

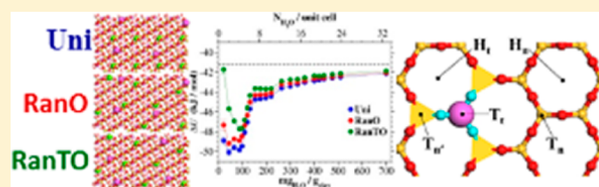
Structural Arrangements of Isomorphous Substitutions in Smectites: Molecular Simulation of the Swelling Properties, Interlayer Structure, and Dynamics of Hydrated Cs–Montmorillonite Revisited with New Clay Models

Brice F. Ngouana W. and Andrey G. Kalinichev*

Laboratoire SUBATECH (UMR-6457), Ecole des Mines de Nantes, 44307 Nantes, France

Supporting Information

ABSTRACT: Three new structural models of montmorillonite with differently distributed Al/Si and Mg/Al substitutions in the tetrahedral and octahedral clay layers are systematically developed and studied by means of MD simulations to quantify the possible effects of such substitutional disorder on the swelling behavior, the interlayer structure, and mobility of aqueous species. A very wide range of water content, from 0 to 700 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$ is explored to derive the swelling properties of Cs–montmorillonite. The determined layer spacing does not differ much depending on the clay model. However, at low water contents up to 1-layer hydrate ($\sim 100 \text{ mg}_{\text{water}}/\text{g}_{\text{clay}}$) the variation of specific locations of the tetrahedral and octahedral substitutions in the two TOT clay layers slightly but noticeably affects the total hydration energy of the system. Using atom–atom radial distribution functions and the respective atomic coordination numbers we have identified for the three clay models not only the previously observed binding sites for Cs^+ on the clay surface but also new ones that are correlated with the position of tetrahedral substitution in the structure. The mobility of Cs^+ ions and H_2O diffusion coefficients, as expected, gradually increase both with increasing water content and with increasing distance from the clay surface, but they still remain 2 to 4 times lower than the corresponding bulk values. Only small differences were observed between the three Cs–montmorillonite models, but these differences are predicted to increase in the case of higher charge density of the clay layers and/or interlayer cations.



1. INTRODUCTION

Clay minerals are ubiquitous in the environment and have many properties, beneficial for their use as natural and engineered barriers in radioactive waste repositories: high sorption and ion exchange capacity, low permeability, swelling in the presence of water, etc.^{1–4} Clay rocks are selected by several national programs as the preferable media for the disposal of medium- and high-level long-lived radioactive waste in deep geological formations.^{5–10}

The most abundant types of clay minerals are illites (nonswelling) and smectites (swelling). Both types belong to the so-called 2:1 clay family.¹¹ Smectites are a major constituent of bentonite¹² used as an environmental buffer separating waste containers from a host rock formation.¹³ Bentonite consists of turbostratically stacked montmorillonite layers forming individual clay particles, and a smaller fraction of accessory minerals.^{14,15} Montmorillonite is also the most widespread and the most common smectite. It has a layered structure made of several stacking TOT units in which one sheet of Al-cations (O) octahedrally coordinated by oxygens and hydroxyls is sandwiched between two sheets of tetrahedrally coordinated Si-cations (T). Isomorphous substitutions (Mg for Al) and (Al for Si), in the octahedral and tetrahedral sheets, respectively, give rise to a negative clay layer charge (typically, between $-0.4|e|$ and $-1.2|e|$ per crystallographic unit cell¹¹ that is compensated

by the presence of cations in the interlayer space and on the surface of montmorillonite particles. The affinity of interlayer cations for water and the relatively weak electrostatic forces holding together montmorillonite sheets lead to an easy hydration of the charge-compensating cations in the presence of water and swelling of the clay.

The properties of cesium-containing clays have already been studied by means of both experimental^{12,16–26} and molecular simulation^{27–39} techniques to investigate hydration and swelling properties, interlayer and interfacial structure and dynamics of Cs^+ ions, and their preferred binding sites in montmorillonite clay. Some of these studies investigated the effect of the total clay layer charge on the above properties.^{12,21,36} Onodera et al.¹² carried out experimental investigations of Cs^+ fixation in smectites. They used several clay samples with variable amounts of tetrahedral and octahedral charge in the clay layers and observed that Cs^+ sorption sites vary with the tetrahedral and octahedral charge amounts. Liu et al.³⁶ used three montmorillonite clay models with different amounts of total layer charge in the octahedral and tetrahedral sheets in their molecular modeling study of the thermodynamic

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70 and structural properties of Cs–smectites. They did not find
71 significant effects of the layer charge distribution on the
72 swelling properties of Cs–smectites, but observed that layer
73 charge distribution have significant influences on the mobility
74 of interlayer Cs^+ and preferred binding sites.

75 The contribution of computational molecular modeling to
76 our understanding of complex materials, including clays, has
77 significantly grown in recent years,^{40–72} and these techniques
78 are capable to quantitatively explore their atomic-scale
79 properties not always accessible to direct experimental
80 measurements. However, more often than not, such simulations
81 are performed using relatively small structural models of clays,
82 which in combination with a relatively low layer charge
83 concentration and the application of periodic boundary
84 conditions (see section 2 below) may result in the exaggeration
85 of the actual degree of ordering in the distribution of the
86 isomorphically substituted sites in the simulated clays. Even the
87 largest simulated models of montmorillonite⁷³ are usually
88 constructed by simply replicating the earlier much smaller
89 structural models to the required size, so the question of the
90 effects of larger-scale disorder in the distribution of octahedral
91 and tetrahedral isomorphous substitutions on the resulting
92 properties of clays remains largely unexplored.

93 The importance of reliable understanding of the predictive
94 capabilities offered by the application of molecular modeling
95 approaches to such systems as Cs-containing clays and of the
96 present limitations of such techniques is only re-emphasized
97 with their use in the wake of the recent Fukushima accident⁷⁴
98 when significant quantities of radioactive materials were
99 released into the environment.^{75–79} Radioisotopes of cesium
100 such as the high-level ^{137}Cs contained in fission products or in
101 spent nuclear fuel were found among these materials.^{80–82}
102 Cesiums isotopes (^{137}Cs , ^{134}Cs , and ^{135}Cs) are normally
103 released as nuclear waste (from nuclear power plants) in
104 which they are one of the most important sources of
105 radioactivity.⁸³ These radioisotopes are highly soluble^{22,84} and
106 highly mobile⁸⁵ in aqueous phase. Hence in the case of their
107 accidental discharge in the environment they are likely to
108 migrate through groundwater into the biosphere²² and have
109 very serious deleterious effects on agriculture, stock farming,
110 and, as a consequence, on human life for many years.⁸⁶

111 The objective of our present work was to systematically
112 investigate any possible correlations between the ordering/
113 disordering in the octahedral and tetrahedral substituted sites of
114 montmorillonite (resulting in the creation of local charge
115 inhomogeneities in the clay structure) and their potential
116 effects on the swelling properties, interlayer structure, and
117 diffusion of Cs^+ ions. For this purpose we have constructed
118 three new larger-scale montmorillonite models with the same
119 atomic composition and the same total amounts of tetrahedral
120 and octahedral charge. The only difference between the three
121 investigated models is in the degree of disorder in the
122 distribution of substituted sites in the octahedral and
123 tetrahedral sheets of the clay layer. The models were
124 constructed taking into account different possible distributions
125 of substituting structural cations in the octahedral^{56,57,87–89} and
126 tetrahedral^{90–92} layers of a dioctahedral smectite. Here we are
127 also taking full advantage of the capability of the CLAYFF force
128 field to more realistically represent the local charge
129 inhomogeneities formed around each specific substituted site
130 in the clay structure.⁵⁸

131 The Cs–montmorillonite system was specifically selected for
132 this study not only because it has significant practical

importance but also because it has already been extensively
studied both experimentally and in molecular simulations.
Because of the particularly low charge/size ratio of the Cs^+
cation, hence its low charge density, one can expect that this
system should have a relatively high sensitivity to the above-
mentioned ordering effects. Such effects observed for the Cs–
montmorillonite system could exhibit themselves less strongly
in the case of other cations, such as Na^+ , Ca^{2+} , or Sr^{2+} .

The construction of the new clay models is fully described in
the next section, followed by the presentation and discussion of
the MD simulation results obtained with these models applied
to the specific case of Cs–montmorillonite. Then we make
detailed comparisons with available experimental data and the
results of previous molecular modeling work for the same
system and formulate several general conclusions for further
investigation.

2. METHODOLOGY

2.1. Structural Models and Force Field Parameters.

The new clay models are based on a $5.16 \times 8.966 \times 9.347 \text{ \AA}^3$
pyrophyllite unit cell structure ($\text{Si}_8\text{Al}_4\text{O}_{20}(\text{OH})_4$) obtained by
single crystal X-ray refinement.⁹³ As in previous simula-
tions,^{27,29,32,35,36,38,40,41,47,54,58,60,68} the clay layer of Wyom-
ing–montmorillonite with the chemical composition was
constructed by introducing the Mg/Al substitutions into one
out of every eight octahedral sites and Al/Si substitutions into
one out of every 32 tetrahedral sites. However, before
introducing these substitutions, the initial pyrophyllite unit
cell was replicated ($4 \times 4 \times 2$) along a , b , and c crystallographic
directions, respectively, resulting in a supercell of two TOT clay
layers of a total of 32 unit cells. This allowed us to prevent
excessive ordering of the substituted sites when the periodic
boundary conditions are applied.

