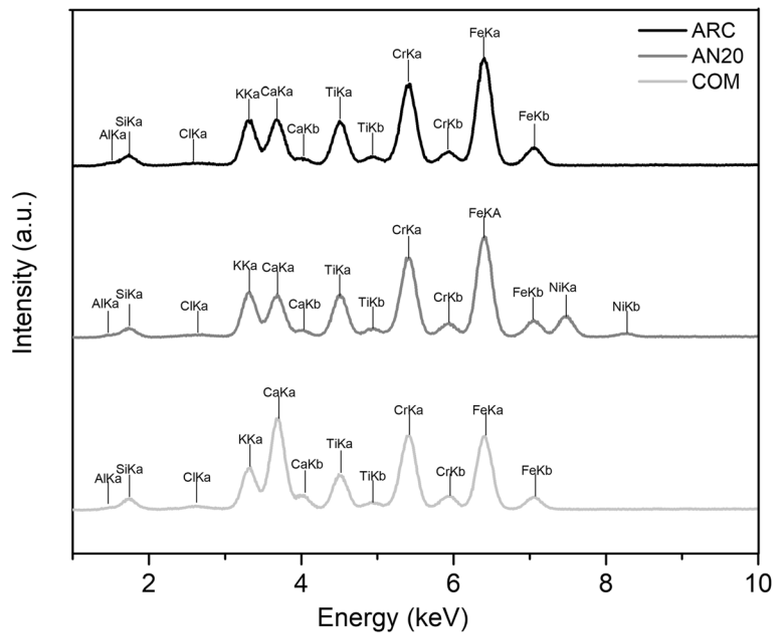


Fig. 14 XRF of ARC, AN20 and COM after calcination at 950 °C

in Table 2. The average values of σ_{str} obtained are within the commercially required standards according to IRAM 12566–1:2005, “the bricks must have a characteristic compressive strength equal to or greater than 4 MPa”. In addition, in Argentina, CIRSOC 501R - 2007 regulates the use in construction, and sets a lower limit of 5 MPa. These

σ_{str} values were lower for AN20 than for ARC due to aggregate biomass. This same behavior is observed in MOR. According to the literature [37], the MOR varies between 10 and 30% of the σ_{str} , so AN20 still has a MOR value within the appropriate parameters of the market (MOR = 23% σ_{str}).

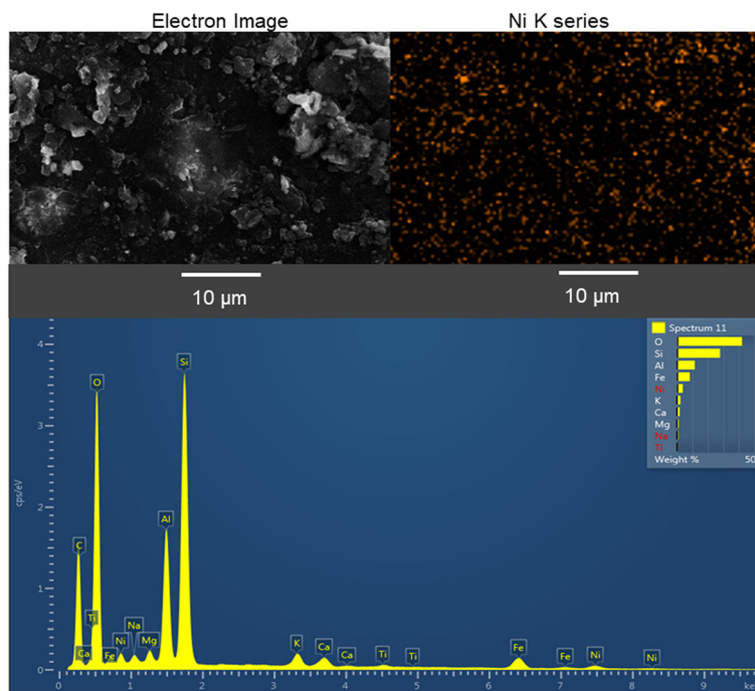


Fig. 15 SEM-EDS of AN20 after calcination at 950 °C

Table 2 Properties of sintered ceramic pieces and a commercial brick

Brick	apD (g cm ⁻³)	apsW (g cm ⁻³)	apP (%)	H ₂ OAbs (%)	apV (cm ³)	LOI (%)	σstr (MPa)	MOR (MPa)
ARC	1.994 ± 0.001	1.997 ± 0.001	26.2 ± 0.2	13.1 ± 0.1	10.12 ± 0.06	8.6 ± 0.1	24.3 ± 0.1	2.8 ± 0.3
COM	1.37 ± 0.01	1.38 ± 0.01	44.2 ± 0.4	32.15 ± 0.05	13 ± 1	–	2.9 ± 0.1	1.7 ± 0.1
AN20	1.854 ± 0.003	1.857 ± 0.003	31.28 ± 0.02	16.85 ± 0.04	11.242 ± 0.006	13.9 ± 0.9	10.5 ± 0.1	2.41 ± 0.01
AZ20	1.84 ± 0.02	1.85 ± 0.02	31.4 ± 0.7	17.0 ± 0.5	10.58 ± 0.02	12.4 ± 0.7	9.7 ± 0.1	1.82 ± 0.07

Very similar values of these properties were obtained for the brick manufactured from sawdust with Zn (II) adsorbed (AZ20) [38]. Also, in Table 2, the values for a COM are presented. AN20 has superior physical and mechanical characteristics compared to COM.

