Experimental and theoretical study of Co sorption in clay montmorillonites

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Abstract
Montmorillonite (MMT) clays are 2:1 layered structures which in natural state may allocate different hydrated cations such as M-nH2O (M = Na, Ca, Fe, etc) in its interlayer space. Depending on the capability for ion sorption, these materials are interesting for environmental remediation. In this work we experimentally study the Co sorption in a natural Na-MMT using UV-visible spectrometry and XRD on semi-oriented samples, and then analyze the sorption ability of this clay by means of ab initio calculation performed on pristine MMT. The structural properties of Na-MMT and Co-adsorbed MMT, and the hyperfine parameters at different atomic sites were analyzed and compared with the experimental ones for the first, and for the case of the hyperfine parameters, presented for the first time for the last. The theoretical predictions based on total energy considerations confirm that Co incorporation replacing Na is energetically favorable. Also, the basal spacing d_{001} experimentally obtained is well reproduced.

1. Introduction
From the beginning of the industrial revolution, environmental pollution by heavy metals appeared as a direct consequence, and its reversal represents a challenge for modern life. In this scenario pollutants generation and the deposition controls constitute a very important topic due to the negative effects that these elements could cause both on the human health and the environmental quality. Among the mechanisms used for heavy metal removal from effluents, ion exchange allows the use of low cost and environmental friendly materials, like clay minerals and its derivatives (modified clays that enhance the sorption process of the sorbent material) [1–3]. Among them, Montmorillonite (MMT), a 2:1 clay mineral, is one of the most promising candidates for decontamination and disposal of wastewater with high-level heavy metal cation content, due to its relatively high specific area and cation exchange capacity [4, 5]. Cobalt is one of the toxic metals affecting the environment. It is widely present in effluents and is originate from different human activities [6]. Its origin is connected with urban storm-water runoff coming from electronic, electroplating, paints and pigments, mining, metallurgical and nuclear industries (principally from wastes released from pressurized water nuclear power reactors and from medical applications) [6–10]. Because it can affect human health and the environment its removal from effluents is essential [6–11].

Some studies were performed testing the efficiency of Co sorption in natural and modified MMT [4, 12–24]. The results indicated that the MMT systems can sorb and retain Co depending on the system itself and the physical and chemical conditions. Although the main expected sorption mechanisms in clays are cationic exchange and surface complexation, sorption mechanisms of Co in MMT are still not clear.

Between the experimental techniques used to enlighten the mechanisms involved in metal/clay complexation, Nuclear Magnetic Resonance (NMR) became an important tool due to its high sensitivity to the local environment of the atoms and the chemical bonds. Clay layers were previously studied with this technique.
using $^{25}$Mg, $^{27}$Al and $^{29}$Si as probe atoms [25–32], whereas for the clay interlayer and swelling behavior, the studies were performed using $^{113}$Cd, $^{133}$Cs, and $^{23}$Na probes [33–37].

Computational quantum simulations have proven to be a suitable tool for understanding the microscopic processes that conduct to the physical and chemical properties of materials. Beyond the simplifications that are made in comparison to the complexity of real materials, they can lead to a better understanding of individual processes involved. Recently ab-initio calculations within the Density Functional Theory (DFT) were used to investigate the mechanisms involved in metal sorption on clays [38–42] and especially in MMT [43–51]. Within this theory some well-tested codes have implemented the determination of the hyperfine parameters [52, 53]. In the present work a combined DFT and experimental study for Co sorption in MMT is reported. The calculations are discussed in term of the energetic and structural features accompanying the Na to Co interlayer exchange in pristine MMT. The electronic properties of MMT and Co sorbed MMT are also analyzed.

The paper is organized as follows: in section 2 a brief description of the system is given. In section 3, the experimental methods and the results for the determination of the Co$^{2+}$ sorption are presented. Section 4 describes the models and computational details of the ab-initio calculation. In section 5 the results of the calculations are shown. This section is divided in three subsections: a-the results concerning the energy formation and cell elongation, b-the predicted hyperfine characterization, and c-the comparison with experimental data obtained from literature. Finally, section 6 is reserved to the conclusions.