Our most disordered model, corresponding to random
distribution of substitutions in both octahedral and tetrahedral
sheets is presented in Figure 1a,b. However, two less disordered
models were also constructed. One of them, labeled here *Uni*
for further discussion, corresponds to a completely uniform
distribution of substitutions in the octahedral sheet of the TOT
layer (see Figure S1a of the Supporting Information). Care was
taken to have at least one Al atom between two Mg/Al
substitutions in the a and b crystallographic directions. Then
Mg atomic positions of the first TOT layer were shifted in a , b ,
and c directions to get Mg substitution positions in the second
TOT layer (see Figure S1b, Supporting Information). This was
done to avoid any symmetry of the octahedral substitution sites
between the two TOT layers and to ensure the proper layer
stacking. Since there are much fewer Al/Si tetrahedral
substitutions compared to the octahedral ones, they were just
placed in the TOT layer far from one another, not adjacent to
an octahedral substitution, and not at the same positions in the
two tetrahedral sheets of the same TOT layer. All these
substitutions were assigned in a manner to obey the
Loewenstein's rule⁹⁴ in proportion of 4 Al per 124 Si and 8
Mg per 56 Al in each TOT layer in order to obtain the
Wyoming–montmorillonite unit cell chemical composition of
 $M^{+}_{0.75}(\text{Si}_{7.75}\text{Al}_{0.25})(\text{Al}_{3.5}\text{Mg}_{0.5})\text{O}_{20}(\text{OH})_4$, where M^+ is a com-
pensating monovalent cation, Cs^+ in our case.

Another clay model, *RanO*, corresponding to a random
distribution of substitutions in the octahedral sheets, was
obtained from the previous one by randomly changing the
positions of the Mg atoms in the octahedral sheets of both
TOT layers, while keeping the Al/Si substitutions in the 193

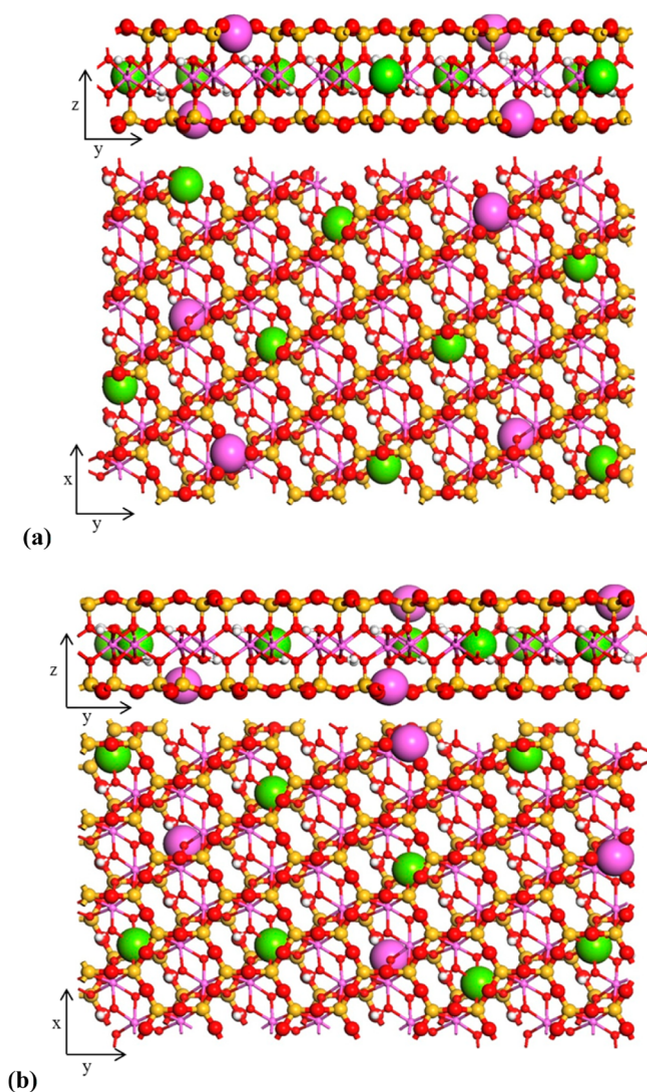


Figure 1. Side and top views of the upper (a) and lower (b) TOT layers of the most disordered clay model *RanTO*. The color code is Si (yellow), Al (pink), Mg (green), O (red), and H (white). Similar structures corresponding to the models *Uni* and *RanO* are provided in the Supporting Information.

194 tetrahedral sheets at the same positions (see Figure S2a,b,
195 Supporting Information).

196 Finally, the clay model *RanTO*, corresponding to random
197 distribution of substitutions in both octahedral and tetrahedral
198 sheets, was obtained from *RanO* by keeping the octahedral
199 substitution positions the same and randomly modifying the
200 positions of tetrahedral substitutions in the TOT layers, while
201 still keeping them obeying the Loewenstein's rule (see Figures
202 1a,b). Chávez-Páez et al.⁵⁴ used a similar approach by
203 modifying the position of one tetrahedral substitution in two
204 clay models.

205 Thus, all three clay models represented a $20.640 \times 35.864 \times$
206 18.694 \AA^3 supercell with 90° between all the pairs of lattice
207 vectors and identical chemical composition. All atomic
208 positions for each of the models are provided in Tables S1–
209 S3 of the Supporting Information.

210 The interatomic interactions in the model systems were
211 described using the partial atomic charges and van der Waals
212 parameters taken from the CLAYFF force field, a fully flexible

general force field suitable for the simulation of hydrated and
213 multicomponent mineral systems and their interfaces with
214 aqueous solutions.⁵⁸ CLAYFF assigns partial charges of
215 different coordinating structural oxygens taking into account,
216 when necessary, their local coordination to the tetrahedral and/
217 or octahedral substituting atoms in the crystal structure⁵⁸ (see
218 Figure 2). This force field has been proven quite successful in
219

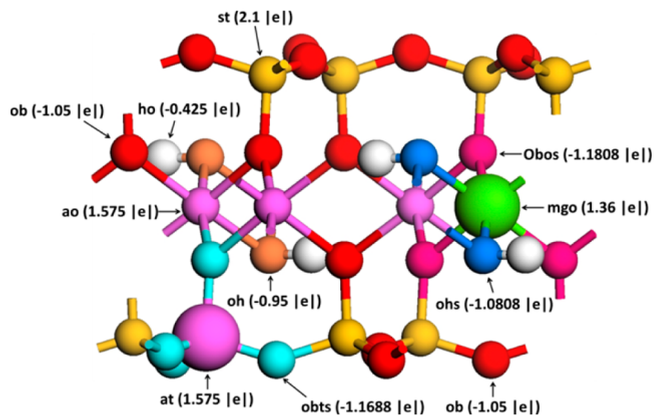


Figure 2. Schematic representation of a montmorillonite unit cell fragment with tetrahedral and octahedral substitutions and different atomic charge distributions according to CLAYFF force field.⁵⁸ O_b (bridging oxygen), O_{bts} (bridging oxygen with tetrahedral substitution), O_{bos} (bridging oxygen with octahedral substitution), O_h (hydroxyl oxygen), O_{hs} (hydroxyl oxygen with octahedral substitution), at (Al in tetrahedral coordination), mgo (Mg in octahedral coordination), ob (Si in tetrahedral coordination).

recent years in extensive molecular simulations of clay and clay-
220 related materials to address a great variety of geochemical and
221 materials science problems.^{36,59,61–63,70–73,95–110} SPC water
222 model¹¹¹ was used to represent the interactions of H_2O
223 molecules among themselves and with the ions of the clay
224 structure for hydrated clays and in solution.
225

The total energy of the systems studied is represented in
226 CLAYFF as the sum of the contributions from the Coulombic
227 (E_{Coul}) and van der Waals (E_{vdW}) interactions of all atomic pairs
228 and the bond stretch (E_{bond}) and angle bend (E_{ang}) interactions
229 for the H_2O molecules and structural OH groups.⁵⁸ The van
230 der Waals interactions are computed by means of the (12–6)
231 Lennard-Jones potential, and the total energy can then be
232 written as
233

$$E_{total} = E_{Coul} + E_{vdW} + E_{bond} + E_{ang} \quad (1) \quad 234$$

with
235

$$E_{Coul} = \sum_{i=1}^{N_{tot}} \sum_{j=i+1}^{N_{tot}} \frac{q_i q_j}{4\pi\epsilon_0 r_{ij}} \quad (1a) \quad 236$$

$$E_{vdW} = \sum_{i=1}^{N_{tot}} \sum_{j=i+1}^{N_{tot}} 4\epsilon_{ij} \left[\left(\frac{\sigma_{ij}}{r_{ij}} \right)^{12} - \left(\frac{\sigma_{ij}}{r_{ij}} \right)^6 \right] \quad (1b) \quad 237$$

$$E_{bond} = \frac{1}{2} \sum_{m=1}^{N_{bond}} k_m (r_m - r_{m0})^2 \quad (1c) \quad 238$$

$$E_{\text{ang}} = \frac{1}{2} \sum_{l=1}^{N_{\text{ang}}} k_l (\theta_l - \theta_{l0})^2 \quad (1d)$$

where N_{tot} , N_{bond} , and N_{ang} are the total number of atoms in the system and total number of bonded and angular terms, respectively. q_i and q_j are the partial charges of the atoms i and j , and r_{ij} is the distance between them. k_m and k_l are the harmonic force constants for bond stretching and angle bending terms for the m th bond and l th angle in the system, respectively, while r_{m0} and θ_{l0} are the equilibrium values of the corresponding bonds and angles. σ_{ij} and ε_{ij} are Lennard-Jones parameters for two interacting atoms i and j , which are obtained from the parameters of the interacting atoms by using the Lorentz–Berthelot combining rules:

$$\sigma_{ij} = \frac{1}{2}(\sigma_i + \sigma_j) \quad (2a)$$

$$\varepsilon_{ij} = (\varepsilon_i \varepsilon_j)^{1/2} \quad (2b)$$

2.2. Simulations Details. The simulated supercell contained 32 clay crystallographic unit cells and two interlayers. Three such supercells were created for each of the three clay models. Before starting the simulations the clay interlayer spaces were increased in the c direction. Then charge-compensating Cs^+ cations ions were placed at the midplane of these interlayer spaces, and water molecules were added randomly (see Figure 3). For each of the three clay models 23

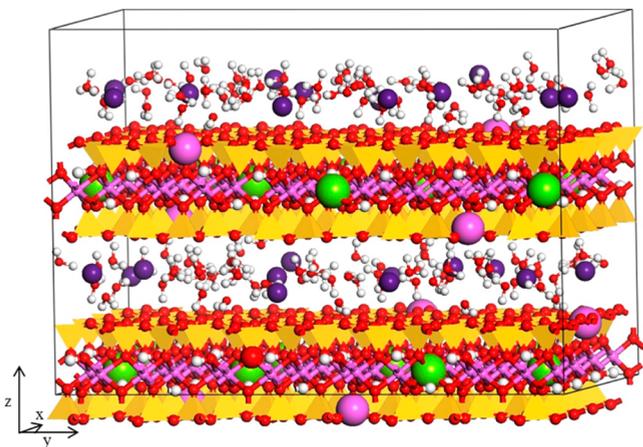


Figure 3. Snapshot of one of the simulated Cs–montmorillonite systems with water content corresponding to 6 H_2O molecules per Cs^+ ion or 5 H_2O per unit cell (1-layer hydrate).

different hydration levels were investigated in a wide range of water contents (from 0 to 700 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$, corresponding to compositions $x = n_{\text{H}_2\text{O}}/(\text{clay unit cell})$ ranging from 0 to 32). Thus, all three models at each hydration level had the same chemical composition of $\text{Cs}_{2.4}(\text{Si}_{2.48}\text{Al}_8)(\text{Al}_{11.2}\text{Mg}_{16})\text{O}_{640}(\text{OH})_{128}\cdot 32\text{H}_2\text{O}$. The total energy of each system for each hydration level was then minimized, allowing initial adjustments of the simulation supercell lengths.

