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Second Harmonic Scattering can Probe Hydration and Specific Ion Effects in Clay Particles

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ABSTRACT: We report Montmorillonite (MMT) particles organization in aqueous solution using polarization resolved second harmonic scattering (SHS). This technique can probe specifically the organization of water in this system. The effect of the colloidal stability of the particle and the nature of the interlayer cations on the SHS Intensity is highlighted. The interlayer hydration has been probed and changes with the order Mg²⁺ > Ca²⁺ > Li⁺, Na⁺, K⁺ > Cs⁺. Real time SHS evolutions during cesium exchange have been monitored, and different pathways are discussed in regards with the experimental results.

TEXT
1. Introduction

Clay minerals and in particular Montmorillonite (MMT) particles are nanostructured materials composed by stacks of negatively charged layers compensated by solvated cations. These materials are currently involved in many environmental and industrial processes because of their catalytic and retention properties. Understanding the fundamentals of clay-water interfaces is a crucial point for the success of those applications. Numerous computational, experimental works using XRD, quasi-elastic neutron scattering or infrared spectroscopy studied the swelling, water adsorption and cation exchange properties on MMT under controlled humidity conditions. Those studies contributed to our current-day understanding of the MMT hydration, thickness and interactions between cations. Roughly, MMT crystalline hydrates with one, two, or three molecular layers of water separating the stack layers depending on the controlled relative humidity and the nature of interlayer cations. In aqueous solution, the behavior of MMT colloidal system is quite more complicated because complex chemo-mechanical properties involving competitive effects, determine the whole organization of the mineral plates into an exfoliated or stacked arrangement. The precise interplay between interfacial water structure and ion adsorption/exchange remains experimentally poorly characterized using in situ and in real time probes that are non-invasive, label-free and specific to interfacial water.

Nonlinear optics and Second Harmonic Scattering (SHS) have been previously used to studied MMT particles in suspension. This first study demonstrated the contribution of the oriented water around the charged plate in the whole SHS intensity and its dependence with the ionic strength. The colloidal transition between the delaminated/stacked plate and the change in the interlayer hydration during cation exchange was not explored in this former work and will be investigated in this article. In particular, the results presented here will be discussed in term of coherent contribution in the SHS Intensity coming from water correlation induced by the MMT
particles. Indeed, in the last three years, numerous works have significantly improved our understanding of the SHS signals concerning the contribution of organized or correlated water in the whole SHS intensity\textsuperscript{13-16}, particularly in the domain of electrolyte solution\textsuperscript{17-20}. Based on this general framework, the global SHS intensity $I$, for $N$ molecules, each located at position $r_i$, can be written as\textsuperscript{18,21}

$$I(v, u) \propto \left\langle \left( e_y \times \sum_{i=1}^{N} \beta_{labo}(\Omega_i) : uu \right) \times e_y \right\rangle^2 + \left\langle \left( e_y \times \sum_{i=1}^{N} \sum_{\omega} \beta_{labo}(\Omega_i) : uu \right) \left( \beta_{labo}(\Omega_j) : uu \right) e^{i\frac{q}{\hbar} (r_i - r_j)} \times e_y \right\rangle^2 \quad (1)$$

Where $\beta_{labo}$ is the hyperpolarizability tensor expressed in the laboratory frame, $q$ is the scattering wavevector $q = k_s - 2k_i$ where $k_i$ is the fundamental incident wave vector, $k_s$ is the harmonic scattered one, $u$ and $v$ are the polarization directions of the incoming and outgoing beams described by the angles $\gamma$, $\Gamma$ (see figure 1 for the notation), and $\Omega$ is the orientation of the molecule, define with three Euler angles, in the laboratory frame (XYZ). The brackets $\langle \rangle$ stand for the time average. The first part of equation 1 is referred as the incoherent contribution and the second term is referred as the coherent contribution in the SHS intensity. In the case of correlated scattered molecule, this second term can overcome the first term, as it will be the case in systems presented here. Otherwise, for the case of uncorrelated molecule, this second term is zero.