2. The system under study

The MMT structure could be described as a 2:1 phyllosilicate constituted by a laminar structure with an elongated cell along the c axis having a triclinic unit cell with space group P1. The layers consist of an octahedral sheet linked with two tetrahedral sheets sharing apical oxygen atoms. The $[(Al/Mg)O_6]$ units form the octahedron layer sandwiched by two layers of $SiO_4$ tetrahedrons. In the interlayer region different cations can be allocated, such as Na and Co, with different hydration degrees. The interactions between layers are supposed to be weak. Pristine Na-MMT have as chemical formula $Na[(MgAl_2O_4)(OH)_2(Si_6O_{18})_2]+m(H_2O)$. Co-sorbed MMT structure, labeled as Co-MMT, was generated by replacing the Na$^+$ with Co$^{2+}$ ions linked with the corresponding H$_2$O molecules, as will be detailed in the next sections. In figure 1 it can be seen the starting structure before internal atomic position relaxation for the case $m=4$ with one Na per unit cell. A second (Na + 4H$_2$O) unit and some $[(Al/Mg)O_6]$ units in the neighbor unit cell in the a axis direction are depicted for clarity.
3. Experimental

The Co\(^{2+}\) sorption experiments were performed using a natural Na-MMT from Río Negro State, Argentina. The sorption studies were performed in batch conditions (V = 25 ml), using a Co\(^{2+}\) initial concentration of 100 mg l\(^{-1}\) (obtained after dissolving the corresponding masses of CoCl\(_2\)6H\(_2\)O, provided by Sigma Co.), a solid/liquid ratio of 1 g l\(^{-1}\), pH = 6 and a contact time of 24 h. Sorption experiments were performed in duplicate.

After the sorption experiment, the liquid and solid phases were separated by centrifugation. The solid phase was dried at 60 °C and stored in a desiccator for its posterior characterization. The remaining Co\(^{2+}\) concentration in the liquid phase was determined and from this value the sorbed Co\(^{2+}\) was obtained. Co\(^{2+}\) determination was performed using a UV-visible spectrometer, following the methodology proposed by Sandell\[54\].

The structure changes of the solid phase after the Co-sorption experiments were determined by XRD technique. Partial x-ray diffraction patterns of semi-oriented clays were collected using a Philips 3020 equipment, operating to 35 mA and 40 kV, with Cu K\(_\alpha\) radiation, from 2° to 10° (2 theta), with 0.02° stepping angle and 10 s/step.

The results of the Co\(^{2+}\) sorption experiments confirm that Co\(^{2+}\) ions were sorbed into the samples. The sorption percentage for this cation resulted as high as 80 \(\pm\) 2%. Before and after the sorption process, XRD analyses were carried out for oriented samples in order to determine the d\(_{001}\) parameter (figure 2). The obtained results also indicate that after sorption experiments Co incorporation in the interlayer was produced, due to an expansion in the c-axis direction. For the determination of the d\(_{001}\) distance, Voight functions were fitted to the XRD spectra. In the case of the sample before the Co\(^{2+}\) sorption procedures, two Voight functions were used to take into account the high asymmetry of the corresponding peak. The determined d\(_{001}\) distance for Co-MMT was 15.71 \(\pm\) 0.06 Å while the corresponding for the natural Na-MMT was 12.58 \(\pm\) 0.04 Å and 13.55 \(\pm\) 0.07 Å, resulting in a high enlargement of the interlayer spacing.

4. Modeling and computational details

Natural samples of MMT may have different low concentrations of impurities (such as Fe, Ca), and also cation exchange between Si, Al and Mg. The effect of such defects in macroscopic structures is out of the scope of this study. The starting atomic positions for the \textit{ab-initio} calculations for the Na-MMT and Co-MMT in combination with the different hydration states were taken from the work of Pirillo \textit{et al}\[46\] with lattice parameters a = 5.2584 Å b = 8.7669 Å c = 12.8511 Å and \(\beta = 95.3443\). Co-MMT structure was generated by replacing the Na with Co atoms. For the initial m-H\(_2\)O units atomic positions we considered starting from symmetric distributions with the H pointing outward the corresponding cation as can be seen in figure 1.

Following the ideas of Berghout \textit{et al}\[45\] and the studied performed by Pirillo \textit{et al}\[46\] for the simulated Na-MMT, we considered three different Na concentration/hydration states: \(a\)-One Na\(^+\) atom surrounded by four H\(_2\)O in the unit cell (NaM1); \(b\)-Two Na\(^+\) atoms, each one of them surrounded by two H\(_2\)O (NaM2); and \(c\)-Two Na\(^+\) atoms, each one of them surrounded by four H\(_2\)O (NaM3). For the simulated Co-MMT we considered the
case of two Co$^{2+}$ atoms, each one surrounded by six H$_2$O in the unit cell (CoM1). For each one of the models, refined atomic positions and lattice constants were obtained as explained below.