The swelling behavior of these Cs–montmorillonite models was studied by performing 2 ns MD runs in the NPT -ensemble at $T = 298$ K and $P = 1$ bar for each of the three clay models and 23 different hydration levels. During these runs, the simulated supercell dimensions were allowed to vary independently of each other and adjust according to the water content. The last 1 ns of each trajectory was taken to

compute statistical averages for a total of 10 equal blocks, to determine the supercell dimensions, the basal spacing, hydration energy, and immersion energy. These properties were computed according to the procedures described in the next subsection. Their error bars were estimated using the block averaging method¹¹² and presented with 95% confidence interval.

The structural and dynamical properties of Cs and water were determined for 4 different hydration levels, after 1.5 ns NVT -ensemble MD simulations at 298 K. The simulation supercells of these NVT runs were adjusted to the average equilibrium supercell dimensions obtained from the previous NPT runs for each of the 4 hydration levels. The equilibrium parts of the NVT trajectories were then analyzed to determine atomic density profiles, radial distribution functions, and diffusion coefficients of the Cs^+ ions and H_2O molecules in the hydrated clay media.

LAMMPS simulation package¹¹³ was used for all MD runs with the integration time step of 1 fs. A spherical cut off of 10.0 Å was applied for short-range van der Waals interactions, while the long-range electrostatic interactions were treated using the Ewald summation method¹¹⁴ with a precision of 10^{-4} in combination with the direct pairwise interactions calculation within a spherical cut off of 10.0 Å.

2.3. Simulation Analyses. **2.3.1. Swelling Properties.** The basal spacing as a function of water content essentially shows how the volume of the system changes with increasing (or decreasing) hydration. At the end of the NPT runs, the basal spacing (d) was calculated as follows:

$$d = \frac{\langle a \rangle \langle b \rangle \langle c \rangle}{2 \langle a \rangle \langle b \rangle} = \frac{\langle c \rangle}{2} \quad (3)$$

where $\langle a \rangle$, $\langle b \rangle$, and $\langle c \rangle$ are the statistically averaged dimensions of the simulation supercell.

In order to quantify the thermodynamics of hydration and swelling for three different Cs–montmorillonite models, we calculated the hydration energy, immersion energy,^{27,36,40,41,96–98,100} and the isosteric heat of adsorption²⁷ at each of the hydration states.

The hydration energy (ΔU) evaluates the energy change associated with water uptake by the dry clay, and can be calculated using the generic formula:

$$\Delta U = \frac{\langle U(N) \rangle - \langle U(0) \rangle}{N} \quad (4)$$

where N is the number of interlayer water molecules for a given hydration level, while $\langle U(N) \rangle$ and $\langle U(0) \rangle$ are the average potential energies of the hydrated clay (with N water molecules) and dry clay ($N = 0$), respectively.

The immersion energy (Q) is the energy consumed or released when the clay system at a given hydration level is brought to another hydration level by adding water to the system:

$$Q = \langle U(N) \rangle - \langle U(N_0) \rangle - (N - N_0)U_{\text{bulk}} \quad (5)$$

where N_0 and $\langle U(N_0) \rangle$ are the number of water molecules and the average potential energy of a reference hydration level. U_{bulk} is the average potential energy of bulk SPC water that was computed in this work from preliminary 500 ps NVT -ensemble MD simulations of 1024 water molecules at 298 K and a water density of 0.9942 g/cm^3 , following a 100 ps equilibration in the NPT -ensemble at the same temperature and pressure of 1 bar.

333 Considering clay at a certain hydration level, the isosteric
334 heat of adsorption is the measure of energy released or
335 consumed by the system when a small amount of water is
336 added or removed from the system. This property is calculated
337 from simulations as follows:

$$338 \quad q_{\text{st}} = RT - \frac{\langle U(N) \rangle - \langle U(N') \rangle}{(N' - N)} \quad (6)$$

339 where N and N' are two consecutive hydration levels, R is the
340 ideal gas constant, and T is the temperature. The factor of RT is
341 needed for conversion between energy and enthalpy.

342 **2.3.2. Structural and Dynamical Properties.** Atom–atom
343 radial distribution functions of the interlayer species and their
344 atomic density profiles perpendicular to the layering were
345 calculated to analyze the structural properties of the swelling
346 clay. Both properties were averaged over the two statistically
347 independent interlayer spaces and over entire trajectories from
348 the equilibrium NVT -simulations. Density profiles, $\rho_{\alpha}(z)$, were
349 determined as the average number of atoms, $\langle N_{\alpha}(\Delta z) \rangle$, of
350 certain type α in a slice of thickness Δz parallel to the clay
351 surface, normalized by the system volume, V :

$$352 \quad \rho_{\alpha}(z) = \frac{\langle N_{\alpha}(\Delta z) \rangle}{V} \quad (7)$$

353 The coordination of atoms in clay interlayers with respect to
354 each other (eq 9) was derived from the radial distribution
355 function (eq 8):

$$356 \quad g_{\alpha\beta}(r_{\alpha\beta}) = \frac{\langle N_{\alpha\beta} \rangle}{4\pi\rho_{\beta}r_{\alpha\beta}} \quad (8)$$

$$357 \quad n_{\alpha\beta}(r_{\alpha\beta}) = 4\pi\rho_{\beta} \int_0^{r_{\alpha\beta}} g(r)r^2 dr \quad (9)$$

358 where $r_{\alpha\beta}$ is the distance between atoms α and β , ρ_{β} is the
359 number density of atoms β , and $\langle N_{\alpha\beta} \rangle$ the average number of
360 atoms β found at the distance $r_{\alpha\beta}$ from atom α .

361 The self-diffusion coefficients of interlayer species were
362 determined from their mean squared displacement (MSD)
363 according to the Einstein relationship:¹¹⁴

$$364 \quad \frac{1}{N} \sum_{i=1}^N \langle |r_i(t) - r_i(t_0)|^2 \rangle = 2dDt \quad (10)$$

365 where N is the number of atoms of interest, $r_i(t)$ is the position
366 of atom i at time t , and the angular brackets indicate the
367 averaging taken over all time origins t_0 along the MD trajectory.
368 D in eq 10 is the diffusion coefficient, and d is the
369 dimensionality of the system equal to 1, 2, or 3 for the
370 calculation of one-dimensional, two-dimensional, or three-
371 dimensional diffusion coefficients, respectively.

372 From this equation we are then able to determine individual
373 special components of the diffusion coefficients of interlayer
374 species. It is known from previous simulations of clays and
375 other layered systems^{3,35,36,38,70,109} that for very long simulation
376 times, the diffusion of aqueous species in the clay interlayers is
377 mostly two-dimensional and occurs along the directions parallel
378 to the clay surfaces (xx and yy). The diffusion is significantly
379 restricted along the direction normal to the layering (zz). This
380 was also observed in our investigated clay systems that were ran
381 for 1 ns each. This means that the two-dimensional D_{xy}
382 diffusion coefficient is a reasonable estimate of the diffusion
383 of aqueous species in the interlayer spaces. However, we

consider that the small value of the restricted normal
component of diffusion is due to the presence of the clay
layers that limit the motions of interlayer species in the
considered directions, and this normal component has to be
taken into account during the determination of the total
diffusion coefficients for comparison with the bulk case. In
particular, when comparison with experimental data is
concerned, we know that these data are determined for three-
dimensional diffusion.

Another important question is the comparison of diffusion
coefficients in the clay interlayers with their corresponding bulk
values between various experimental and simulation sources. In
this case, following the previous work of Kosakowski et al.³⁵
and Bourg and Sposito,⁷⁰ we are using relative diffusion
coefficients to minimize the uncertainties introduced by the
potential inaccuracy in reproducing experimentally observed
bulk diffusion coefficients by different force field models for
 H_2O and ions and to most directly compare the retardation or
acceleration of molecular mobility in the clay interlayers with
respect to the bulk solution:

$$D_t = \frac{D_{\text{interlayer}}}{D_{\text{bulk}}} \quad (11) \quad 404$$

In section 3.6, we are presenting such 3D relative diffusion
coefficients to compare the results obtained using our three
new clay models with previous simulations and experimental
data.

The MSDs used to derive the diffusion coefficients were
calculated from the last 500 ps of our 1 ns equilibrium MD
trajectories. A time interval of 0.5 ps was used between two
consecutively saved states. The quasi-linear part of the MSDs
was divided into 4 blocks. The diffusion coefficients and their
statistical uncertainties were then determined by block
averaging of the 4 independent MSD slopes and assuming d
= 3 in eq 10. Diffusion coefficients of Cs^+ ions and H_2O
molecules in bulk aqueous solution were also determined
following the previously described procedure from 100 ps NPT
and 500 ps NVT MD simulations of a system of 16 (Cs^+ , Cl^-)
pairs dissolved in 1024 water molecules using the same force
field parameters as for the clay simulations. All the diffusion
data from this work are shown with a $\pm 2\sigma$ confidence interval.