2. Experimental Methods

2.1. *Experimental Optics setup*:

The figure 1 depicts the notation together with the experimental setup\textsuperscript{22}. Briefly, this setup was built on a femtosecond Ti-sapphire oscillator laser source providing pulses with a duration of about 100 fs at a repetition rate of 80 MHz (coherent, model Chameleon ultra II). After passing through a low-pass filter, the fundamental beam, set to a fixed wavelength at 800 nm and an
averaged power of 600 mW, was focused by a microscope objective (Ealing x10, numerical aperture 0.25.) onto the sample at 2 mm close to the output cell window (Hellma QS 10 mm). The SH light was collected at 90° by a 10 cm focal length lens and separated from its fundamental counterpart by a high-pass filter. The SH light was detected with a water-cooled CCD camera (Andor, Newton) placed after a spectrometer (Andor, Shamrock 193). The fundamental input beam was linearly polarized and the input polarization angle $\gamma$ was selected with a rotating half-wave plate. An analyzer, placed in front of the spectrometer, was used to separate two polarization states: vertical output state further named $I(\gamma, V_{out})$, or horizontal output state referred to $I(\gamma, H_{out})$. In order to facilitate the reading and to be coherent with other works$^{13, 15, 18-19}$, the SHS intensity $I(0^\circ, V_{out})$ is also referred to $I_{SSS}$ and $I(0^\circ, H_{out})$ to $I_{PSS}$. figure 1A summarizes these notations. The experimental data of figure 1 were recorded during 10 seconds for each point. The experimental data of figure 2 were recorded during 10 seconds, each point are repeated 5 times, and the errors bar represents +/- 2 standard deviation. In figure 3, each point is recorded during 10 s in the $I_{SSS}$ configuration and the global time monitoring is about 4500 s. All experiment were conducted at 20°C and under stirring condition (a magnetic stirrer operates in the cell measurement). For figures 1 and 2, the solution were equilibrated 12h prior the measurement.

2.2. Materials

In all experiments, the MMT particles are prepared in 18 MΩ.cm water (MilliQ) at the same concentration of 0.25 g/L and two types of MMT particles were used. The particles named in the manuscript as Na-MMT are the SWy-3 montmorillonite from the Source Clay Repository of The Clay Mineral Society with structural formula $[(Al_{2.99} Fe_{0.43} Mg_{0.52})(Si_{7.97} Al_{0.03})O_{20}(OH)_4] M^{+0.7}$. This clay mineral exhibits a low octahedral charge and extremely limited tetrahedral
substitution. Size fractionation was performed by centrifugation to extract the <1 µm size fraction. Prior to the measurement, the particles were sonicated during 10 min and kept under permanent agitation. The particles named Ca-MMT in the manuscript were prepared by ion-exchange reaction of Na-MMT at room temperature with 1 mol/L aqueous solutions of CaCl₂. The suspensions solution was first allowed to stir for 24 h and then, the solid fraction was removed from the saline solution by centrifugation and dispersed again in a fresh saline solution at the same concentration. These steps were repeated three times. The excess of CaCl₂ was removed by rinsing several times the solid using distilled water until no Cl anions were measured in solution as indicated by a negative silver nitrate test. For the figure 2 data, the following ions are added in the solution: LiNO₃, KNO₃, CsNO₃, NaNO₃, Ca(NO₃)₂, Mg(NO₃)₂, from Sigma Aldrich. For the figure 3, ions addition is carried out using a microvolume injection (18 µL) of a 0.8 M solution of NaNO₃ in a 2 ml suspension of 0.25 g/L of Na-MMT (figure 3A) or 0.8 M solution of CsNO₃ in a 2 ml suspension of 0.25 g/L of Ca-MMT (figure 3B), in such way, the dilution factor is negligible. The solution is rapidly homogeneous thanks to a magnetic stirrer place in the measuring cell.

3. Results and discussion
Figure 1. Sketch of the SHS experiment, SHS intensities are recorded at a scattering angle of 90° for an input polarization described by the angle $\gamma$, and output polarization states described by $\Gamma=V$ (Vertical) or $H$ (Horizontal). The four circles represent the different systems probed: A) Na-MMT particles at 0 M ionic strength, B) Na-MMT particles at $10^{-3}$ M ionic strength, C) pure water and D) Ca-MMT particles at 0 M ionic strength. In the four graphs on the right, the full circles represent the experimental $I(\gamma,V_{\text{out}})$ and the empty squares the experimental $I(\gamma,H_{\text{out}})$. The full and dashed lines are fits using equation (2).