All the simulations of the electronic structure were performed within DFT. So, as starting step the density functional first-principles simulations were performed based on the numerical Linear Combination of Atomic Orbitals (LCAO) method as implemented in the Siesta Code [53]. The calculations have been carried out within the Generalized Gradient Approximation (GGA), using the functional parametrized by Perdew, Burke and Ernzerhof (PBE) to simulate the electronic exchange and correlation potentials [55]. All the core electrons were replaced by ab-initio norm-conserving pseudopotentials generated using the Troullier-Martins scheme [56], in the Kleinman-Bylander fully non-local separable representation [57]. The pseudopotentials were generated scalar-relativistically. The size of the basis set chosen was double-$\zeta$ plus polarization for the valence states of all the atoms. An equivalent plane wave cutoff of 200 Ry was used to represent the charge density. During geometry optimizations, a 7 Å k-grid cutoff for all the Brillouin zone integrations was used. For the structural optimizations, the atomic positions were allowed to relax until the maximum component of the force on any atom was smaller than 0.01 eV Å$^{-1}$, and the maximum component of the stress tensor was smaller than 0.0001 eV Å$^{-3}$. Finally, the whole internal atomic positions and lattice parameters a, b, c, \(\alpha\), \(\beta\) and \(\gamma\) were relaxed according to the method described above.

Once optimized the atomic positions and cell parameters with the above described method, they were used as input files for the codes available in Quantum ESPRESSO package [52]. The electronic correlation and exchange terms were calculated using the same GGA-PBE functionals. Interaction of valence electrons with nuclei and core electrons were treated by the projector augmented-wave (PAW) method [58]. The energy cutoff for the plane-wave basis set was fixed to 60 Ry and the electron density of 550 Ry. A Monkhorst and Pack [59] k-point grid of 2 \(\times\) 2 \(\times\) 2 was used to sample the first Brillouin zone of the cells. The simulation of NMR parameters was performed with the gauge including projector augmented wave (GIPAW) approach [60, 61].

## 5. Results and discussion

### 5.1. Formation energy and cell elongation

The calculated lattice parameters and \(d_{001}\) values after optimization of the initial parameters of the different cell considered are listed in Table 1. From this table and figure 2 it can be seen that the experimental and theoretically predicted \(d_{001}\) for the Co-MMT are in good agreement. The lattice was strongly expanded in the c direction after the procedure from an initial value of \(c = 12.851\) Å to the final \(c = 15.842\) Å, and the cell stress are nearly zero. With the final lattice constants, the resulting \(d_{001}\) resulted 15.81 Å, which should be compared with the experimentally obtained \(d_{001exp-Co} = 15.5 \pm 0.6\) Å. Regarding the Na-MMT structures, the resulting \(d_{001}\) obtained values are: for NaM1 \(d_{001NaM1} = 12.89\) Å; for NaM2 \(d_{001NaM2} = 12.74\) Å; and for NaM3 \(d_{001NaM3} = 14.19\) Å. These results can help to distinguish which model is closer to the experimental sample. The \(d_{001}\) for the case of two Na atoms with 8 H$_2$O (4 H$_2$O for each Na) NaM3 is much higher than those obtained in this work 12.58 \pm 0.4 Å and 13.55 \pm 0.07 Å. So, NaM3 does not reproduce well the sample before the Co$^{2+}$ sorption. Regarding the models that take into account 4 molecules of H$_2$O, NaM1 and NaM2, on the basis of the \(d_{001}\) it is not easy to reveal the more appropriate model. Both structures give \(d_{001}\) values close to one of the experimentally obtained 12.58 \pm 0.4 Å, thus describing well enough part of the Na-MMT sample, but strictly comparing the obtained numbers, together with the obtained lattice parameters, the NaM2 is closer. It must be mentioned that in real samples coexistence of different hydration states can occur. This effect have been observed and discussed in previous works [63]. As it was mentioned, in our samples two Voight functions were needed to take into account the asymmetry of the XRD peak. The site with \(d_{001} = 13.55 \pm 0.07\) Å was well reproduced for any of these models.

