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### FORMATION AND RESTACKING OF DISORDERED SMECTITE OSMOTIC HYDRATES

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Abstract—Clay swelling, an important phenomenon in natural systems, can dramatically affect the properties of soils and sediments. Of particular interest in low-salinity, saturated systems are osmotic hydrates, forms of smectite in which the layer separation greatly exceeds the thickness of a single smectite layer due to the intercalation of water. In situ X-ray diffraction (XRD) studies have shown a strong link between ionic strength and average interlayer spacing in osmotic hydrates but also indicate the presence of structural disorder that has not been fully described. In the present study the structural state of expanded smectite in sodium chloride solutions was investigated by combining very low electron dose, highresolution cryogenic-transmission electron microscopy observations with XRD experiments. Wyoming smectite (SWy-2) was embedded in vitreous ice to evaluate clay structure in aqua. Lattice-fringe images showed that smectite equilibrated in aqueous, low-ionic-strength solutions, exists as individual smectite layers, osmotic hydrates composed of parallel layers, as well as disordered layer conformations. No evidence was found here for edge-to-sheet attractions, but significant variability in interlayer spacing was observed. Whether this variation could be explained by a dependence of the magnitude of long-range cohesive (van der Waals) forces on the number of layers in a smectite particle was investigated here. Calculations of the Hamaker constant for layer-layer interactions showed that van der Waals forces may span at least five layers plus the intervening water and confirmed that forces vary with layer number. Drying of the disordered osmotic hydrates induced re-aggregation of the smectite to form particles that exhibited coherent scattering domains. Clay disaggregation and restacking may be considered as an example of oriented attachment, with the unusual distinction that it may be cycled repeatedly by changing solution conditions. Key Words-Clay Swelling, Cryogenic Transmission Electron Microscopy, Montmorillonite, van der

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#### INTRODUCTION

Waals Forces.

Despite their widespread importance in soils, sedi-37 ments, oceans, the atmosphere, and for technological 38 applications, scientific understanding of the behavior of 39 swelling clays, such as smectite, is incomplete (Low and 40 Margheim, 1979; Laird, 2006). Clay swelling influences 41 significantly the structure and permeability of soils and 42 subsurface sediments including clay-rich shales 43 (Anderson et al., 2010). Smectite comprises a variety 44 of 2:1 layer silicates with sufficiently low charge to 45 enable them to swell, accommodating water molecules 46 and organic compounds in the interlayer region (Foster, 47 1955; Laird, 2006). At elevated temperature and very 48 49 dry conditions, all interlayer water may be removed. In 50 humid air or high-ionic-strength solutions, smectites can 51 accommodate between 1 and 3 ordered layers of water, 52 forming "crystalline hydrates" with basal spacings that 53

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vary in discrete steps from 1.1 to 1.9 nm (Dazas *et al.* 36 2015).

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In low-ionic-strength solutions, smectites can incor- 38 porate a much larger amount of water into the interlayer 39 region than would be the case for high-ionic-strength 40 solutions. In this regime, expansion of the clay is driven 41 by an osmotic process (Norrish, 1954) justifying the 42 description of the consequent structured material as 43 "osmotic hydrates" (cf. Luckham and Rossi, 1999). In a 44 landmark paper, Norrish (1954) reported that smectite 45 undergoes a discontinuous (first-order) transition from 46 the crystalline to osmotic hydrate state as solution ionic 47 strength is decreased. At concentrations of  $<\sim 0.25$  M 48 NaCl for Wyoming bentonite, the separation between 49 layers shows a linear dependence on  $N^{-\frac{1}{2}}$ , where N is the 50 salt concentration. Remarkably, this trend was observed 51 out to periodicities of ~14 nm and was subsequently 52 shown to be a function of salt type and concentration, 53 clay concentration and layer charge, and temperature 54 (Viani et al., 1983; Slade et al., 1991; Zhang et al., 55 1993; Faisandier et al., 1998; Amorim et al., 2007; 56 Svensson and Hansen, 2013). 57