3. RESULTS AND DISCUSSION

3.1. Layer Spacing. Figure 4 illustrates the variation of
basal layer spacing in Cs –montmorillonite as a function of
water content from our simulations and provides a comparison
with available experimental results.^{17–19,23} There are no
observable differences on the calculated layer spacing between
our three clay models with different distribution of
substitutions. For all three models the dry clay layer spacing
of 10.60 Å is similar to the simulation (10.50–10.70 Å) and 5%
lower than experimental values (11.2 Å) that are provided in
Table 1. For all three clay models, a plateau corresponding to
the formation of a stable monolayer hydrate is clearly
observable at ~ 12.6 Å, in good agreement with the range of
12.3–12.6 Å reported from experiments^{17–19,23} and 12.2–12.9
Å obtained from previous simulations.^{27,31,35,36} In our
simulations, this monolayer hydrate corresponds to $x = 4.5$,
which is in the range of 4.0–5.5 observed in previous
simulation studies (see Table 1).

Table 1 also shows comparison between our data and other
simulation results for layer spacings corresponding to stable 2-
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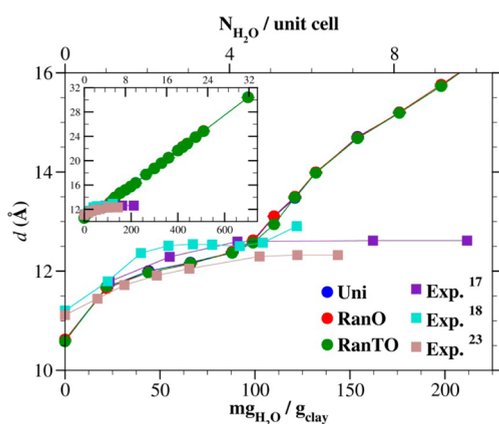


Figure 4. Swelling curves of Cs–montmorillonite simulated with three new clay models *Uni* (blue), *RanO* (red), and *RanTO* (green). Available experimental data are shown for comparison. The error bars for the simulated values, calculated with a 95% confidence interval, are within the size of the symbols.

Table 1. Basal Spacing (in Å) of 1-, 2-, 3-, and 4-Layer Hydrated Cs–Montmorillonites

hydration state	present simulation results	other simulations	experiments
0W	10.60	10.53 ²⁷ 10.60–10.70 ³¹ 10.50 ³⁶	11.2 ^{18,23}
1W	12.60	12.68 ²⁷ 12.68 ²⁹ 12.7 ³¹ 12.7, 12.9 ³⁵ 12.6–12.7 ³⁶ 12.22 ³⁷	12.57 ^{17,18} 12.3 ²³
2W	16.34	15.74–16.33 ²⁷ 12.0–12.5 ²⁸ 15.1 ³⁵ 16.3 ³⁶ 14.88 ³⁷	
3W	19.58	18.36 ³⁵ 17.23 ³⁷	
4W	22.23	22.01 ²⁷ 21.87 ³⁵	

442 layer hydrate ($x = 10.0$ in our simulations and 8.0 – 11.0 for
443 other simulations), 3-layer hydrate ($x = 15.0$ in our simulations
444 and 12.0 – 15.0 for Kosakowski et al.³⁵), and 4-layer hydrate (x
445 = 19.1 in our simulations and $x = 20$ in ref 35). The equilibrium
446 values of layer spacing for these multilayer hydrates cannot be
447 determined based on the swelling curve alone (Figure 4). In
448 addition, the calculations of the energetic characteristics of Cs–
449 montmorillonite swelling were used, as discussed in the next
450 subsections. On average, our layer spacing values for these
451 multilayer hydrates are somewhat higher than the ones
452 observed in other simulations, but they are very close to the
453 simulation data that also used the CLAYFF force field.³⁶ The
454 differences with other simulation data can be attributed to the
455 different force fields used,^{27–31,35,36,38} to the slightly different
456 clay composition with no tetrahedral charge,^{28,31,35} or the
457 presence of both Na^+ and Cs^+ counterions in the clay
458 interlayers.^{31,35}

459 In the range from $x = 0$ to $x = 4.5$ (0 to 100 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$) the
460 simulated layer spacings are always very close to the

experimental ones, being just slightly lower^{17–19} or slightly
higher,²³ depending on the sample. Calvet^{18,19} and Mooney et
al.¹⁷ performed their measurements for montmorillonite
samples having no tetrahedral charge and with total layer
charges of 0.78lel and 0.66lel , respectively. The composition of
the montmorillonite sample used in the more recent experi-
ments of Berend et al.²³ was the closest to ours (the tetrahedral
charge of 0.22lel and octahedral charge of 0.526lel). This may be
the reason why for most of the points our results are closer to
the ones of Berend et al.²³ and the small differences may be due
to different temperatures used (25 °C in this article and 20 °C
for the mentioned experimental works).

Beyond $x = 4.5$ (100 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$) experimental data indicate
invariable layer spacings unlike the simulation swelling curves
that increase. One has to remember, however, that the
experimentally measured water contents include contributions
not only from the H_2O molecules in clay interlayers but also
from the ones adsorbed on the external surfaces of clay particles
and from the interparticle pore-space (see, e.g., ref 27).
Therefore, when the clay interlayer is stabilized ($x = 4.5$) the
additional water molecules will preferably adsorb onto the
external surfaces, resulting in no noticeable change in the layer
spacing value as seen in the experimental data (Figure 4).
Besides, one may also consider that clay swelling occurs
through the formation of mixed-layer hydrates so that the
observed layer spacing is averaged over the interstratified
structure,¹¹⁵ and this may be different from the calculated layer
spacing of the uniformly expanded hydrated clay from the
simulations that are only due to interlayer water.

3.2. Hydration Energy. The hydration energies of our Cs–
montmorillonite models are presented in Figure 5 together

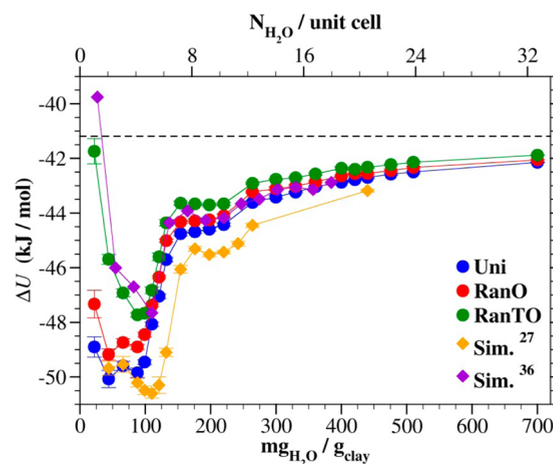


Figure 5. Hydration energy curves for Cs–montmorillonite simulated with the three new clay models and their comparison with previous simulations. The error bars are shown with a 95% confidence interval.

with the results of two other simulations.^{27,36} All results exhibit
two minima for water contents of $x = 4.5$ (~ 100 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$)
and $x = 10.0$ (~ 220 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$). As expected,^{27,107} the
hydration energies tend to approach the value of internal
energy of bulk water as the water content increases. A
comparison between our three clay models shows that the
model *Uni* has lower hydration energies compared to the two
more disordered models, *RanO* and *RanTO*. However, the
energies of *Uni* and *RanO* models are closer to each other than
any of these two models with the *RanTO* model. These
observations suggest that the variation in the positions of

503 octahedral substitutions has a lesser effect on the hydration
 504 energy (from *Uni* model to *RanO* in Figure 5) than a similar
 505 variation in the tetrahedral sheet (from *RanO* model to *RanTO*
 506 model in Figure 5). In addition, one can see that modifying the
 507 substitution positions in both octahedral and tetrahedral sheets
 508 affects the hydration energy most (from *Uni* model to *RanTO*
 509 model) and that the observed differences are very prominent
 510 especially at lower water contents before decreasing with
 511 increasing hydration. Nevertheless, the compositions of the
 512 monolayer and bilayer hydrates are the same for all three
 513 models. Further analysis of the swelling energetics based on the
 514 calculation of the immersion energy and isosteric heat of
 515 adsorption provides additional information for the clear
 516 identification of the water contents corresponding of the stable
 517 1W and 2W hydration states and can also help to assign water
 518 contents for the hypothetical 3W and 4W hydration states.

519 The compositions of the monolayer and bilayer hydrates
 520 obtained by Liu et al.³⁶ and by Smith²⁷ are found, respectively,
 521 at water contents of $x = 5$ and $x = 10$ (110 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$ and 220
 522 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$) and $x = 5.5$ and $x = 11$ (120 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$ and 230
 523 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$). From our hydration energies calculations, the
 524 stable monolayer and bilayer compositions correspond to water
 525 contents of $x = 4.5$ (100 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$) and $x = 10.0$ (220
 526 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$). This means that slightly less water is required in
 527 our case to complete the formation of the stable monolayer and
 528 bilayer hydrates as compared to the results of Smith.²⁷ A
 529 comparison of the monolayer hydration energies between the
 530 three simulations shows that Smith's²⁷ value is lower than the
 531 others, while the energies obtained with the two models labeled
 532 *RanO* and *RanTO* are very close to the values of Liu et al.³⁶
 533 This observation is the same for the bilayer and for higher water
 534 contents. This is not surprising since Liu et al.³⁶ used the same
 535 CLAYFF force field (and the SPC model of water) just as we
 536 did, but with a set of initial atomic positions⁴⁴ that differs from
 537 ours. In contrast, Smith²⁷ has employed the SPC/E water
 538 model and Skipper's force field and atomic positions within a
 539 rigid clay model.^{43,44} Hence the differences/similarities in the
 540 initial atomic configuration and force field used can be
 541 responsible for the differences/similarities observed between
 542 our results and the other simulation results presented.

