Water Mobility within Compacted Clay Samples: Multi-Scale Analysis Exploiting $^{1}$H NMR Pulsed Gradient Spin Echo and Magnetic Resonance Imaging of Water Density Profiles

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ABSTRACT: $^{1}$H NMR pulsed gradient spin echo attenuation and water density profile analysis by magnetic resonance imaging are both used to determine the mobility of water molecules confined within a porous network of compacted kaolinite clay sample (total porosity of ~50%). These two complementary experimental procedures efficiently probe molecular diffusion within time scales varying between milliseconds and few hours, filling the gap between the time scale of diffusion dynamics measured by traditional quasi elastic neutron scattering and through-diffusion methods. Furthermore, magnetic resonance imaging is a nondestructive investigation tool that is able to assess the effect of the local structure on the macroscopic mobility of the diffusing probe.

1. INTRODUCTION

Clay minerals are ubiquitous at the surface of the earth and play important roles in a large variety of environmental applications, including pollutant retention and nuclear waste storing, by exploiting their physicochemical properties (low hydraulic conductivity, large anisotropy, high specific surface and adsorbing power, ionic exchange capacity). In this context, numerous studies have been performed to determine the mobility of water molecules and ions confined within the complex porous network of dense clay samples. For this purpose, quasi elastic neutron scattering (QENS) and through-diffusion measurements are the two complementary approaches generally used to quantify the mobility of water molecules confined within the clay porous network. While QENS is an ideal probe of local water motions performed during a limited diffusion period (less than 1 ns), through-diffusion experiment probes macroscopic displacements of confined fluids at a time scale larger than a few minutes. Because of the multi-scale organization of clay aggregates within the dense samples, it appears useful to investigate fluid mobility at intermediate time scales. Such investigations are necessary to efficiently relate the apparent macroscopic mobility of the confined fluid to the organization of the clay sample. For this purpose, we investigate the potentialities of the two applications of NMR, i.e., measurement of the water self-diffusion tensor by using pulsed gradient spin echo (PGSE) attenuation and analysis of time evolution of water density profiles by exploiting magnetic resonance imaging (MRI). This approach contributes significantly to characterize the multi-scale mobility of confined water molecules because PGSE-NMR adequately probes diffusion times larger than 1 ms, whereas diffusion times larger than a few seconds are easily investigated by MRI, thus filling the above-mentioned gap between the time scales investigated by QENS and through-diffusion experiments. Previous studies have demonstrated the applicability of PGSE-NMR and MRI to investigate the mobility of water molecules within aqueous dispersions and dense pastes of synthetic minerals. The application of the same NMR experiments for dense clay samples is never obvious because of the presence of iron within natural clay particles, strongly enhancing the NMR relaxation of the nuclear magnetization pertaining to the protons of the confined water molecules. To settle this question, we selected natural clay (kaolinite) whose composition (0.039 iron per unit cell) and size distribution were already characterized. Furthermore, by contrast with other natural clays (like montmorillonite), kaolinite (like illite) is a nonswelling clay mineral, thus avoiding the occurrence of some interlayer porosity after saturation of the clay sample by water molecules. As a consequence, the porous network accessible to water molecules is limited to interparticle porosity.

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In this study, $^1$H PGSE-NMR attenuation measurement is first used to determine the components of the tensor describing the water self-diffusion within the water-saturated sample of kaolinite. MRI is then used to determine the water concentration profiles after the addition of a controlled amount of heavy water on the top of the water-saturated clay sample. The exchange between the added heavy water and initially confined water molecules is quantified by integrating the water concentration profiles whose asymptotic limit may be used to determine the porosity of the clay sample. Numerical modeling performed by Brownian dynamics (BD) is finally used to interpret the observed exchange kinetics on the basis of the clay sample porosity and the longitudinal component of the water self-diffusion tensor previously measured by PGSE-NMR, thus demonstrating the agreement between both experimental procedures.