In osmotic hydrates, equilibrium layer-layer separa-1 tions derive from a competition between repulsive 2 electrostatic forces and attractive van der Waals (vdW) 3 forces between the layers (Laird, 2006). Because both 4 interactions are reduced by the shielding effect of 5 interlayer ions, the interlayer spacing exhibits the 6 observed dependence on ionic strength. The same 7 8 concepts underpin the theory of colloid dispersions predicted by the theory of Derjaguin and Landau, 9 Verwey and Overbeek (DLVO), and hence many 10 treatments of this regime apply DLVO theory 11 (Greathouse et al., 1994; Liu, 2013). For smectite with 12 low structural charge (such as montmorillonite) and 13 monovalent ions, these approaches can reproduce trends 14 in interlayer spacing with ionic strength. For example, 15 Quirk and Marčelja (1997) showed DLVO theory to 16 correctly predict the interlayer separation of Li-mont-17 morillonite from 1.8 to 12 nm as a function of pressure 18 and LiCl concentration. DLVO failed, however, to 19 predict accurately the ionic-strength threshold for 20 aggregation (Missana and Adell, 2000) and swelling 21 pressure (Viani et al., 1983). Moreover, all models based 22 on DLVO theory fail to depict accurately behavior for 23 highly charged surfaces or non-monovalent ions such as 24  $Ca^{2+}$  (Kleijn and Oster, 1982). 25

All XRD and small-angle X-ray scattering (SAXS) 26 studies of smectite osmotic hydrates show evidence of 27 28 disorder in the interlayer spacings (Quirk and Marčelja, 1997; Faisandier et al., 1998). Whether this is caused by 29 intrinsic heterogeneity of natural smectites, with cation 30 substitutions leading to local variations in layer charge or 31 whether it is simply a consequence of natural fluctuations 32 due to, e.g. Brownian motion, is not known. Molecular 33 simulations have predicted that alternative stacking 34 configurations, such as edge-to-face, could prevail under 35 certain conditions (Jönsson et al., 2008). Tests of such 36 predictions are currently lacking, however. 37

The present authors suggest that osmotic hydrates 38 contain structural disorder that has not been fully 39 characterized by any prior study and that a better 40 understanding of this disorder would advance knowledge 41 of the forces between clay layers. Conventional methods 42 are unable to reveal detailed information about hydrated 43 smectite structure. Despite recent advances in the 44 analysis of X-ray scattering data, this approach has 45 limited the ability to characterize disorder in suspensions 46 of anisotropic crystallites. Accurately quantifying the 47 48 distributions of different layer hydration states in crystalline hydrates is possible from XRD data (Pons 49 et al., 1981; Ferrage et al., 2005; Dazas et al., 2015), but 50 not the distributions of spacings in osmotic hydrates. 51 Even fewer attempts have been made to quantify 52 disorder in osmotic hydrates. Several groups have used 53 SAXS to explore the average distances between layers in 54 completely exfoliated smectite as a function of solution 55 conditions (e.g. Paineau et al., 2011). Interpretation of 56 SAXS data beyond quantification of mean interlayer 57

spacings is very model dependent, however, and direct 1 observations of layer disorder, useful for validating 2 proposed models, have not been reported to date. 3

Only transmission electron microscopy (TEM) has 4 the necessary spatial resolution for imaging smectite 5 particles. Guthrie and Veblen (1989) showed that it was 6 possible to measure coherent stacking sequences in 7 mixed-layer illite-smectite. The ultrahigh vacuum conditions required for TEM can cause complete loss of 9 interlayer water, however. Many groups have studied 10 smectites that were expanded by alkylammonium intercalation and preserved in epoxy (*e.g.* Schumann *et al.*, 12 2014) but epoxy intrusion into the interlayers can alter interlayer spacing relative to values observed by XRD 14 for hydrated systems. These conventional TEM methods 15 are even less suitable for studying osmotic hydrates. 16