543 The small differences observed between Liu et al.'s³⁶ results
 544 and ours show that the swelling energetics can be quite
 545 sensitive to variations in the specific clay structure with different
 546 locations of substitution sites, even when the total layer charge
 547 and the chemical composition remain the same. The
 548 observation stands for all three clay models discussed in this
 549 work since the observed differences can only be attributed to
 550 the differently distributed substitution sites among the
 551 tetrahedral and octahedral layers because the three models
 552 were all treated with the same procedure.

553 **3.3. Immersion Energy and Isosteric Heat of**
 554 **Adsorption.** Figures 6 and 7 show the immersion energies
 555 and isosteric heat of adsorption, respectively, for all three
 556 models. These two observables provide additional useful
 557 information helping to identify the stable hydration states.
 558 Apart from the completely dry state corresponding to the first
 559 points on the plots, the immersion energies calculated at all
 560 other water contents are very similar between the three clay
 561 models. Figure 6 allows us to better distinguish the previously
 562 observed monolayer and bilayer hydrates through the local
 563 minima for $x = 4.5$ ($\sim 100 \text{ mg}_{\text{water}}/\text{g}_{\text{clay}}$) and $x = 10.0$ (~ 220
 564 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$) for all simulation data presented. Two hypo-
 565 thetical 3-layer and 4-layer hydrates can also be located at water

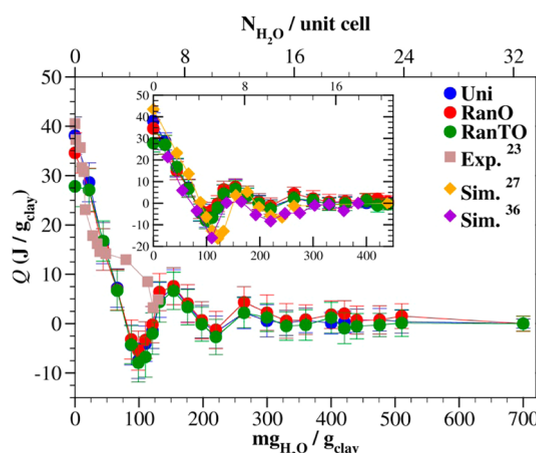


Figure 6. Immersion energy curves simulated for the three new Cs–montmorillonite models and their comparison with experimental data and previous simulations. The error bars are shown with a 95% confidence interval.

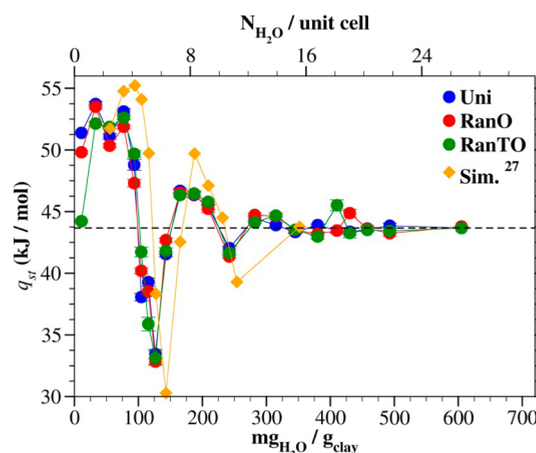


Figure 7. Isosteric heat of adsorption simulated for the three new Cs–montmorillonite models. The error bars are shown with a 95% confidence interval.

566 contents of 330 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$ ($x = 15.0$) and 420 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$ ($x = 19.1$), that are visible for all three clay models. Though, it is
 567 clear on Figure 6 that the immersion energies found beyond
 568 300 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$ do not vary much.

569 However, the hypothetical 2W hydrates have been earlier
 570 observed in the simulation studies,^{27,36} and in our work, the
 571 hypothetical 3W and 4W hydrates. Cs–montmorillonites are
 572 known to form stable monolayer hydrates in water.¹¹⁵ Our
 573 calculations of hydration and immersion energies confirm that
 574 the monolayer hydrate is the most stable hydration state. The
 575 hydration states beyond 1-layer hydrate (and to some extend 2-
 576 layer hydrate) are not observed in experiments under ambient
 577 conditions. However, they can be easily probed in the
 578 simulations through “forced hydration” of the interlayer spaces
 579 by adding more H_2O molecules to each interlayer.^{27,36} For
 580 certain water contents the addition of a small amount of H_2O
 581 molecules can induce not only the expansion of the clay layers
 582 as shown by the increase in layer spacing (Figure 4) but also
 583 reorientations and rearrangement of the interlayer water
 584 molecules resulting in a small decrease of the total energy of
 585 the system and the formation of hypothetical 3W and 4W
 586 hydrates (minima in Figure 6 and specified intersections in 587

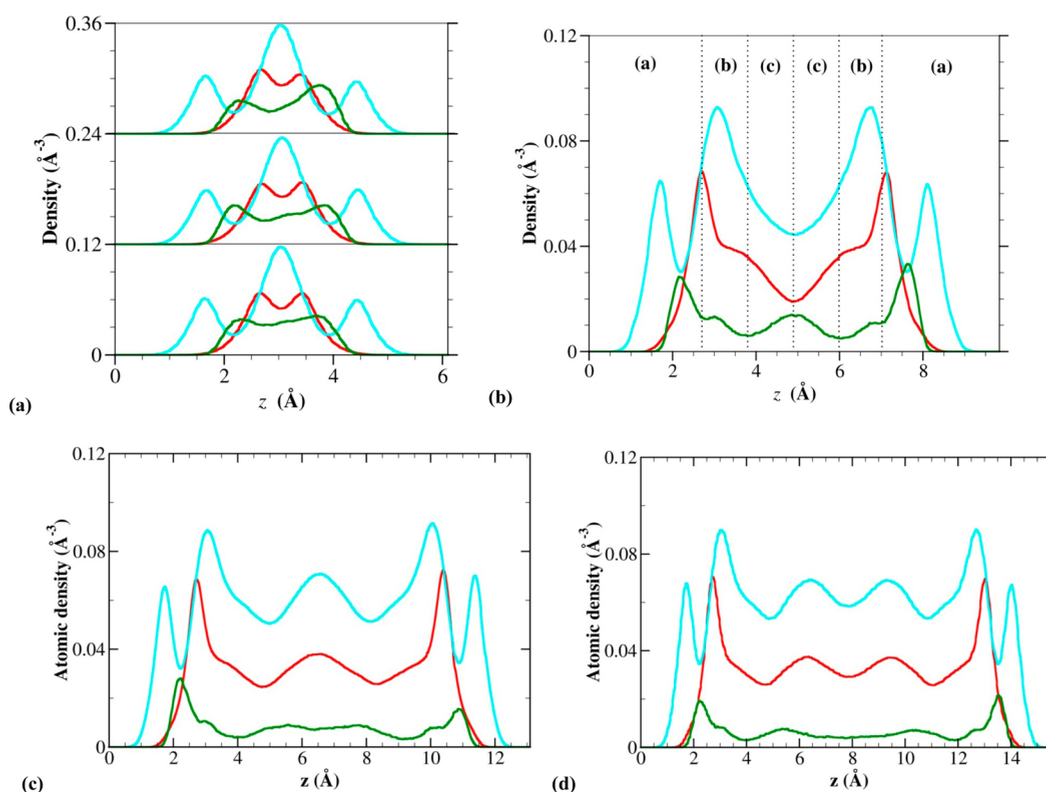


Figure 8. Atomic density distributions along the direction normal to the clay surface, for Cs^+ ions (green), water oxygens (red), and water hydrogens (cyan). The monolayer (a) is shown for three clay models, *RanTO* (topmost), *RanO* (middle), and *Uni* (undermost). The bilayer (b), trilayer (c), and tetralayer (d) hydrates are also shown only for the clay model *RanTO*. Different regions are defined according to Cs^+ ion distributions as shown in panel b.

588 Figure 7 around 330 and 420 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$). Unlike the previous
 589 simulation studies,^{27,36} these hypothetical 3W and 4W hydrates
 590 could be observed in our work because we have systematically
 591 probed hydration at water contents up to $x = 32$ (700 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$). They are studied here as a sensitive probe to see how
 592 much the substitution distribution could affect the formation of
 593 successive water layers of the swelling clay.

594 The calorimetric data of Berend et al.,²³ one of the rare cases
 595 when experimental data are available for direct comparison, are
 596 displayed in Figure 6 together with our simulation data. These
 597 data were digitized from published plots of ref 23 and rescaled
 598 to present the immersion enthalpy as a function of water
 599 content rather than the original relative humidity variable. In
 600 order to be consistent with those data for which bulk water was
 601 used as the reference state for immersion enthalpy calculations,
 602 and following the work of Liu et al.,³⁶ the highest water content
 603 (700 $\text{mg}_{\text{water}}/\text{g}_{\text{clay}}$, or $x = 32$) was taken as the reference state in
 604 eq 6. We also recalculated the immersion energies from Smith²⁷
 605 using the same consideration (Figure 6).

606 One can see in Figure 6 that the simulation closely follows
 607 experiment²³ for dry and nearly dry clays, but deviates when
 608 approaching the monolayer range. The immersion energy
 609 decreases close to zero (slightly less than zero for our results
 610 and slightly more for the experimental ones) where the stable
 611 monolayer hydrate is being formed. As explained by Smith,²⁷
 612 for a clay that forms monolayer hydrates, the value of the
 613 immersion energy would oscillate around the 1-layer hydrate,
 614 indicating that small changes in the energy value are associated
 615 with external surface hydration. The differences observed
 616 between our results and other simulation results^{27,36} can be
 617 attributed, as before, mostly to the different force field used

and, to some extent, to the location of the substituted sites in
 the clay layers. An addition source of discrepancy is introduced
 by the average value of bulk water energy, which is used in
 deriving the immersion energy as shown by eq 6 and which
 differs between different water models. In our case this value is
 equal to $-41.2 \text{ kJ mol}^{-1}$ (SPC water), while Smith²⁷ used the
 value of $-41.4 \text{ kJ mol}^{-1}$ (SPC/E water), and Liu et al.³⁶ just
 used the experimental value of $-43.9 \text{ kJ mol}^{-1}$ in their
 calculations.