On the other hand, the comparison of the predicted Na-MMT and Co-MMT \(d_{001}\) shows that the Cobalt incorporation in the MMT lattice produces an expansion of the \(d_{001}\) plane, shifting the \(d_{001}\) about 3.23 Å, close

### Table 1. Lattice parameters, \(d_{001}\) [Å] and angles [°] obtained from the structure relaxation for the different configurations. (*) data from this work.

| Experimental parameters for Na-MMT from [46] | a  | b  | c  | \(\alpha\) | \(\beta\) | \(\gamma\) | \(d_{001}\)Na-MMT | \(d_{001}\)Co-MMT |
|---------------------------------------------|----|----|----|----------|----------|----------|------------------|------------------|
| NaM1 (1 Na + 4 H$_2$O)                     | 5.258 | 8.767 | 12.851 | 90 | 95.3 | 90 | 12.60, 12.58 \pm 0.4 Å (*) | 15.5 \pm 0.6 Å (*) |
| NaM2 (2 Na + 4 H$_2$O)                     | 5.256 | 8.767 | 12.795 | 90 | 95.3 | 90 | 12.74 | 14.19 |
| NaM3 (2 Na + 8 H$_2$O)                     | 5.210 | 8.857 | 14.191 | 90 | 89.1 | 90.0 | 12.74 | 14.19 |
| CoM1 (2 Co + 12 H$_2$O)                    | 5.258 | 8.930 | 15.842 | 88.2 | 93.3 | 89.9 | 15.5 | 15.81 |
to that determined experimentally. So, from this correspondence it can be inferred that Co sorption in MMT incorporates the Co atom in the interlayer of the MMT clay, and not as a superficial adsorption, being this fact up to now not discarded.

In order to quantitatively evaluate if the Co sorption is favored, total energies obtained from ab-initio calculation for MMT system with Na and Co performing using the GIPAW code with and without spin polarization were performed. For the case of Co-MMT, the spin-polarized case was that with lower energy, so all the energy differences are calculated referred to the spin-polarized case. We considered the formation energy and the bond energy as:

\[
\Delta E_{\text{form}} = E_{\text{lat}} - (E_{\text{layer}} + 2E_X + nE_{\text{H}_2\text{O}})
\]

\[
\Delta E_{\text{bond}} = E_{\text{total}} - (E_{\text{layer}} + E_{\text{interlayer}})
\]

Where \( X \) stands for Na or Co. For the calculation of the total energy \( E_X \), there were computed the corresponding to the bulk compounds, divided by the number of atoms of the structure. For the \( E_{\text{H}_2\text{O}} \) term, the total energy of the free molecule was calculated, and \( n \) represents de number of molecules for each case. For the \( E_{\text{layer}} \) term, it was computed the total energy of the crystalline structure but without the atoms of the interlayer (figure 3), and vice versa for the \( E_{\text{interlayer}} \) term. In MMT the layers are negatively charged, so the presence of the cation in the interlayer stabilizes the structure. Then, the \( E_{\text{bond}} \) can give information of the charge stability. The resulting energies using equations (1) and (2) are shown in table 2. It can be seen that the Co incorporation related to the formation of the Co-MMT phase is favored with respect to the Na incorporation in the Na-MMT system, and the energy related with the charge stability remains the same. So, it can be concluded that Co-Na exchange is energetically favored.

5.2. Hyperfine characterization

As it is known the external magnetic field used in NMR experiments at probe sites are normally shielded by a shielding coefficient which depends on the local environment (\( \sigma_{\text{sample}} \)). The determined experimental chemical shift is related with the isotropic shielding \( \sigma_{\text{sample}} \) and could be compared with calculated ones taking into account the standard expression:

\[
\delta_{\text{chem}} = \sigma_{\text{Ref}} - \sigma_{\text{sample}}
\]

Starting from the optimized cell parameter, the NMR parameters were evaluated on each atomic site using the GIPAW code. The obtained magnetic shielding, the mayor component of the electric field gradient tensor represented in the local principal axis \( V_{zz} \) and the corresponding asymmetry parameter \( \eta = \frac{V_{xy} - V_{yx}}{V_{zz}} \) are summarized in table 3. Regarding the O atoms, it can be seen in figure 1 that the SiO\(_4\) tetrahedra are oriented in such a way that each tetrahedron has one basal plane approximately parallel to the interlayer, so, these are the oxygen atoms capable to bond with the Na/Co + nH\(_2\)O units, although just one or two of them will effectively
act as bonding atom. On the other hand, on the opposite vertex of the tetrahedron the corresponding atom is