II. MATERIALS AND METHODS

II.1. Sample Preparation. The kaolinite clay sample (i.e., KGa-2 kaolinite reference from the Source Clay Repository of The Clay Minerals Society) was used to prepare a cylindrical compacted clay sample (diameter 6.5 mm, height 12.3 mm) for NMR measurements. For this purpose, the original kaolinite sample was Na-saturated using three saturation cycles in a 1 mol/L NaCl solution, washed in distilled water, and dried at room conditions. The obtained powder was then placed within a poly(tetrafluoroethylene) (Teflon) cylinder. The size of the NMR detection coil and underwent a uniaxial compression to reach a porosity value close to $\sim$0.5. Information on the chemical composition and properties of the kaolinite sample are available in the literature.\textsuperscript{2} Transmission electron microscopy (TEM) images (see Figure 1a,b) are recorded after the dispersion of the sample into ethanol to illustrate the size distribution of the stacks composed of individual clay platelets. The porosity of the dry clay sample ($\phi = 0.52$) is determined by helium pycnometry, corresponding to an average grain density of 2.75 g/cm$^3$. The PGSE-NMR experiments are first performed with the water-saturated clay sample. The kinetics of the water/heavy water exchange is determined by $^1$H MRI profile analysis after the addition of 0.23 mL of D$_2$O on the top of the hydrated clay sample. The compactness of the clay sample was obtained by uniaxial compression to reach the same porosity ($\phi \sim 0.5$). The clay sediment is carefully saturated by water and equilibrated during 2–3 days before performing PGSE measurements. However, no boundary condition is applied to the sediment after compression. As a consequence, the porosity of the clay sample must be determined independently by helium pycnometry for the dry sample and after hydration by MRI. Some difference is expected to occur because no boundary is applied to restrict the swelling of the clay sample.

II.2. $^1$H NMR Measurements. NMR measurements were performed using a Bruker DSX100 Bruker spectrometer with a static field of 2.35 T and equipped with a saddle detection coil. A typical pulse duration of 5 $\mu$s is required for transferring the equilibrium longitudinal magnetization into the detection plane perpendicular to the static magnetic field $B_0$ (also called $\pi/2$ pulse). Figure 2 illustrates the pulse sequence used to perform the PGSE-NMR attenuation measurements.\textsuperscript{2,3} The encoding of transverse magnetization is performed by a pulse sequence composed of an initial $\pi/2$ pulse used to create transverse magnetization, a set of bipolar field gradients (duration $\delta$, strength $G$) applied along any desired direction, and a final $\pi/2$ pulse to transfer back magnetization in the longitudinal direction. After the evolution period ($\Delta$), transverse magnetization is refocused by using a pulse sequence similar to the encoding one, but with opposite field gradients. NMR relaxation times are critical parameters limiting the feasibility of this PGSE-NMR experiment. Because of the paramagnetic impurities (iron) present in the atomic network of kaolinite, the NMR relaxation time of the longitudinal magnetization of the confined water molecules ($T_1 = 0.053$ s) is fifty times smaller than that of bulk water ($T_1 = T_2 \approx 3$ s). Furthermore, the transverse relaxation time of the same confined water molecules ($T_2 = 1.2 \times 10^{-3}$ s) is again reduced by more than 1 order of magnitude because confinement is responsible for large differences between the NMR relaxation rates of liquids ($T_2 < T_1$). The purpose of the stimulated echo used in our PGSE-NMR experiment is to transfer the transverse magnetization encoded by the first set of pulsed gradient into the longitudinal direction parallel to the stimulated echo.
the static magnetic field $B_0$. As illustrated in Figure 2, the stimulated echo allows them to probe the mobility of the confined water molecules during roughly 150 ms since the longitudinal magnetization becomes negligible after the diffusion time $\Delta_{\text{max}} \sim 3T_1$. Another limitation arises from the time required to build the field gradients (typically 60 $\mu$s).