The immersion energies of the dry clay calculated for our
 three models all lay in the range 25–40 J/g_{clay} with the *Uni*
 model being the closest to the experimental value of $\sim 40 \text{ J/g}_{\text{clay}}$.
 The immersion energy value for a dry clay was not provided by
 Liu et al.³⁶ After recalculation, Smith's dry clay immersion
 energy gives $\sim 43 \text{ J/g}_{\text{clay}}$, which is closer to experimental value
 than the value of 60 J/g_{clay} reported in the original paper²⁷
 where the reference hydration level did not correspond to the
 highest water content. This clearly indicates that the most
 convenient way to evaluate immersion energies from
 simulations using eq 5 would be to take the highest water
 content as the reference hydration level, $\langle U(N_0) \rangle$, since it will
 be closer to bulk water than the 1-layer hydrate used by
 Smith.²⁷

The isosteric heat of adsorption has been introduced and
 determined before²⁷ and is calculated here for our three new
 clay models. It is equivalent to the negative of the differential
 hydration enthalpy, which, in turn, corresponds to the change
 in enthalpy of a clay of given water content, per mole of water,
 upon adsorption of an additional infinitesimal amount of water.
 The points where the heat of adsorption crosses the bulk
 vaporization enthalpy of water with negative slope correspond

650 to stable hydration states.²⁷ We can then clearly observe the 1-,
651 2-, 3-, and 4-layer hydrates in Figure 7.

652 The swelling curves do not show significant differences
653 between the three clay models used, and all of them yield a
654 good prediction of the experimental ones. The swelling
655 energetics clearly shows that immersion energies are only
656 weakly sensitive to the actual distribution of substitutions in the
657 clay sheets, especially at very low water contents. This effect is
658 more pronounced for the hydration energies (Figure 5), but it
659 always weakens as the water content increases.

660 **3.4. Atomic Density Profiles in the Clay Interlayer.** The
661 distribution of interlayer water molecules and Cs⁺ ions as a
662 function of their distance from the surface was determined for
663 1-, 2-, 3-, and 4-layer hydrates ($x = 5, 10, 15,$ and $20,$
664 respectively), as shown in Figure 8. Experimental studies of
665 Cs–montmorillonites did not observe the hydration states
666 higher than the monolayer hydrate. Hence, as discussed above,
667 the three other hypothetical hydration states are investigated in
668 this study to see whether the proposed clay models could have
669 different behavior at different hydration states and also to make
670 better comparisons with other simulation studies. The atomic
671 distributions shown in Figure 8 were calculated by averaging
672 over two statistically independent interlayer regions of each
673 model containing the same amounts of water molecules and Cs
674 ions. Cs density in Figure 8 is magnified by a factor of 5 for
675 clarity.

676 In the monolayer hydrate, for our three clay models, Cs⁺ ions
677 and oxygens of H₂O molecules (O_w) have a double-peaked
678 distribution in the interlayer regions with Cs⁺ found around 2.3
679 Å from the clay surface (defined here by the time-averaged
680 positions of the bridging oxygens of the surface siloxane rings
681 (see Figures 2 and 3)). The two distinguishable peaks of these
682 distributions indicate the molecular organization near the clay
683 surfaces. Smith²⁷ and Marry et al.³¹ have found that Cs⁺
684 distributes as a single sharp peak in the middle of the interlayer
685 for water content equivalent to a monolayer hydrate, thus
686 showing a lower affinity of Cs⁺ to the clay surface than observed
687 in our simulations. Both these previous simulations have used
688 Skipper's rigid clay model and the force field,^{43,44} which does
689 not represent the charge delocalization around the substituted
690 sites sufficiently accurately.⁵⁸ Moreover, even if Smith²⁷ used
691 the same amounts of tetrahedral and octahedral substitutions as
692 we did, Marry et al.³¹ used a montmorillonite model with only
693 octahedral substitutions. However, it is well-known that the
694 amount of tetrahedral substitutions influences the distribution
695 of ions in montmorillonite interlayers.^{32,44} These reasons for
696 the observed disagreements can also be confirmed by the
697 consistency of our results with those of Liu et al.,³⁶ who had
698 also used the CLAYFF force field in their simulations.

699 For all three clay models, as we increase the water content to
700 the bilayer hydration state, Cs⁺ ion distribution splits into two
701 pairs of symmetric peaks plus a central peak. The highest peak
702 is located around 2.1 Å from the clay surface indicating a
703 tendency for Cs⁺ ions to move closer to the clay surface with
704 increased water content and form relatively strong inner-sphere
705 surface complexes, as observed in previous experimental^{116,117}
706 and simulation^{3,27,31,35,36} studies. These inner-sphere complexes
707 can occupy hexagonal cavities and triangular sites^{27,116,118,119}
708 (see Figure 9). Therefore, Cs⁺ ions found at ~ 2.17 Å from the
709 clay surface form inner sphere complexes located in the
710 hexagonal cavities, while inner-sphere complexes formed with
711 the oxygens of the triangular Si or Al sites are indicated by the
712 second lower peak at the clay surface found around 3.1 Å. The

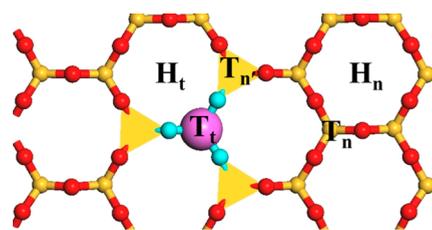


Figure 9. Identification of different binding sites on the clay surface: “normal” hexagonal site without substitution (H_n), hexagonal site adjacent to tetrahedral substitution (H_t), “normal” triangular site without substitution (T_n), substituted triangular site (T_t), and triangular site adjacent to tetrahedral substitution (T_n).

713 third Cs⁺ peak is found at ~ 5.0 Å and represents the outer
714 sphere complexes formed by hydrated cesium ions.

715 As the water content increases from bilayer to tri- and quad-
716 layer hydrate, the location of the first and second peaks of the
717 Cs⁺ density profiles remain about the same, whereas the third
718 peak now shifts to 5.3–5.4 Å from the clay surface, indicating
719 that these Cs⁺ ions are not in a relatively stable outer-sphere
720 coordination to the surface but rather in a diffuse aqueous
721 layer.¹¹⁷ There are no experimental data available for
722 montmorillonite for comparison with these simulations.
723 However, available synchrotron X-ray reflectivity measure-
724 ments¹²⁰ locate Cs⁺ around ~ 2.15 Å from the (001) plane of
725 the muscovite surface. Muscovite also has a 2:1 clay structure
726 with a layer charge almost 3 times higher than montmorillonite
727 and entirely concentrated in the tetrahedral layers.¹¹⁵ Because
728 of its favorable crystal cleavage properties it is a good model for
729 experimental studies of phenomena occurring at water–mineral
730 interface, unlike montmorillonite.⁶⁷ With its high and fully
731 tetrahedral layer charge then, we expect to find Cs⁺ ion slightly
732 closer to muscovite surface than to montmorillonite surface,
733 indirectly indicating good agreement of our simulations with
734 available experimental data.

735 For the hydration states above the monolayer hydrate ($x =$
736 10 to 20) and for all three clay models, Cs⁺ ions mostly form
737 inner-sphere complexes represented by the peaks at ~ 2.3 and
738 ~ 3.1 Å. This is comparable with other simulation data that may
739 have used either different atomic data source or force field and
740 that report Cs–montmorillonite distances of $\sim 2.8,$ ³¹ $\sim 3.0,$ ³⁵
741 ~ 2.2 – $2.5,$ ³⁶ and ~ 2.8 Å³ at water contents higher than the
742 monolayer.

743 **3.5. Atom–Atom Pair Correlation Functions, Lateral**
744 **Atomic Density Distributions, and Cs⁺ Coordination**
745 **Numbers in the Clay Interlayers.** According to the density
746 profiles presented and discussed above, the clay interlayer space
747 can be divided into 3 regions differing by their representative
748 distance from the surface, where Cs⁺ are preferentially located.
749 They are labeled (a), (b), and (c) in Figure 8b. The region (a)
750 (between 0 and ~ 3 Å from the surface includes Cs⁺ ions
751 forming inner-sphere complexes with the hexagonal sites on the
752 clay surface (sites H_n and H_t of Figure 9). The region (b)
753 between ~ 3 and ~ 4 Å from the surface is characterized by the
754 Cs⁺ ions in the inner-sphere coordination to the triangular sites
755 on the clay surface (sites T_n, T_n, and T_t of Figure 9). The
756 region (c) (from ~ 4 to ~ 5 Å and more from the surface,
757 depending on the hydration state) comprises cesium ions in the
758 outer-sphere coordination to the surface and/or in the diffuse
759 aqueous layer. The region (a) exists for all the hydration states
760 studied, while the regions (b) and (c) do not develop for the
761 monolayer hydrate and below. Unlike other simulation

Table 2. Positions of the First Maxima (R_{\max}) and Minima (R_{\min}) of the RDF and the Contribution of Different Oxygen Atoms (O_b , O_{bts} , and O_w) to the First Coordination Shell of Cs^+ Ions in the Interlayer of Hydrated Montmorillonite

hydration state	region	R_{\max} (Å)				R_{\min} (Å)				N_O				
		O_b	O_{bts}	O_w	O_w (bulk)	O_b	O_{bts}	O_w	O_w (bulk)	O_b	O_{bts}	O_w	O_{tot}	O_w (bulk)
1W	a	3.42	3.15	3.05	3.08	4.30	4.00	4.00	3.96	5.50	0.90	4.92	11.32	8.32
2W	a	3.40	3.15	3.05		4.30	4.00	4.00		4.06	1.37	5.67	11.07	
	b	3.50	3.25	3.05		4.20	4.00	3.98		2.17	0.15	7.62	9.94	
	c			3.07				4.00				8.74	8.74	
3W	a	3.40	3.15	3.05		4.30	4.00	3.95		4.07	1.43	5.66	11.16	
	b	3.50	3.20	3.05		4.20	4.05	3.98		2.29	0.25	7.37	9.91	
	c			3.07				4.00				8.88	8.88	
3W	a	3.40	3.15	3.05		4.30	4.00	3.96		4.14	1.32	5.74	11.20	
	b	3.50	3.20	3.05		4.18	4.05	3.98		2.41	0.22	7.66	10.29	
	c			3.07				4.00				8.95	8.95	

studies,^{31,35,36} we decided to carry out a more detailed investigation of the local structure of Cs^+ ions in each of these regions separately.