The results for the last group are not shown. As it is known, the hyperfine parameters (HP) are highly sensitive to local distortions due to their dependence on the charge density near and at the nuclei positions. So, in perfect symmetric tetrahedron or octahedron units it should be expected similar values in the

| Atom | NaM1 | NaM2 | CoM1 |
|------|------|------|------|
| Na1  | 569.4| 562.73| —   |
| Na2  | —    | —    | —    |
| Co1  | —    | —    | —    |
| Co2  | —    | —    | —    |

| Atom | Na1  | NaM2 | CoM1 |
|------|------|------|------|
| Al1  | 541.66| 541.08| 541.75 |
| Al2  | 544.81| 538.02| 543.54 |
| Al3  | 540.09| 541.07| 541.85 |
| Al4  | 542.84| 538.04| 544.74 |
| Mg1  | 546.93| 551.82| 551.73 |
| Mg2  | 553.55| 555.33| 556.22 |
| Si1  | 464.15| 428.16| 426.21 |
| Si2  | 426.28| 432.81| 424.46 |
| Si3  | 425.98| 429.18| 441.61 |
| Si4  | 413.55| 430.75| 424.74 |
| Si5  | 426.64| 429.18| 434.85 |
| Si6  | 415.71| 430.75| 421.53 |
| Si7  | 409.22| 428.15| 436.34 |
| Si8  | 439.24| 428.15| 414.26 |

| Atom | Naphal | Optical | OH1 |
|------|--------|--------|-----|
| O1   | —      | 9.03   | 9.07 |
| O2   | 194.65 | 177.54 | 190.04 |
| O3   | 169.81 | 1.4    | 87.78 |
| O4   | —      | 44.79  | 87.12 |
| O5   | 162.68 | 121.1  | 152.11 |
| O6   | 134.49 | 121.8  | 151.21 |
| O7   | 164.34 | 182.02 | 167.25 |
| O8   | 160.59 | 163.31 | 160.71 |
| O9   | 162.31 | 181.94 | 169.77 |
| O10  | 162.5  | 163.28 | 157.26 |
| O11  | 85.12  | 121.19 | 163.64 |
| O12  | 165.86 | 121.87 | 140.64 |

Table 3. Predicted hyperfine parameters $V_{zz}$ and $\eta$ and isotropic shielding $\sigma_{\text{sample}}$ for NaM1, NaM2 and CoM1 models. Oxygen atoms are divided into three groups according to their unit positions.
HP of the constituent O atoms, and on the contrary, differences in the HP are signals of strong local deformations of the units. For example, in table 3 it can be seen that, for the NaM1 structure, the $V_{zz}$ of the $O_{basal}$ are mainly in the $9\times10^{21}$ \text{V m}^{-2} range, except for O1, O4 and O11. In figure 4(a) it can be seen in correspondence that O1 form the chemical bond with the Na $+$ $4\text{H}_2\text{O}$ unit, resulting the corresponding tetrahedron strongly deformed. In the upper part of the interlayer, O4 displaces from the basal plane towards the interlayer in a sort of $O$–$H$–$O$ bridge, thus also producing a strong deformation of the tetrahedron. Finally, O11 is shared between the basal plane of the corresponding tetrahedron and a free H. In correspondence, the $V_{zz}$ for the $O_{apical}$ are mainly in the $4.6\times10^{21}$ \text{V m}^{-2}, except for O16 and O18, which are the apical oxygen atoms of the deformed SiO$_4$ tetrahedra that share the basal O4. A similar reasoning can be done for the HP parameters at the Si atoms in these structures: the SiO$_4$ tetrahedrons of Si3 and Si5 are geometrically similar, as those of Si4 and Si6, then the corresponding $V_{zz}$ and $\eta$ values are near the same. In the lower plane of tetrahedra in figure 4(a) it can be seen that they are all geometrically different due to proximity of the (Na $+$ $4\text{H}_2\text{O}$) unit and the different bonding, consequently the HP of the corresponding Si atoms spread over a range of values. Interestingly, it should be noted that $V_{zz}$ and $\eta$ have much more sensitivity to the mentioned deformations than the $\sigma_{sample}$ for the case of Si, and the opposite occurs for the O. For the case of the Al and Mg atoms, it should be noted that $\sigma_{sample}$ has nearly constant values around 545 ppm for both type of atoms, whereas the corresponding $V_{zz}$ and $\eta$ have very different values between atoms in same the group.