As a consequence, the duration (4$r = 3.04$ ms) of the coding and refocusing pulse sequences (see Figure 2) reaches milliseconds (i.e., the order of magnitude of the transverse relaxation rate $T_2$), leading to important attenuation of the NMR magnetization according to the exponential law $\exp(-4r/T_2) = 0.08$. Finally, the intensity of the NMR signal varies according to the relationship$^{32,33}$

$$E(\vec{G}, \Delta) = C_{\text{te}} \exp \left[ -4\pi^2 \frac{q^2}{\gamma_G} \vec{g} \cdot \vec{c}_G (\Delta + \frac{3}{2} \tau - \frac{\Delta}{\delta} - \frac{4r}{T_2} - \Delta \frac{T_1}{T_2} \right]$$

(1)

where, $q = \gamma \delta G/\pi$, $\gamma$ is the gyromagnetic ratio ($2.6752 \times 10^8$ rad/s for $^1$H), $G$ is the strength of the field gradient (see Figure 2) and $\vec{c}_G$, its director, $D$ is the self-diffusion tensor, and $\delta$, $\Delta$, and $\tau$ are time delays illustrated in Figure 2. Since our PGSE-NMR echo attenuations are recorded for a unique set of time delays ($\delta = 500 \mu$s, $\Delta = 20$ ms, and $\tau = 760 \mu$s), the components of the self-diffusion tensor are obtained by fitting the attenuation of the NMR signal measured for a set of strengths of the field gradient $G$ according to a Gaussian propagator$^{33}$

$$E(\vec{G}, \Delta) = E(0, \Delta) \exp \left[ -4\pi^2 \frac{q^2}{\gamma_G} \vec{c}_G \cdot \vec{G} \left( \Delta + \frac{3}{2} \tau - \frac{\Delta}{\delta} \right) \right]$$

(2)

This Gaussian propagator of self-diffusion$^{33}$ is equivalent to the intermediate scattering function$^{36}$ probed by neutron scattering experiments (QENS). The maximum resolution of spatial domains probed by these PGSE-NMR measurements is given by the maximum strength of the magnetic field gradient$^{33}$ ($G_{\text{max}} = 1.6$ T/m)

$$\text{resolution}_{\text{max}} = \frac{1}{q_{\text{max}}} = \frac{\pi}{\gamma \delta G_{\text{max}}} \approx 15 \mu m$$

(3)

where, $q$ is equivalent to the wave vectors probed by QENS.

$^1$H magnetic resonance imaging is also used to study the exchange between the initially confined water molecules and excess bulk heavy water added to the saturated clay samples. Figure 3 illustrates the pulse sequence used for these MRI experiments.$^{33}$ After an initial $\pi/2$ excitation pulse, a magnetic field gradient (noted $G_z$) is applied during the time period $\delta/2$ along the vertical direction ($O_z$) corresponding to the cylindrical axis of the clay sample. Inversion pulse (noted $\pi$) is then applied and the same field gradient $G_z$ is applied during the evolution time $\delta = 2.56$ ms with a simultaneous acquisition of the NMR signal. This sequence is selected to generate an echo of the NMR signal after the evolution period $\delta/2$, leading again to a noticeable attenuation (exp($-\delta/(2T_2)$) $\approx 0.34$) of the NMR signal pertaining to the confined water molecules without significantly altering the NMR intensity of free water (see $T_2$ values reported above). Our MRI measurements were performed for an evolution period $\delta$ composed of 256 elementary time steps of 10 $\mu$s. As the time domain is used to sample the NMR signal, the corresponding frequency domain obtained by Fourier transform is also composed of 256 elementary frequency steps. As a consequence, the resulting longitudinal MRI profiles are digitalized into 256 superposed sheets. To easily investigate the water molecules located in the hydrated clay sample (length $L_1 = 12.3$ mm) and the amount of added heavy water (length $L_2 = 7$ mm), we want to probe a total sample length of 25 mm, leading to a spatial resolution of 97.7 $\mu$m (since resolution $= 25/256$ mm). This spatial resolution corresponds to a wave vector $q$ of $1.024 \times 10^4$ m$^{-1}$ obtained by applying a magnetic field gradient of 0.094 T/m (see eq 3).