Quasi-two-dimensional radial distribution functions (RDF) and coordination numbers (CN) for $Cs-O_w$, $Cs-O_b$, and $Cs-O_{bts}$ pairs quantifying the local coordination of Cs^+ ions separately for each of the three regions defined above were determined from monolayer to quad-layer hydrate structures and for each of our three clay models. For a chosen region, only Cs^+ ions located in this region were considered. The results obtained for our three clay models are very similar, with only small differences within the statistical errors of our calculations. Therefore, in the Supporting Information (Figures S3 and S4) we are only showing $Cs-O_b$, $Cs-O_{bts}$, and $Cs-O_w$ RDFs of the most general model *RanTO*. One should note that Cs^+ ions within the region (c) are already too far from the clay surface and only H_2O molecules are forming their first coordination shell. Data in this region are not plotted but are given in Table 2, which provides all quantitative details of the composition of the first coordination shell of Cs^+ in all three interfacial regions (a), (b), and (c) for the four hydration states considered.

The positions of the first maxima on Figures S3, Supporting Information, indicate that Cs^+ ions in the region (a) always bind more strongly to the oxygens coordinating the Al/Si substituted sites, O_{bts} ($Cs-O_{bts}$ distance is 3.12–3.18 Å not far from the $Cs-O_w$ distance slightly less than 3.1 Å), than to the oxygens coordinating nonsubstituted Si sites, O_b ($Cs-O_b$ distance is 3.39–3.43 Å). This observation holds for all hydration states studied and also for the region (b) and can be explained by the local electrostatic inhomogeneity around the substituted site: O_{bts} are slightly more negatively charged than O_b (see Figure 2), as it is reflected by the CLAYFF charge assignment.⁵⁸ Unfortunately, similar direct comparisons cannot be made with the results of other simulation studies where the $Cs-O_{clay}$ distributions were calculated without distinguishing the O_b and O_{bts} atom types. However, all previously reported $Cs-O_{clay}$ distances fall in the range between 3.3 and 3.8 Å,^{31,35,36} which is consistent with our results. Figure 10 region (c) shows that the $Cs-O_w$ distance is found in the range 3.02–3.10 Å matching well with our calculated bulk value of 3.08 Å and other calculated $Cs-O_w$ distances ranging from 3.0 to 3.21 Å in clay interlayers at different hydration states^{13,31,35,36} or in bulk aqueous solution.^{121,122} The poor statistical quality of the $Cs-O_{bts}$ radial distribution functions in region (b) is due to the relatively small number of Cs^+ ions located in this region as one can see in Figures 8b–d.

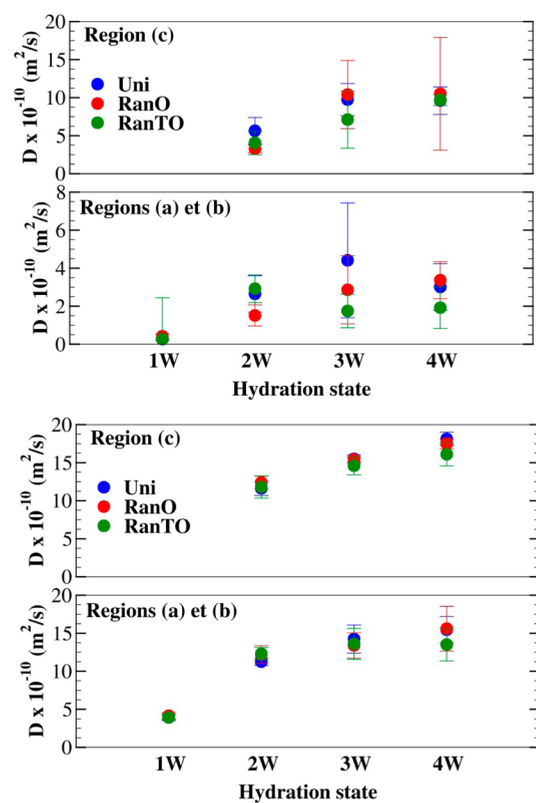


Figure 10. Three dimensional self-diffusion coefficients of Cs^+ ions (a) and water oxygens (b) in the interlayers of hydrated Cs -montmorillonite calculated for the three clay models. The self-diffusion coefficients of Cs^+ and O_{H_2O} calculated in this work for a dilute bulk solution are $(21 \pm 4) \times 10^{-10} \text{ m}^2/\text{s}$ and $(31 \pm 1) \times 10^{-10} \text{ m}^2/\text{s}$, respectively. The error bars are shown with a 95% confidence interval.

The running coordination numbers and the time-averaged 808 populations of various types of oxygen atoms in the first 809 coordination shell of Cs^+ ions in the montmorillonite 810 interlayers are shown in Table 2 (see also Figures S3 and S4 811 of the Supporting Information). The values in Table 2 were 812 determined using a cutoff value of $R_{\min} = 4.0 \text{ Å}$ for all three $Cs-$ 813 O_b , $Cs-O_{bts}$, and $Cs-O_w$ pairs. Surprisingly, even though the 814 $Cs-O_{bts}$ distances are shorter, the coordination of Cs^+ with O_b 815 atoms at the surface is much higher. This can be easily 816 explained by the fact that the Al/Si substitutions leading to the 817 presence of O_{bts} atoms are quite rare on the montmorillonite 818

819 surface. Only one out of every 60 oxygen atoms are O_{bts}
820 according to the selected clay composition. Therefore, 6 Cs^+
821 ions that can potentially adsorb on each surface (assuming that
822 the 12 interlayer ions are equally distributed to both clay
823 surfaces) have much higher chances to be coordinated by an O_{b}
824 atom rather than by an O_{bts} atom.

825 The number of O_{bts} coordinated to Cs^+ ions in region (a) is
826 equal to 0.90 for the monolayer, 1.37 for the bilayer, 1.43 for
827 the trilayer, and 1.32 for the quad-layer hydrate. In contrast,
828 there are 5.50, 4.06, 4.07, and 4.14 O_{b} atoms coordinating Cs^+
829 in the region (a), respectively, for the monolayer, bilayer,
830 trilayer, and quad-layer hydrates. These observations suggest
831 that for the monolayer hydrate ($x = 5$) Cs^+ ions are all found at
832 the hexagonal sites (H_{n} and H_{t} in Figure 9). Since the number
833 of H_{n} sites (no O_{bts} atoms in the hexagonal ring) are larger than
834 H_{t} sites (2 O_{bts} atoms in the hexagonal ring), this results in a
835 higher $\text{Cs}-O_{\text{b}}$ coordination compared to $\text{Cs}-O_{\text{bts}}$ coordina-
836 tion. As the water content increases to 2-, 3-, and 4-layer
837 hydrates, some of the Cs^+ ions are detached from the hexagonal
838 sites, and most of the remaining Cs^+ ions are located at the H_{t}
839 sites (Figure 9), so that the averaged local coordination of Cs^+
840 ions at the clay surface is made of about 4 O_{b} atoms and more
841 than 1 O_{bts} atom. The charge difference between Al (+1.575|e|)
842 and Si (+2.11|e|) structural cations in the tetrahedral sheets as
843 well as between the three O_{bts} atoms around the substituting Al
844 atoms (-1.1688|e|) and the O_{b} around the Si atoms (-1.05|e|)
845 explain the preference of Cs^+ ions for the H_{t} sites (see Figures 2
846 and 9). Even though there are 26/32 H_{n} sites and only 6/32 H_{t}
847 sites available on each clay surface for 6 Cs^+ ions, the H_{t} sites
848 are still electrostatically preferable for Cs^+ sorption.

849 Cesium ions in region (a) are coordinated to approximately 5
850 to 6 water molecules bringing the total coordination number
851 close to more than 11. This coordination number is higher by 2
852 or 3 than in a bulk aqueous solution for which we found a value
853 of 8.32 in agreement with previous findings.¹²¹⁻¹²⁴

854 In the region (b) and for the different water contents shown,
855 Cs^+ ions are almost always coordinated to 3 surface oxygens
856 with a higher contribution from O_{b} atoms (more than 2 O_{b})
857 compared to O_{bts} atoms (close to 0 O_{bts}). The contributions
858 from the two different types of surface oxygen atoms indicate
859 that Cs^+ ions found in region (b) are located mostly at T_{n} sites
860 and in some cases at $T_{\text{n}'}$ sites (28/32 and 3/32 of all such sites,
861 respectively). The T_{t} sites only marginally contribute to the Cs -
862 surface coordination within region (b) because the Cs^+ ions
863 preferably adsorbed at the neighboring H_{t} sites (see Figure 9
864 and the above discussion) would repel other Cs^+ ions and
865 prevent them from adsorbing at the nearby T_{t} sites. In addition,
866 there are only 1/32 such tetrahedral sites on each surface.
867 Therefore, the small charge delocalization around the silicon
868 atoms having two O_{b} and one O_{bts} atoms in their tetrahedra
869 ($T_{\text{n}'}$ sites), which are adjacent to the tetrahedrally substituted
870 T_{t} sites, makes the $T_{\text{n}'}$ sites favorable for the adsorption and
871 binding of Cs^+ ions in the region (b).