Here a first imaging study of smectite in the osmotic 17 hydrate state using cryogenic transmission electron 18 microscopy (cryo-TEM) is reported. Rapid vitrification 19 of samples enabled electron microscopy imaging of 20 particles in aqueous suspensions. The flash freezing 21 approach has been proven to preserve the structure of 22 hydrated biological macromolecules such as ribosomes 23 for which independent atomic-resolution structures exist 24 (see Cheng, 2015, and references therein). Cryo-TEM 25 has previously been used to determine the structure of 26 microbial mineral precipitates (Comolli et al., 2011) and 27 the structure of fragile, extended aggregates of Fe 28 oxyhydroxide nanoparticles (Yuwono et al., 2010; 29 Frandsen et al., 2014; Legg et al., 2014). Cryo-TEM 30 was used by Segad et al. (2012) to compare the 31 aggregate structures of Na- and Ca-equilibrated mont- 32 morillonite suspensions, but without investigating the 33 osmotic hydrate regime. 34

In the current study a state-of-the-art low-dose cryo- 35 TEM was used to characterize osmotic hydrates formed 36 by a montmorillonite from a Wyoming bentonite 37 (SWy-2). The bentonite was either studied without 38 treatment, or was treated with standard methods to 39 obtain a purified, Na-equilibrated montmorillonite frac- 40 tion of sub-2 µm particles. Image details led the authors 41 to speculate that variations in interlayer spacings arise 42 neither from intrinsic layer charge heterogeneity nor 43 from random Brownian motion, but rather from the 44 action of vdW forces which extend over multiple 45 interlayer distances. This length scale is not typically 46 considered in theories of smectite swelling. Because 47 configurational disorder prevented direct validation of 48 this concept in the present study, the idea was tested 49 using a recently developed computational method for 50 prediction of vdW forces in multi-layer geometries. 51 Despite lack of the materials-properties data required for 52 accurate predictions for layered aluminosilicates in high- 53 salinity solutions, calculations in the present study 54 suggest strongly that equilibrium interlayer spacings in 55 osmotic hydrates are influenced by layer-layer vdW 56 forces that extend across multiple smectite layers. 57

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#### Experimental methods

Clay treatments. The experiments used samples of the 4 Source Clay SWy-2 from the Source Clays Repository of 5 The Clay Minerals Society (http://www.clays.org/ 6 SOURCE%20CLAYS/SCBackground.html). SWy-2 is a 7 Na-rich bentonite from Crook County, Wyoming, in 8 9 which the smectite fraction is montmorillonite. A multistep treatment of this material was performed to 10 create a homoionic Na-saturated suspension free of 11 mineral impurities referred to as Na-montmorillonite. 12 The steps are described in more detail in the 13 Supplementary Information section (deposited with the 14 Editor in Chief and available at http://www.clays.org/ 15 JOURNAL/JournalDeposits.html), and included (1) the 16 removal of calcite impurities using a 1 M sodium 17 acetate/0.564 M glacial acetic acid solution, (2) the 18 equilibration of the clay with a 1 M sodium chloride 19 solution, (3) the removal of excess Na-salts with 20 Nanopure water, (4) the separation of quartz and 21 feldspar impurities from clay particles by centrifugation 22 while retaining the  $<2 \mu m$  fraction, and (5) oven-drying 23 of the purified clay mineral phase at 45°C (~2 weeks). 24 Steps 1-3 were performed by dialysis. The dried clay 25 was recovered and ball milled for 2 min and stored dry 26 until its resuspension in NaCl solutions. 27

MATERIALS AND METHODS

28 Dried smectite was pressed into powder pellets, 29 carbon coated, and analyzed by electron probe microanalysis (EPMA) with a Cameca SX-51 in the 30 31 Department of Earth and Planetary Science at the University of California, Berkeley, USA. Data acquisi-32 tion, analysis, and correction procedures were conducted 33 with the software, Probe, for EPMA, as described in the 34 Supplementary Information section. 35

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Cryo-TEM specimen preparation and instrumentation. 37 As-received or treated samples of dried and powdered 38 Wyoming montmorillonite were suspended in water or 39 NaCl solutions at salt concentrations below the crystal-40 line-to-osmotic transition threshold (~0.25 M). Samples 41 were equilibrated overnight. Aliquots of 5 µL were 42 taken directly from the suspensions, placed onto 43 200 mesh lacey carbon formvar or 300 mesh lacey 44 carbon Cu-grids (Ted Pella, Incorporated, California, 45 USA, #01881 and #01895, respectively), manually 46 blotted with filter paper and plunged into liquid ethane 47 48 at liquid nitrogen temperature. The TEM grids had 49 previously been glow discharged.