Figure 1 presents the polarization resolved SHS intensity for different systems: pure water, initially exfoliated or delaminated Na-MMT particles at various Ionic Strength (IS) and initially stacked Ca-MMT particles. The experimental data in figure 1 have been normalized by the SHS intensity of pure water, it means that all points have been divided by the $I_{\text{SSS}}$ water intensity. The derivation of equation 1 gives the following expression for the polarization resolved SHS intensity:

$$I_{\text{SHS}}(\gamma) \mu \left( a_{\text{inc}} + a_{\text{coh}} \right) \cos^4(\gamma) + \left( b_{\text{inc}} + b_{\text{coh}} \right) \cos^2(\gamma) \sin^2(\gamma) + \left( c_{\text{inc}} + c_{\text{coh}} \right) \sin^4(\gamma) \ (2)$$
where $\Gamma=V$ or $H$. The subscript ‘inc’ and ‘coh’ means -respectively- incoherent and coherent contribution and the different values obtained by the fits presented on figure 1 are given in SI.

When MMT particles are introduced in water, the SHS intensity is globally enhanced by a factor around two, which is explained by coherent contribution of water coming from its organization around and inside the mineral sheet. The coherent term of equation 1 is thus not zero and contributes in the same order of magnitude as the incoherent term coming from “bulk” water. No contribution in the SHS intensity of the mineral materials is considered here, as it will be justified below with the cesium results. When the ionic strength is modified, the behavior of the polarization plot changes with a small decrease in the $I(\gamma,H_{\text{out}})$ intensity whereas the $I(\gamma,V_{\text{out}})$ remains more or less stable. For the case of stacked Ca-MMT particles, the $I(\gamma,V_{\text{out}})$ intensity is higher compared to exfoliated Na-MMT system. To study the transition between exfoliated/stacked particles and the possible specific ion effects, figure 2 shows the SHS intensity evolution for initially exfoliated Na-MMT particles by varying ionic strength with different cations.
Figure 2. Evolution of the normalized SHS intensity in the PSS and SSS polarization states as a function of the ionic strength for different mono and divalent cations. On the top, illustration of the different organization probed in the experiment. On the right, a microscopic view showing the different hydration states in the interlayer domains.

To interpret those curves, two domains of ionic strength (IS) can be considered, low IS (IS<10⁻³ M) and high IS (IS>10⁻³ M). In the first domain, no specific ion effects are observed since all curves seem to follow globally the same evolution: the intensity I_{SSS} is nearly constant and intensity I_{PSS} decreases. These evolutions are explained by the interference between water molecules oriented by the electric field induced by the surface charge onto the platelet. This contribution is referred as Diffuse Double Layer (DDL) contribution¹⁵. This evolution in the 10⁻⁵
- $10^{-3}$M IS range is in agreement with both theoretical model\textsuperscript{15} describing the interference in the DDL and with previous experimental work presented by Eisenthal\textsuperscript{12}. At higher IS ($10^{-3}$-$10^{-2}$ M), the DDL contribution tends toward zero and the colloidal stability of the MMT particles changes with a transition to a stacked organization\textsuperscript{24-25}. This transition is observed both on the two polarization states $I_{\text{PSS}}$ and $I_{\text{SSS}}$. Depending on the cation added in the system, the experimental data exhibit different behaviors coming from the supposed cation exchange and leading to different water organization in the system.

When Li\textsuperscript{+} or K\textsuperscript{+} ions are added, no significant deviations are observed experimentally in the SHS evolution compared to the case of sodium ion. When Ca\textsuperscript{2+} or Mg\textsuperscript{2+} ions are introduced, the $I_{\text{PSS}}$ and $I_{\text{SSS}}$ increase significantly. The cation exchange induced a stacking of the layer with more correlated water in the interlayer domain. The data obtained at $10^{-2}$ ionic strength may be compared with the case of Ca-MMT presented in fig 1D. The global polarization plots are consistent even if prepared Ca-MMT intensity is a little bit higher, 2.5 compared to 2.2 with the data coming from the in situ Ca-exchange. In the case of prepared Ca-MMT, the stacking is likely better than the organization obtained by in situ cation exchange. When Cesium ion are added, the SHS intensity decreases and the ratio $I/I_{\text{water}}$ falls very close to 1 for $I_{\text{SSS}}$ and close to 1.4 for $I_{\text{PSS}}$. This probably means that the coherent contribution in equation 1 becomes very small compared to the initial condition. It is interesting to note that the ratio $I_{\text{PSS}}$ is slightly higher but nevertheless close to the value measured by Roke and coworker in electrolyte solution at high ionic strength\textsuperscript{19}. These two ratios may be interpreted as the electrolyte contribution and the contribution of organized water around or inside Cs-MMT particles seems negligible. These results likely show that few correlated water molecules are involved in the interlayer domain and around the particles. This is in agreement with a zero water model in the interlayer and with the “cavity charge matching effect” which has been recently described for the cesium ions.\textsuperscript{26}
Moreover, the cesium results are also consistent with a negligible contribution of the mineral material compared to water contribution. Indeed if the material itself exhibits an intrinsic significant nonlinear response, this contribution would be measured in that case.