In the case of the oxygen atoms in the $O_{basal}$ group of the NaM2, O1, O3 and O4 are the atoms that show out of range values. In figure 4(b) it can be seen that O1 and O3 are the bonding atoms between the (SiO$_4$) layer and the (Na $+$ $2\text{H}_2\text{O}$) units, thus producing an elongation in the tetrahedral faces. O4 stands for an $O$–$H$–$O$ bridge as in the previous case. In this case the similar deformed tetrahedra are those containing Si1 and Si7, and the pair Si2–Si8. Thus the corresponding apical oxygen O19–O13 and O20–O14 have very similar HP. Other pairs of similar $O_{apical}$ bonding can be observed from table 5.

In figure 5 it can be observed the relaxed cell of the CoM1. As a difference with the NaM1 and NaM2 cases, it can be observed that the elongation of the cell in the $c$-direction causes the interlayer (Co $+$ $6\text{H}_2\text{O}$) units to be displaced further away from the $O_{basal}$ atoms than in the previous cases, i.e., the enlargement is practically all contained in the interlayer spacing. As a result, in this structure do not appear Co–$O_{basal}$ chemical bonds similar to the Na–O1 in the NaM1 or the Na–O1 $+$ Na–O3 of the NaM2. Nevertheless, O4 and O3 are displaced towards the interlayer, thus producing strongly deformed tetrahedra. This fact is reflected in their corresponding HPs which are out of the mean values also for $V_{zz}$ and $\sigma_{sample}$ (table 3). Regarding the (Co $+$ $6\text{H}_2\text{O}$) units, it appears an unexpected behavior after relaxing the atomic positions. It can be observed that the corresponding units to Co1 and Co2 are different. Co1 maintains a nearly regular octahedron of 6 H$_2$O molecules, but Co2 is located in such a way that it is closer to one of the water molecules originally from Co1 (W1 in figure 5) than to one water molecule from its original unit (W2). So, this forms a strongly deformed octahedron, linked to the corresponding Co1 with a Co1–O–Co2 bridged structure, and releasing a water molecule.
5.3. Comparison with experimental results

From the \textit{ab-initio} results the chemical shift for each probe site of the MMT can be determined. On one hand, for the use of equation (3) the corresponding $\sigma_{\text{Ref}}$ values must be provided, nevertheless a search in the literature does not shield unique values. On the other hand, other authors use an empirical model in which there is a linear relation between the chemical shift $\delta_{\text{chem}}$ and the isotropic shielding $\sigma_{\text{sample}}$, in which the coefficients for the linearization are obtained from a linear regression from measurements on many different compounds:

$$
\delta_{\text{chem}}(\text{ppm}) = A \sigma_{\text{sample}} + B
$$

(4)

Also for this approach, after a selected search within this model at least two pair of values can be found for the linear coefficients A and B. In table 4 there are shown the data collected from the literature.

Unfortunately, up to our knowledge there are no reports for measurements using the $^{17}\text{O}$ isotope as probe atoms for this type of MMTs, nor NMR experiments reported on Co-MMT. Thus, for the comparison with the experiment we will restrict to the HP on Al, Mg, Si and Na for the NaM1 and NaM2 structures. In table 5 there are shown the HP obtained from the literature from NMR experiments on Na-MMT. From the predicted HP (table 3) we calculated the chemical shift for each one of these atoms, using equations (3) and (4). Thus we obtained a set of values that depend on the model and the experimental $\sigma_{\text{Ref}}$ and the A and B set of parameters. For $\sigma_{\text{sample}}$ for the different Al, Si, Na and Mg atoms we used those obtained from the corresponding average of the predicted values from table 3. For the determination of $V_{ZQ}$ form NMR data we used:

$$
C_Q = \frac{V_{ZQ} Q e}{\hbar}
$$

(5)

$$
P_Q = C_Q \sqrt{1 + \frac{\eta^2}{3}}
$$

(6)
by means of the corresponding quadrupolar moment Q after averaging the obtained moduli of the $V_{zz}$ for the same type atoms.