Images were acquired on a JEOL-3100-FFC electron 50 51 microscope equipped with a field emission gun (FEG) electron source operating at 300 kV, an Omega energy 52 filter, and a cryo-transfer stage. Images were recorded on 53 a Gatan 795 4k × 4k charge-coupled device (CCD) camera 54 (Gatan Inc., Pleasanton, California, USA) mounted at the 55 exit of an electron decelerator operated at 200 kV, 56 resulting in images formed by a 100 kV electron beam 57

at the CCD. The stage was cooled to 80 K with liquid 1 nitrogen during the acquisition of all data sets. 2

Cryo-TEM image acquisition and analysis. Over 100 4 two-dimensional (2D) images were recorded at nominal 5 magnifications of 50 k  $\times$ , 80 k  $\times$ , 120 k  $\times$ , and 200 k  $\times$  6 giving pixel sizes of 0.112 nm, 0.07 nm, 0.047 nm, and 7 0.028 nm at the specimen, respectively. Underfocus 8 values ranged between  $\sim 1000 \pm 100$  nm and 35  $\pm$  5 nm 9 (measured from minimum contrast), and energy filter 10 widths were typically ~30 eV. The survey of the grids 11 and the selection of suitable targets were done in low- 12 dose defocused diffraction mode to minimize radiation 13 damage, as established for biological samples (Luef et 14 al., 2012). Images were acquired under low-dose 15 conditions, typically with doses of ~125-250, 320, 16 720, and 1000  $e^{-}$ Å<sup>-2</sup> per image for nominal magnifica- 17 tions of 50 k×, 80 k×, 120 k×, and 200 k×. The 18 software ImageJ 1.38x, National Institutes of Health 19 (NIH) http://rsb.info.nih.gov/ij/) was used for analysis 20 and measurements of the 2D image projections. 21

X-ray diffraction. Powdered samples of treated SWy-2 23 were suspended in NaCl solutions or ultrapure water at a 24 smectite concentration of 200 µg/mL for 1 week. A 25 small aliquot of the suspensions was placed in 300-µm 26 internal-diameter quartz capillaries for synchrotron XRD 27 analysis at beamline 12.3.2 at the Advanced Light 28 Source. For improved powder statistics, samples within 29 the capillary were scanned over a 100  $\mu$ m  $\times$  100  $\mu$ m 30 (5  $\mu$ m steps) area through a 2 × 10  $\mu$ m 10 keV beam, 31 equivalent to a wavelength of 1.2398 Å. Powder patterns 32 were recorded in 600-s exposures using a Pilatus-1M 33 area detector (Dectris, Baden-Dättwil, Switzerland) 34 positioned at  $20^{\circ}2\theta$  and ~190 mm distance. The experi- 35 mental geometry was calibrated using Al<sub>2</sub>O<sub>3</sub> National 36 Institutes of Standards and Technology (NIST) standard 37 material (SRM 676a). Initial suspensions were too dilute 38 to provide either interlayer or in-plane diffraction peaks, 39 so the capillaries were spun at 10 krpm for 2 h in a 40 special insert machined to fit an Eppendorf centrifuge. 41 Centrifugation caused a visible increase of clay density 42 in the salt suspension and permitted analysis by XRD. A 43 portion of the same sample was dried on a filter, 44 removed gently and re-powdered, and placed into an 45 identical capillary. The low-angle  $(1.8-8^{\circ}2\theta)$  data were 46 processed by fitting and removing an exponential 47 function corresponding to scattering from water. The 48 wide-angle  $(5-35^{\circ}2\theta)$  data were processed by subtract-49 ing the scattering from an empty capillary. 50

#### Calculations

Van der Waals force calculation. The freely available 53 Gecko Hamaker code (Rajter *et al.* 2013) was used for 54 simulating the vdW interactions within multi-layered 55 models of stacked smectite in water. The Gecko code 56 calculates the Hamaker constant  $A_{123}$  for the interaction 57