II.3. Numerical Simulations. Numerical simulations of Brownian dynamics$^{41}$ (BD) are used to analyze the macroscopic exchange between the water molecules initially confined within the clay sample and the added heavy water as investigated by profile analysis obtained by $^1$H MRI. To our knowledge, determination of water diffusivity by direct analysis of the time evolution of the water concentration profiles is not possible. For this reason, we proceeded to numerical simulations of BD to perform a quantitative comparison between the experimental and numerical data, thus extending to hours the time-scale initially probed by PGSE-NMR spectroscopy (typically a few milliseconds). For this purpose, $N_1$ $H_2O$ molecular probes ($N_1 = 386 571$) are uniformly distributed within an ideal parallelepiped (longitudinal length $L_1 = 12.3$ mm, section $S_1 = 13.75$ mm$^2$) in contact through its upper surface with another parallelepiped (longitudinal length $L_2 = 7$ mm, section $S_1 = 25$ mm$^2$) containing $N_2$ $D_2O$ molecular probes ($N_2 = 400 000$). These dimensions and water contents have been selected to reproduce the volumetric ratios between the diffusing spaces inside the clay sample (labeled 1) and the added bulk water (labeled 2). The self-diffusion coefficients of both water and heavy water molecules are set equal to the value determined by $^1$H PGSE-NMR attenuation measurements, whereas their values in bulk water are set equal to $2.1 \times 10^{-9}$ $m^2/s$, in agreement with the available experimental data. Simple reflecting conditions are applied when a molecular probe hits any wall of the simulation cell except for the surface connecting the two independent sub-volumes: when a probe, initially diffusing in the cell corresponding to the clay sample (labeled 1), hits the separating surface, its transfer to the cell labeled 2 is always considered. By contrast, a molecular probe initially diffusing in the cell corresponding to bulk water (labeled 2) and hitting the separating surface will be considered to potentially transfer to the clay sample with a probability equal to the porosity of the clay sample. Adequately formulating the condition for accepting these potential transfers between the two sub-cells is the only difficulty for modeling such water exchange between

![Figure 3. Schematic view of the pulse sequence used to perform magnetic resonance imaging (MRI) of the water concentration profiles inside the compacted kaolinite clay sample (see text Section II.2).](image-url)
the two sub-cells with different sizes, water concentrations, and mobilities. An interesting suggestion was the use of detailed balance conditions. Unfortunately, such a procedure is valid only for describing stationary states. By contrast, our problem is typically a transient dynamical process for which detailed balance condition becomes inadequate. For this purpose, we selected to use a systematic control, at each step of the BD iteration, of the water flux across the separating interface between the two sub-cells. We simply impose that the total number of \( H_2O \) and \( D_2O \) molecules leaving a given sub-cell must be perfectly compensated by the same number of \( H_2O \) and \( D_2O \) molecules leaving the other sub-cell. Such condition corresponds to some extension, at the microscopic scale, of the macroscopic incompressibility of liquid water. An equivalent procedure was already used to describe by Brownian dynamics the ionic exchange between ions confined within clay interlayers and freely diffusing in aqueous solution.

The displacement of the molecular probes (labeled i) is described by the Langevin equation

\[
\frac{d\vec{\xi}(t)}{dt} = -m_i \vec{\xi}(t) + \vec{R}_i(t)
\]

where, \( m_i \) and \( \vec{\xi}_i \) are respectively the molecular mass and friction coefficient, and \( \vec{R}_i(t) \) is a random force resulting from thermal collisions within the liquid. This random force must satisfy statistical requirements: it must be Markovian and time independent (see eq 5a), with zero mean (eq 5b), without correlation with the velocities (eq 5c), and distributed along a Gaussian law (eq 5d)

\[
\langle \vec{R}_i(0) \times \vec{R}_j(0) \rangle = 0
\]

\[
\langle \vec{\xi}_i(t) \times \vec{\xi}_j(t) \rangle = 0
\]

\[
P(\vec{R}_i) = \frac{1}{2\pi \sqrt{\langle \vec{R}_i^2 \rangle}} \exp \left( -\frac{\vec{R}_i^2}{2\langle \vec{R}_i^2 \rangle} \right)
\]