As it can be seen, the calculated $^{29}$Si chemical shift is in good agreement with the reported chemical shift in the literature. Regarding $^{23}$Na, the comparison of the calculated $\delta_{\text{chem}}$ with the experimental one becomes difficult due to the strong dependence on the A and B empirical parameters selection. Specially, the B parameter presents strong differences between the data form [64] and [37], and this difference particularly affects the results for chemical shifts near zero, as in this case. On the other hand, the $V_{zz}$ obtained values are not far from the experimental data. For Mg, the calculated $\delta_{\text{chem}}$ using [37] resulted higher than that observed in [26]. Also, the $V_{zz}$ are higher for both models NaM1 and NaM2. On the contrary, the obtained $\eta$ are in good agreement with the experiment. For $^{27}$Al it can be seen that $V_{zz}$ are in good agreement with the range extracted from the work of Takahashi et al (the range comes from the indetermination $\eta$, taking the extreme values 0 and 1) for both NaM1 and NaM2. Nevertheless it must be observed that the comparison is performed with the mean value of the $V_{zz}$ module for all the Al sites. While for NaM2 all the Al sites have uniform $V_{zz}$ values, the corresponding values in NaM1 present strong fluctuations. These fluctuations are not reflected in the individual $\sigma_{\text{sample}}$ Nonetheless the obtained chemical shifts resulted above the experimental result using equation (3) or equation (4).

6. Conclusions

Cobalt sorption on natural Na-MMT was experimentally studied. From the results of the XRD spectra and analysis of the solid and residual liquid phase it was concluded that Co$^{2+}$ was introduced in the sample and as a result the layered structure was strongly enlarged in the direction perpendicular to the interlayer.

DFT calculations can be an important tool to contribute to resolve issues relating to the structure of natural and synthetic layered materials. This theory was successfully used to reproduce the changes introduced in MMT structure upon substitution of Na by Co as it was observed. Regarding the total energy values, ab initio calculation predicts that the replacement, though sorption, of Na by Co in Na-MMT is energetically favorable. Three different models with different hydration/Na-concertation state were tested to reproduce the Na-MMT samples and one for the obtained Co-MMT. Within these models of Na-MMT, two of them with 4 H$_2$O and one or two Na atoms give d$_{001}$ parameters compatible with the natural sample, whereas the third model with 2 Na atoms and 8 H$_2$O have been discarded due to the high d$_{001}$ values obtained at equilibrium. The Co-MMT model with 2 Co atoms and 6 H$_2$O resulted in a d$_{001}$ compatible with that obtained experimentally in this work.

In this work we observed that the substitution of the Na-n(H$_2$O) by Co-m(H$_2$O) produces not only an enlargement of the cell, but also a re-organization of some of the O atoms of the SiO$_4$ tetrahedrons that affect the geometry of the sheets, thus changing their properties.

The hyperfine parameters on all models were calculated giving a good approach to the experimentally determined by NMR. The equilibrium configuration together with the tetrahedra and octahedral geometry of the (SiO$_4$) and (Al/MgO$_6$) units were discussed and related with the corresponding hyperfine parameters.

### Table 5. Experimental and calculated chemical shift, $V_{zz}$, and $\eta$.

| Probe | $Q$ [meV] | $\delta_{\text{chem}}$ | $V_{zz}$ | $\eta$ | $\delta_{\text{chem}}$ | $V_{zz}$ | $\eta$ | $\delta_{\text{chem}}$ | $V_{zz}$ | $\eta$ |
|-------|-----------|------------------------|----------|--------|------------------------|----------|--------|------------------------|----------|--------|
| $^{27}$Al | 140.2$^a$ | 6.9$^e$ | 2.7–3.12 | 11.98$^b$, 14.05$^c$ | 2.43 | 0.47 | 14.72$^b$, 16.85$^c$ | 2.52 | 0.48 |
| $^{29}$Si | $-93.7$ natural MMT$^f$ | $-88.6$ synthetic MMT$^f$ | | | | | | | | |
| $^{23}$Na | 109$^b$ | $-6^d$ | 1.44$^d$ | $-3.25^c$ | 0.93 | $-2.97^c$ | 1.20 | |
| $^{25}$Mg | 199.4$^b$ | 16$^d$ | 0.7 | 29.3$^h$ | 1.52 | 0.76 | 26.09$^h$ | 1.12 | 0.61 |

$^a$ using coefficients from [64].
$^b$ using coefficients from [17].
$^c$ using coefficients from [63].
$^d$ data from [33].
$^e$ data from [39].
$^f$ data from [26].
$^g$ data from [65].
$^h$ data from [66].
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