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between two structured objects 1 and 3 separated by 1 medium 2 of thickness D. The approach was described 2 fully by Podgornik and Parsegian (2004) and Parsegian 3 (2006) and is based upon the theory of Lifshitz (1956) 4 that links the complex optical properties of two solids to 5 the dispersion forces between them. The vdW force per 6 unit area,  $P_{vdW}$ , is then calculated from: 7

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$$p_{\rm vdW} = \frac{A}{6\pi D^3}$$

A series of calculations was designed to test the 11 hypothesis that long-range vdW interactions (beyond 12 next nearest neighbor layers) made a significant 13 contribution to the overall cohesion energy of a smectite 14 particle with a uniform interlayer spacing that was fixed 15 at either 4 or 5 nm. Given the lack of optical-property 16 measurements for layer silicates, a silica-corundum-17 silica stack was used to approximate one individual 2:1 18 layer, using material constants for corundum  $(Al_2O_3)$ 19 and amorphous silica (SiO<sub>2</sub>) from the Gecko database 20 (French et al. 1998; Tan et al. 2005). A full calculation 21 22 of clay particle cohesion energy as a function of all relevant parameters will require new dielectric measure-23 24 ments for layer silicates and for salt solutions.

The Hamaker constant for inner- and outermost pairs 25 of clay layers was calculated as a function of the total 26 number of layers in a clay particle, from 2 to 14. For 27 28 these calculations the intervening solution was pure 29 water because the dielectric functions for electrolyte 30 solutions are not known accurately. A portion of the 31 vdW forces are effectively screened by electrolyte 32 solutions, but the most significant part involves electronic fluctuations which occur on a timescale that is far too 33 short to be effectively screened by the motion of ions in 34 solution (Davies and Ninham, 1972). No previous study 35 of ionic-strength effects on vdW interactions between 36 minerals could be found by the present authors. 37 However, Marra (1985) showed that the strength of 38 vdW interactions between digalactosyl-diglyceride 39 bilayers decreased by ~50% in 0.2 M NaCl compared 40 to water. Therefore, the 0.1 M solution used in the 41 present study was assumed to reduce the vdW strength 42 by half relative to that predicted for pure water. This 43 correction factor was not applied to the calculations. 44

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Electrostatic force calculation. For comparison with the 46 magnitude of the vdW interaction forces, the electro-47 static repulsion pressure between two model 'mont-48 49 morillonite' layers in 0.1 M NaCl was calculated. The 50 pressure,  $P_{\rm EDL}$ , was calculated from the force between overlapping electrical double layers (EDL): 51

$$P_{\text{EDL}} = \frac{1}{2} \varepsilon_r \varepsilon_0 \kappa^2 \Phi_0^2 \left(\frac{1}{\cosh\frac{\kappa D}{2}}\right)^2$$

55 where  $\kappa = (\varepsilon_r \varepsilon_0 R T/2z^2 F^2 c)^{\frac{1}{2}}$  is the inverse Debye length, 56  $\varepsilon_r \varepsilon_0$  is the dielectric permittivity of water, R is the ideal 57

gas constant, T = 298 K is temperature, z = 1 is the ionic 1 valence,  $\Phi$  is Faraday's constant, and c is the bulk 2 electrolyte concentration (in ions/cm<sup>3</sup>). The surface 3 potential,  $\Phi_0$ , was fixed at 250 mV (Miller and Low, 4 1990; Hou et al. 2009). 5

#### RESULTS

Both intact and treated SWy-2 formed visually clear 9 suspensions readily after equilibration in 0.1 M NaCl 10 solution at smectite concentrations of 50 µg/mL or 11 lower. Representative low-magnification defocused dif- 12 fraction imaging of cryo-TEM preparations are shown in 13 Figures S1-S3 (deposited with the Editor in Chief and 14 available at http://www.clays.org/JOURNAL/ 15 JournalDeposits.html). The suspensions contained a 16 dispersion of individual layers consisting of one 17 octahedral and two tetrahedral sheets (2:1 layers), the 18 fundamental structural unit of smectite, and particles 19 containing multiple stacked 2:1 layers (smectite parti- 20 cles). Higher-magnification lattice-fringe images cap- 21 tured individual layers, smectite particles, and osmotic 22 hydrates that lay within the depth of focus in the vitreous 23 ice sample (estimated to be >100 nm for 80 k  $\times$ 24 magnification). These images revealed that the majority 25 of smectite in both the untreated and treated clays had 26 expanded beyond all crystalline hydrate states and 27 formed osmotic hydrates, as expected for the low NaCl 28 concentrations. All images revealed conformational 29 disorder including heterogeneity in smectite layer 30 spacings. 31