where, \( P(\vec{R}_i) \) is the distribution law of the modulus of the random displacements, \( \vec{R}_i \) is the Boltzmann constant, and \( T \) is the temperature. Equation 5a results from the fluctuation–dissipation theorem ensuring the balance between the increase of the velocities induced by the random force and its decrease resulting from the intermolecular friction. Various algorithms have been proposed to solve the set of eqs 4 and 5a–5d, depending on the ratio between the friction coefficient (\( \vec{\xi}_i \)) and the time step (\( \Delta t \)) selected to simulate the molecular trajectories. If the time step is much larger than the velocity correlation time (i.e., \( \vec{\xi}_i \Delta t \gg 1 \)), the solution of eqs 4 and 5a–5d becomes

\[
\vec{\xi}_i(t + \Delta t) = \vec{\xi}_i(t) + \vec{R}_i
\]

where, \( \vec{R}_i \) is the local self-diffusion coefficient of the molecular probes. We can use this simplified equation since the time step \( \Delta t \) used to integrate eq 6a is equal to 1 s, while the friction coefficients (\( \vec{\xi}_i \)) are equal to \( 15 \times 10^{13} \) and \( 6.55 \times 10^{13} \) s\(^{-1} \) for confined and bulk water, respectively. The average molecular displacements (see eq 6b) of the confined water molecules reach \( 45 \) \( \mu m \), i.e., are compatible with the spatial resolution probed by our MRI analysis of the water density profiles.

III. RESULTS AND DISCUSSION

Figure 4 illustrates the results obtained by the measurements of the PGSE-NMR attenuation of the \(^1H\) NMR signal for the water saturated sample of kaolinite. As detailed in eq 2, the decrease in the relative intensity of the NMR signal as a function of the strength of the magnetic field gradient \( G_r \) is perfectly described by a Gaussian relationship, leading to the apparent water self-diffusion coefficient along any direction \( O_x \) of the applied gradient \( G_r \). This field gradient is constructed along six non-collinear directors, noted \( \vec{e}_1 = (1,0,0), \vec{e}_2 = (0,1,0), \vec{e}_3 = (0,0,1), \vec{e}_4 = (1,1,0), \vec{e}_5 = (1,0,1), \vec{e}_6 = (1,1,1) \), where the longitudinal director \( \vec{e}_1 \) is parallel to the cylindrical axis of the clay sample (i.e., \( O_x \)). The six corresponding components of the water self-diffusion tensor are easily determined (see Figure 4) and the principal axes describing the water mobility are identified by tensor diagonalization. A small difference is then detected between the water self-diffusion coefficient in the direction parallel (\( D_{long} = 0.92 \pm 0.02 \times 10^{-9} \) m\(^2\)/s) and perpendicular (\( D_{trans} = 1.05 \pm 0.02 \times 10^{-9} \) m\(^2\)/s) to the cylinder axis which coincides with the compression axis of the dense clay sample (see Section III.2). By dividing the self-diffusion coefficient of bulk water (\( D_0 = 2.1 \times 10^{-9} \) m\(^2\)/s) by its value measured within the dense sample, we obtain an evaluation of the tortuosity \( \theta \) of the clay, noted \( \theta = 2.28 \pm 0.05 \) in the longitudinal direction and \( \theta = 2.00 \pm 0.05 \) in the direction perpendicular to the cylinder axis.

It should be interesting to perform diffusion measurements at various time delay \( \Delta t \) to probe the structural properties of the porous network. As displayed in Figure 5, the measured self-diffusion coefficient is independent of the time delay \( \Delta t \) in the range of accessible values (i.e., between 2 and 150 ms). As a consequence, the water self-diffusion coefficient measured by
relaxation times, the concentration pro-
can be directly compared. Secondly, the MRI signal results
of the clay sediment exhibit some increase in their longitudinal
molecules located in the bulk phase near the upper surface
of the NMR probes. Firstly, as explained in Section II.2,
because of the large difference between their transverse relaxation times, the concentration profiles of bulk \( T_1 = 0.053 \text{ s} \) and confined \( (T_1 = 3 \text{ s}) \text{ whose magnetization is then saturated by this short recycling delay. As a consequence, qualitative analysis must focus}

PGSE-NMR describes the macroscopic mobility of the water molecules diffusing within the porous network of the dense clay sediment.