A common concern when adding contaminated waste to fired brick is the possible migration of these contaminants into the environment. In this sense, it is important to evaluate the leachability of the manufactured AN20 ceramic matrices. National Law 24,051 and its regulatory decree establish that leachates in dangerous products must be analyzed in accordance with TCLP. The measurements were obtained in triplicate following the protocol (using fragments of AN20 and a solution of pH 4.93 ± 0.05 as extraction solvent, determined from the sample’s alkalinity according to TCLP). Table 3 shows the average Ni (II) result obtained in the leached liquid. This value was less than below detection level, that is, less than 0.05 mg L⁻¹ of nickel and was compared with the maximum concentrations allowed according to Decree No. 2020 of 2007 of Law No. 2214 of the City of Buenos Aires (CABA), US EPA Code of Federal Regulations 2012 and the regulations in force in Spain and China. Therefore, the metal concentrations obtained by TCLP of AN20 are below the established maximum. A previous investigation has a similar behavior in AZ20 bricks [38]. These tests demonstrate that during the brick firing process, biomass residues are destroyed and heavy metals could encapsulate in the clay matrix, form oxides, and interact with silicon, iron, or aluminum oxides in the clay to form minerals stable, according to the literature [29, 39]. The phase transformations and the evolution of the heavy metals during the firing of the ceramic pieces determine the fixation of the contaminants. However, the mechanism of immobilization of heavy metals during the sintering process has not yet been understood [40].

The calculation of the efficiency of the retention is made according to the Eq. (3) [41].

$$ER (\%) = \frac{(Mmib - Mml) \times 100}{Mmib} \tag{3}$$

where, ER = efficiency retention of metal in the brick (%), Mmib = mass of metal incorporated in AN20 and Mml = mass of metal extracted in the leachate. Mmib is estimated from the nickel mass adsorbed by pine sawdust in the ratio of highest efficiency selected and corrected for the mass of sawdust added as pore-forming agent in AN20. Mml is estimated from the nickel concentration in the leachate corrected for the mass of the entire brick. ER for AN20 bricks is almost 100%.

Therefore, it is possible to immobilize Ni (II) in fired clay bricks and thus reuse the pine sawdust previously used as an adsorbent of these contaminants. In addition, a common paint such as latex, could act as an additional “barrier” that guarantees the non-leaching of Ni from inside the brick [28].

Conclusions

Pine sawdust generated by the timber industry was evaluated as an adsorbent of synthetic Ni (II) solutions. This residue, without a specific use at present, showed the ability to retain the ions of the heavy metal studied. In this way, pine sawdust constitutes a simple and low-cost system for wastewater treatment. The adsorption tests show that the most efficient adsorbate/adsorbent combination was found to be 50 mL of 1 M NiCl₂ and 20 g L⁻¹ of sawdust.

The novelty is to use this new contaminated biomass in the manufacture of porous clay bricks, immobilizing heavy metals and minimizing the impact on the environment.

The sintered bricks presented values of apP and mechanical characteristics within the market requirements. In addition, TCLP tests show an ER close to 100%, which shows an excellent immobilization of Ni (II) within the ceramic pieces.

Table 3 Results of the TCLP standard procedure of AN20 bricks. Comparison with limit concentrations of Ni (II)

Heavy metal	TCLP of AN20 (mg L ⁻¹)	Decree No. 2020 of 2007 of Law No. 2214 of the City of Buenos Aires (CABA) (mg L ⁻¹)	US EPA Code of Federal Regulations 2012 (mg L ⁻¹)	UNE-EN 12457-4 (Spanish standard) (mg L ⁻¹)	GB 5085.3-2007 (Chinese standard) (mg L ⁻¹)
Ni	< 0.05	5.0	Not Given	2.0-0.4	5.0

With these results, it would be possible to respond to the needs of the industry that generates contaminated effluents and of the agribusiness that generates biomass waste without a specific use and considerably reducing the negative implications on the environment.

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Authors' contributions

DS carried out the experimental work, processed the data obtained and wrote the document. NQ participated in the discussion of the results. SG collaborated with the experimental work and participated in the discussion of the results. RP collaborated with the experimental work and participated in the discussion of the results. AC coordinated the investigative work and wrote the document. The author(s) read and approved the final manuscript.

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Availability of data and materials

All data generated or analyzed during this study are included in this published article or request the corresponding author.

Competing interests

The authors declare they have no competing interests.

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