An image recorded from a suspension of untreated 32 SWy-2 illustrates the tendency of the expanded smectite 33 to form oriented, parallel stacking configurations in the 34 osmotic hydrate state (Figure 1a). In other cases, despite 35 the formation of pairs or triplets of parallel smectite 36 layers, these units were rotated with respect to each 37 other within the larger ensemble (Figure 1b). Partial 38 osmotic hydrates that also include regions of crystalline 39 hydrates with interlayer spacings consistent with both 40 two- and three-water layers are shown in Figure 2. 41 Completely delaminated individual clay layers with little 42 evident curvature were also observed (Figure S4) as 43 were highly disordered aggregates of layers (Figure S5). 44

The treated SWy-2 sample delaminated more com- 45 pletely than the untreated sample following equilibration 46 with 0.1 M NaCl solutions. More individual layers were 47 observed and no examples of crystalline hydrates were 48 found. As shown in Figure 3, the expanded smectite was 49 frequently observed as parallel, stacked layers, just as 50 observed for untreated SWy-2 suspended in 0.1 M NaCl. 51 Fully delaminated layers, either with evident curvature 52 or with a highly linear morphology, were observed 53 (Figure S6). Also observed were disordered osmotic 54 hydrates (Figure S7A) and parallel assemblies rotated 55 relative to one another (Figure S7B), as seen for the 56 untreated sample. 57

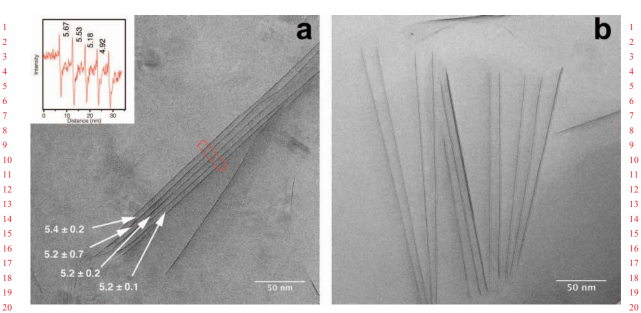


Figure 1. Cryo-TEM images of untreated SWy-2 smectite equilibrated in 0.1 M NaCl at 50 µg/mL (a). This is an example of a more ordered stacking configuration in the osmotic hydrate state showing the average interlayer spacing and the standard deviation (nm) from five measurements along the osmotic hydrate (cf. Figure S9). Inset: example intensity profile from one indicated area used to determine interlayer spacing (nm). (b) Example of a partially ordered stacking configuration. The arrow indicates a pair of layers in a crystalline hydrate configuration. 

As shown by the EMPA data (Table S1, deposited http://www.clays.org/JOURNAL/JournalDeposits.html), 28 with the Editor in Chief and available at the long dialysis period decreased slightly the Si:Al 29

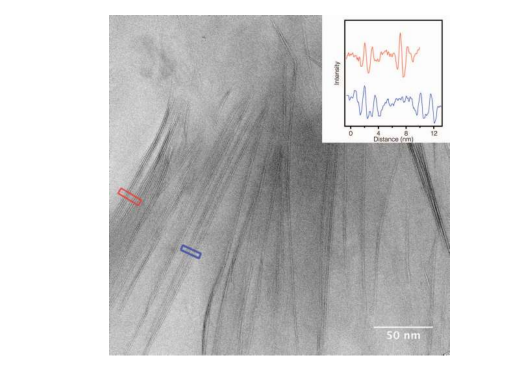


Figure 2. Cryo-TEM image of untreated SWy-2 smectite equilibrated in 0.1 M NaCl at 50 µg/mL. The image shows evidence of a partially expanded osmotic hydrate that incorporates 2- and 3-water crystalline hydrates. Inset: intensity profiles from the areas indicated reporting the interlayer spacing (nm). 