The water concentration profiles are measured by \(^1\text{H} \) MRI after the addition of bulk heavy water on the top of the water saturated clay sample (see Section II.2). Typical time evolution of the water profiles is illustrated in Figure 6a. As for PGSE-NMR spectroscopy, the intensity of the NMR signals is attenuated by the longitudinal and transverse relaxation times of the NMR probes. Firstly, as explained in Section II.2, because of the large difference between their transverse relaxation times, the concentration profiles of bulk \( (T_2 = T_1 = 3 \text{ s}) \) and confined \( (T_2 = 1.2 \times 10^{-3} \text{ s}) \) water molecules cannot be directly compared. Secondly, the MRI signal results from the averaging of 16 scans with a recycling delay of 0.5 s to sufficiently probe the kinetics of the water exchange between the two media. Since the longitudinal relaxation time of the confined water molecule is short enough \( (T_1 = 0.053 \text{ s}) \), each scan records 100% of magnetization pertaining to the protons from the confined water molecules. This condition is not fulfilled by the protons from the bulk water \( (T_1 = 3 \text{ s}) \) whose magnetization is then saturated by this short recycling delay. As a consequence, quantitative analysis must focus on the relative intensity of the water concentration profiles within the clay sample. Next problem occurs because the total size of the sample slightly exceeds the spatial domain of complete excitation and detection probed by the NMR coil. For this purpose, we must discard magnetization pertaining to water molecules located at distances smaller than 8 mm from the bottom of the detection window whose magnetization is not fully detected. Finally, artifacts also occur at the boundary between the two media, further excluding from the profile analysis the contribution from water molecules located at distances smaller than 2 mm from the interface between the clay sample and the bulk liquid. We are not totally sure about the physical origin of the reported artifact, already detected within dilute clay dispersions.\(^{39}\) We expect that water molecules located in the bulk phase near the upper surface of the clay sediment exhibit some increase in their longitudinal NMR relaxation rate because of the proximity of the paramagnetic impurities (iron) of kaolinite. As a consequence, saturation of their magnetization is reduced with reference to that of "free" bulk water. Another explanation could be the contrast of the magnetic susceptibility between bulk water and the hydrated clay sediment. Obviously, all these artifacts affecting the experimental data are totally excluded from the numerical data, restricting the sample area used for a direct comparison between the data obtained by MRI NMR measurements and BD numerical simulations.

Figure 6 illustrates the evolution of the fraction of confined water molecules obtained by integrating the water concentration profiles between 8.3 and 13.3 mm from the bottom of the detection window. As displayed in Figure 7, the experimental data are fitted by an empirical biexponential law, giving two characteristic times \( (7800 \pm 200) \) and \( (39000 \pm 2000) \) quantifying the time evolution of the integrated water concentration profiles (see Figure 7). These two time scales have no physical meaning since the biexponential fit is simply used for allowing quantitative comparison between the experimental and numerical data. Figure 7 also displays a long-time asymptotic limit \( (0.49 \pm 0.01) \) which may be used to evaluate the porosity (noted \( \phi \)) of the clay sample since it must satisfy the relationship

\[
\text{asymptote} = \frac{\text{total } H_2O}{\text{total } H_2O + D_2O} = \frac{L_1 \phi}{L_1 \phi + L_2} = 0.49
\]

(7)

where, \( L_1 = 12.3 \text{ mm} \) and \( L_2 = 7 \text{ mm} \) are the lengths occupied within the poly(tetrafluoroethylene) (Teflon) cylinder (see

**Figure 5.** Variations of the water self-diffusion coefficient \( D_3 \) (diffusion along the direction \( Oz \)) measured within the saturated clay sample as a function of the time delay \( \Delta \) (see the text).