To summarize this part, increasing ionic strength in the system induces a decrease in the DDL SHS contribution, which is a non-specific ion effect, and induces also a transition from exfoliated to stacked particles. When particles are stacked, the water organization in the interlayer domains exhibits specific ion effect, with an hydration effect varying in the order \( \text{Cs}^+ < \text{K}^+, \text{Na}^+ \approx \text{Li}^+ < \text{Ca}^{2+} < \text{Mg}^{2+} \). To get in situ information about the transition state during the cation exchange and the global organization of the system, the SHS real time intensity is recorded during the ion exchange.
Figure 3. Real time evolution of $I_{SSS}$ for two cases: Cs injection on exfoliated Na-MMT at 0 ionic strength (A), and Cs injection on stacked Ca-MMT at 0 ionic strength (B). In all curve, the initial time is the injection time. Top, a microscopic view showing the different steps during the Cs exchanges process on the exfoliated Na-MMT. Bottom, a microscopic view showing the different steps during the Cs exchanges process on stacked Ca-MMT.
Figure 3 shows the real-time evolutions when Cs is injected in the system at zero ionic strength for the two different kinds of MMT particles: exfoliated Na-MMT or stacked Ca-MMT. In both cases, the final ionic strength is $7 \times 10^{-3}$ M, and SHS signal evolutions with time are well reproduced. For the case A, the time dependent plot exhibits a first increase during around 1000 seconds. This duration could be considered as the characteristic time of exfoliated/stacked transition. It contains both the ions diffusion/adsorption signal and the stacking platelet characteristic time. This characteristic time is consistent with the SHS evolution obtained for the case of calcium injection onto exfoliated Na-MMT (see additional in the SI). This first increase is followed by an abrupt decrease, which is interpreted as a dehydration step where water is expelled from the interlayer domains. This time duration is likely in the same order of magnitude observed for the previous diffusion/exchange/stacking characteristic times. The equilibrium is reached in around 3000 seconds. In the case B, the evolution shows a more fluctuating decrease and reached an equilibrium state of the same order of magnitude than in case A. These fluctuations are the highest during the time interval 1000-3000 seconds and are comprised in the range between 2.5 and 1.5 value. We interpreted these fluctuations as a signature of the cation exchange: the cesium cations close progressively the sheet when calcium cations are expelled. Those results are remarkable in the context of recent works about cesium adsorption in clays concerning the discussion between different pathway referred as “direct exchange” or “decollapse–driven exchange”. In the case of exfoliated Na-MMT (figure 3A), the Cs-exchange seems to follow the sequence « collapse induced dehydration ». In the case of initially stacked Ca-MMT the nature of the fluctuation in the experimental data seems to sign the coexistence between calcium hydrated / cesium dehydrated stacked domain in the system. The sequence “decollapse driven exchange” as illustrated on bottom figure 3, may be in agreement with this fluctuating evolution. Nevertheless, these data cannot clearly determine if one pathway between
“direct exchange” or “decollapse–driven exchange » is really preferred. These real time SHS data show that for the two different initial states studied here, the characteristic time of ion exchange is nearly the same. In other words, the “opening” of the stacked arrangement (see bottom figure 3) seems not to be a limiting step in the kinetics.

4. Conclusions

In conclusion, this work demonstrates that SHS is able to probe MMT particles organization and interlayer hydration. The nature of the compensating cation strongly changes this interlayer hydration. Real time SHS has been used to monitor the cation exchange for two different initial states. For initially exfoliated particles, a specific evolution with two steps involving a collapse before dehydration has been highlighted. For initially stacked particles, fluctuations in the SHS intensity are observed during the cation exchange. Future works involving other type of clays particles, for example clays nanotube, are currently planned in order to study the effect of other geometrical water arrangements in such material. Overall, we hope that these studies will stimulate the use of SHS to probe hydration and molecular organization around and inside porous materials.

ASSOCIATED CONTENT

Supporting Information. Additional information about the fit parameters presented in figure 1 and complementary real time SHS evolution. The following files are available free of charge.

AUTHOR INFORMATION

Notes
Any additional relevant notes should be placed here.

The authors declare no competing financial interests.

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TOC GRAPHICS