872 In this article we combined H_{n} and H_{o} sites (Figure 9) into
873 H_{n} sites³⁶ unlike Smith²⁷ who discussed each type of site
874 separately and found that there is no particular correlation
875 between the location of the H_{o} sites (located above a Mg/Al
876 substitution in the octahedral sheet below) and the adsorption
877 of Cs^+ ions on the clay surface. However, the preferential
878 binding to H_{t} rather than H_{n} that we observe in our simulations
879 was also observed in these two previous works.^{27,36} One should
880 also note, however, that Liu et al.³⁶ did not distinguish between
881 different binding sites as a function of their relative distance

from the surface, as proposed by Smith²⁷ and followed in this 882
work. Nevertheless, Liu et al.³⁶ have also found a very small 883
contribution from T_{t} sites compared to H_{t} sites in the 884
monolayer hydrate, indicating a strong binding preference for 885
the H_{t} sites as it was observed by Smith²⁷ for 1-, 2-, and 4- 886
layer hydrates and in our work for 1-, 2-, 3-, and 4-layer 887
hydrates. Indeed, this is due to a combination of favorable steric 888
and electrostatic reasons: the large area of the hexagonal sites 889
compared to triangular sites, the favorable 6-oxygen coordina- 890
tion compared to only 3-oxygen coordination on the triangular 891
site, the stronger repulsion from Al or Si of the triangular sites 892
compared to the hydroxyl below the hexagonal site,³⁶ and lower 893
adsorption energy on hexagonal sites compared to triangular 894
sites.¹¹⁹ No distinction was made between T_{n} and $T_{\text{n}'}$ sites in 895
the previously mentioned papers^{27,36} in which both types of 896
sites were termed T_{n} . Still they found a higher contribution 897
from T_{n} sites compared to T_{t} sites as we can now more 898
accurately quantify and explain with our findings. 899

In this region there are approximately two more H_2O 900
molecules added to the coordination sphere of Cs^+ as 901
compared to region (a) for $x = 10, 15,$ and 20 (see Table 2). 902
This indicates that the inner-sphere Cs^+ complexes formed on 903
the triangular sites need additional water molecules in the 904
hydration shell to compensate for the lacking coordination to 905
the surface oxygens, as compared to the region (a). 906

In the region (c) Cs^+ ions are only coordinated by oxygens of 907
water molecules with coordination numbers varying from 8.74 908
to 8.95 for $x = 10$ to 20 . This is still somewhat higher than the 909
coordination number of Cs^+ in bulk aqueous solutions and can 910
probably be explained by the effect of confinement in the clay 911
interlayer that may induce more densely packed H_2O molecules 912
than in the bulk. Our calculated coordination numbers agree 913
well with previous simulations.^{29,31,35} 914

3.6. Cs^+ Mobility in Montmorillonite Interlayers. The 915
diffusion coefficients of Cs^+ ions and H_2O molecules (O_{w}) were 916
determined according to eq 10 separately for the regions shown 917
in Figure 8b, and for the entire interlayer space. The regions (a) 918
and (b) were taken together since the structural analysis 919
presented above clearly indicates that Cs^+ forms only inner- 920
sphere surface complexes in both of them. For MSD 921
calculations in each of the defined regions, we considered 922
only atoms belonging to the selected region and those that do 923
not leave this region during the time period between t_0 and t 924
(see eq 10.³ 925

The 3D self-diffusion coefficients of Cs^+ ions (Figure 10a) 926
and H_2O (Figure 10b) are plotted as a function of water 927
content for the different regions defined according to Cs^+ 928
distribution in the interlayer space. For different regions, the 929
diffusion coefficients of H_2O always increase with increasing 930
interlayer water content. Adding more water molecules in the 931
interlayer increases the mobility of Cs^+ ions and H_2O molecules 932
at the clay surfaces. The higher number of water molecules 933
compared to Cs^+ ions allows us to have better statistics. This is 934
reflected in the error bars of our calculated diffusion coefficients 935
that are larger for Cs^+ ions than for H_2O molecules. 936

The mobility of Cs^+ ions in regions (a) and (b) is 2 to 4 937
times slower compared to region (c), while water diffusion 938
coefficients only slightly vary between these two regions. The 939
attractive effect of the clay surfaces is indeed more important in 940
the case of Cs^+ ions than for water molecules. The calculated 941
diffusion coefficients of water are 5 to 8 times higher than the 942
values obtained for Cs^+ ions in regions (a) and (b), and about 2 943
times higher in region (c). 944

945 Although the model *RanTO* exhibits a somewhat better
 946 statistical accuracy than the other two models, present results
 947 do not show any systematic trend explaining how the variation
 948 in substitution positions affects the mobility of the interlayer
 949 species. What can be clearly seen though is that even if the
 950 substitution location slightly affects the diffusion coefficient of
 951 Cs^+ ions, it almost does not in the case of water molecules.
 952 Table 3 compares the relative diffusion coefficients obtained
 953 from our simulations with experimental data and previous

Table 3. Relative Diffusion Coefficients of Cs^+ Ions and H_2O Molecules in the Interlayers of Montmorillonite

	data source	Cs^+	H_2O	
1W	simulations	present work	0.02–0.03	
			0.02–0.07 ³⁵	
			0.05–0.08 ³¹	
			0.006 ⁷⁰	
			0.01 ³⁸	
			0.03 ³⁶	
	experimental data		$\leq 0.008^{20,126}$	
			$\leq 4 \times 10^{-619}$	
		2W	present work	0.14–0.19
				0.18–0.25 ³⁵
3W	present work	0.53 ³¹	0.42 ³¹	
		0.03 ⁷⁰	0.18 ⁷⁰	
		0.02 ³⁸	0.42 ³⁸	
		0.31–0.44	0.48–0.52	
4W	present work	0.30–0.42 ³⁵	0.48–0.54 ³⁵	
		0.05 ⁷⁰	0.29 ⁷⁰	
		0.06 ³⁸	0.69 ³⁸	
		0.34–0.41	0.52–0.60	
		0.37 ³⁵	0.67 ³⁵	

954 simulation results. In most cases, previous simulations provided
 955 D_{xy} , the total diffusion coefficient of the species they calculated.
 956 According to eq 11, we then divided these values by the
 957 corresponding bulk self-diffusion coefficients provided or
 958 mentioned in each of the original papers. In the other cases,
 959 the relative diffusion coefficient was already provided, but given
 960 in terms of D_{xy}/D_{bulk} . Considering that D_{xy} is almost 3/2 of D_{xyz}
 961 we then divided these values by 3/2 to get the data comparable
 962 to ours. For the experimental data, we simply divided the
 963 various experimental interlayer diffusion coefficients by the bulk
 964 experimental values of Cs^+ ions⁸⁵ and water.¹²⁵

965 As discussed above, only the monolayer hydrate was
 966 investigated in experiments. There is a very large scatter of
 967 experimental values, and our relative diffusion coefficient is 2 to
 968 4 times higher than the maximum value obtained from diffusion
 969 experiments in a bentonite at different compaction densities
 970 and ionic strengths.^{20,126} One possible source of such a
 971 discrepancy is that unlike the simulations, which only look at
 972 the adsorption and diffusion in the interlayers, in real clay
 973 samples significant adsorption can happen at the clay particle
 974 edges, further retarding the diffusion of Cs^+ ions in
 975 montmorillonite. Indeed, our calculations show very good
 976 agreement with other similar simulation data, especially for the
 977 monolayer hydrate. The small differences observed increase
 978 from 2- to 4-layer hydrates, with our data being always lower
 979 than the others. Since we have already minimized the effects of
 980 the force used to describe water and ions by using the relative
 981 diffusion coefficient, the observed discrepancies between our

simulations and others may come from the presence of both
 Na^+ and Cs^+ ions in the clay interlayer and the absence of
 tetrahedral substitution in the clay composition,^{31,36,70} or the
 different force field parameters used for the clay struc-
 ture.^{31,36,38}

4. CONCLUSIONS

We have developed three new models of montmorillonite clay
 based on different degrees of disorder in the location of Al/Si
 and Mg/Al isomorphous substitutions in their tetrahedral and
 octahedral sheets. The models were thoroughly tested in MD
 simulations of Cs–montmorillonite using the CLAYFF force
 field to take advantage of its ability to model local charge
 inhomogeneities around the substituted sites and to investigate
 how various distributions of these inhomogeneities affect the
 thermodynamics of clay swelling and the structure and mobility
 of aqueous species in clay interlayers.

We have demonstrated that the specific localization of
 substitutions in the TOT layers do not affect the layer spacing
 at all, but at lower water contents up to 1-layer hydrate,
 changing the locations of octahedral substitutions in the two
 TOT layers of montmorillonite can induce a variation of
 hydration energy from 1 to 2 kJ/mol in absolute values, and
 similar changes in the tetrahedral sheets can lead to the
 variation in hydration energy from 1 to 8 kJ/mol in absolute
 values. However, the calculated values of the immersion energy
 and the isosteric heat of adsorption demonstrate much less
 sensitivity to these modifications of the local clay structure.

Detailed investigations of the structure of interlayer Cs^+
 ions allowed us to identify and probe four different binding sites
 on the basal clay surface that are identical for all the clay models
 studied. The analysis of diffusional dynamics of interlayer
 species at different distances from the clay surface demon-
 strates, as expected, that the mobility Cs^+ ions and H_2O
 molecules increases both with increasing interlayer water
 content and increasing distance from the clay surface. Only
 very small differences within the statistical errors of our
 calculations were observed between the three clay models. Our
 simulated results are in good agreement with available
 experimental data on the thermodynamics and structure of
 Cs–montmorillonite and also consistent with the results of
 previous simulations.

In summary, we can conclude that the specific localization of
 isomorphous substitutions in the structure of Cs–montmor-
 illonite has only a minor effect on the thermodynamic,
 structural, and transport properties of the system, given the
 same clay composition, the same total layer charge, and the
 same distribution of this charge between the octahedral and
 tetrahedral layers of clay. Since clay properties strongly depend
 on its composition and the nature of interlayer cations, we
 expect the effects of disorder in the distribution of the layer
 charge inhomogeneities to be much stronger in the case of the
 clays with higher total layer charge (higher concentration of
 substituted sites) and for divalent and trivalent interlayer
 cations. The MD simulations to quantify these effects are
 currently in progress and will be reported separately.

■ ASSOCIATED CONTENT

📄 Supporting Information

Additional snapshots illustrating the construction of the new
 clay models and additional results of their structural analysis.
 The atomic configurations of the three new montmorillonite

1041 models are given in separate CIF files. This material is available
1042 free of charge via the Internet at <http://pubs.acs.org>.

1043 ■ AUTHOR INFORMATION

1044 Corresponding Author

1045 *(A.G.K.) E-mail: kalinich@subatech.in2p3.fr.

1046 Notes

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