**Figure 6.** Time evolution of the water concentration profiles: (a) measured by magnetic resonance imaging after the addition of bulk heavy water on the top of the water saturated sample and (b) obtained by numerical simulations of Brownian dynamics (see the text).
between the intrinsic mobility measured by both experimental procedures, validating the use of $^1$H PGSE-NMR to directly determine the intrinsic self-diffusion tensor of confined fluids. Finally, as displayed in Figure 8, a simple straight line roughly reproduces the variation of the intrinsic mobility of the fluid as a function of the porosity of the confining network. The scaling curve displayed in Figure 8 totally validates the use of $^1$H PGSE-NMR to directly measure the water self-diffusion within such porous materials saturated by water molecules.

$^1$H pulsed gradient spin echo attenuation and magnetic resonance imaging of water density profiles were shown to adequately probe the water mobility within natural compacted clay samples. These experimental procedures give additional information on the water mobility within porous networks. On one hand, PGSE-NMR is able to directly measure the various components of the tensor describing the macroscopic equilibrium self-diffusion of water molecules within the hydrated clay sample, by the direct determination of the corresponding self-diffusion propagators. On the other hand, the time evolution of water concentration profiles detected by MRI allows to investigate the kinetics of water exchange between the water saturated clay and added heavy water. In addition, MRI is a powerful tool to determine the impact of sample heterogeneity on the mobility of the confined water molecules. However, in contrast with PGSE-NMR attenuation experiments directly quantify the intrinsic self-diffusion coefficient $D$ of the fluid confined within various porous networks, through-diffusion measurements are sensitive to the fluid permeability of the macroscopic porous media, leading to an effective diffusion coefficient $D_e$. The corresponding intrinsic mobility of the fluid inside the porous network may be evaluated by the simple relationship

$$D = \frac{D_e}{\phi} \quad (8)$$

where, $\phi$ is the porosity of the porous network. This approximation neglects the topological and morphological properties of the porous media. As a consequence, its validity is restricted to homogeneous and isotropic porous network. Figure 8 nevertheless exhibits clear equivalence between the intrinsic mobility measured by both experimental procedures, validating the use of $^1$H PGSE-NMR to directly determine the intrinsic self-diffusion tensor of confined fluids. Finally, as displayed in Figure 8, a simple straight line roughly reproduces the variation of the intrinsic mobility of the fluid as a function of the porosity of the confining network. The scaling curve displayed in Figure 8 totally validates the use of $^1$H PGSE-NMR to directly measure the water self-diffusion within such porous materials saturated by water molecules.

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measurements, the extraction of quantitative data on the water self-diffusion requires a modeling of the H$_2$O/D$_2$O exchange in relation to the time evolution of the water concentration profiles within the clay samples. Since these two complementary NMR measurements are able to investigate dynamical processes over a broad time scale (typically between milliseconds and hours), they lead to additional information on dynamical process investigated by QENS or through-diffusion experiments. Although this study concerns only water and kaolinite clay, it could be easily extended to other molecular and ionic NMR probes diffusing within a large variety of porous networks. Kaolinite, a nonswelling clay, was selected in this study to probe its feasibility because the water molecules are then localized between the clay aggregates and are easily detected by NMR spectroscopy. The same behavior is expected to occur for other nonswelling clays like illite. By contrast, in the case of swelling clays like montmorillonite, the water molecules localized within the interlamellar space is indeed expected to prohibit their detection by PGSE and MRI. The same behavior is easily detected by NMR spectroscopy. The same behavior is expected to occur for other nonswelling clays like illite. By contrast, in the case of swelling clays like montmorillonite, the water molecules localized within the interlamellar space 42,45 cannot be detected by PGSE or MRI NMR spectroscopy. A strong confinement within such interlamellar space 9,43 is indeed expected to enhance the transverse NMR relaxation rate of the water molecules, prohibiting their detection by PGSE and MRI. By contrast, we are quite sure to easily measure, by the same NMR procedure, the mobility of the water molecules located between the aggregates of montmorillonite platelets. In the near future we plan to measure, by the same NMR procedure, the mobility of water molecules and added ions (cations and anions) diffusing within the porous network of dense sediments resulting from the compression of such swelling clays.

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Notes

The authors declare no competing financial interest.

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