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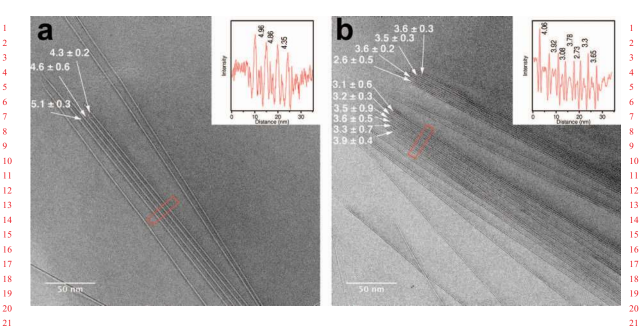


Figure 3. Cryo-TEM images of treated SWy-2 smectite equilibrated in 0.1 M NaCl at 20% w/v. Also given are the average interlayer spacing and the standard deviation (nm) from five or eight measurements along the osmotic hydrate. Insets: example intensity profiles from the regions indicated used to estimate interlayer spacing (nm).

ratio. This could be an indication of smectite dissolution 26 and the release of Al<sup>3+</sup> ions that are known to be able to 27 displace Na<sup>+</sup> from the interlayer regions. Extensive 28 29 release of trivalent Al ions would be expected to collapse the smectite, however, which was not observed. 30 31 For smectite particles that showed the expected 32 osmotic hydrate structure, the interlayer spacings were measured. Because the smectite layers are oriented at an 33 unknown tilt with respect to the image plane, such 34 measurements could overestimate the interlayer spacing. 35 For example, the TEM asymmetric intensity profile 36 shown in Figure 1a probably indicates tilt of the layers. 37 As shown in Figure S8, however, analysis of a tilt series 38 showed that a lattice-fringe imaging condition exists only 39 for tilt angles within  $\pm 15^{\circ}$  or less of the beam direction. 40 Thus, the measured spacings could be up to 5% too large. 41 The mean interlayer spacings from ordered osmotic 42 hydrates are shown in Figures 1-3 as well as are the 43 standard deviation determined from 5-8 measurements 44 at ~50-nm intervals (Figure S9). A significant range in 45 interlayer spacings was observed, from 2.6 to 5.4 nm. 46 Although the sample size is very small, the different clay 47 particles in the same sample exhibited different average 48 49 spacings. One plausible hypothesis is that the stoichiometric composition of the clay (and hence layer charge) 50 51 was locally homogeneous but varied throughout the 52 sample. The observations suggested an alternative hypothesis, however: the mean interlayer spacing could 53 be a function of the number of layers in the particle if 54 vdW interactions extend beyond a single interlayer 55 distance. Thus, a series of numerical calculations, 56 designed to test the idea that long-range vdW interac-57

tions could significantly alter the cohesion energy of a 26 stack of 2:1 layers, was performed. 27

The calculations of the vdW interactions between 28 pairs of model 2:1 layers in an osmotic hydrate (Figure 4 29 inset) assumed pure water in the interlayer space because 30 accurate dielectric functions for NaCl solutions were not 31 available. As discussed above, the NaCl probably 32 reduces the strength of the vdW interactions by  $\sim 50\%$  33 relative to pure water (Marra, 1985). Because no 34 accurate measurement of vdW shielding for saline 35 solutions currently exists, however, no correction factor 36 was applied to the calculations reported in Figure 4. 37 Although the effect of salinity on the absolute magnitude 38 or range of the vdW interaction forces is not known 39 precisely, this uncertainty does not invalidate the 40 calculation, which aimed to test the dependence of 41 vdW interactions on layer number. 42

The calculations of layer-layer attraction,  $P_{\rm vdW}$ , 43 summarized in Figure 4, clearly predict that the vdW 44 interaction forces between a pair of model clay layers in 45 a particle are affected by the presence of additional 46 layers. For pure water and a fixed spacing of 4 nm, the 47 results predict that long-range interactions increase the 48 total cohesion energy of the system by >10%. Beyond 49 ~5 layers, vdW interactions contribute little and, thus, 50 can be ignored. An estimate of the electrostatic repulsion 51 pressure,  $P_{\rm EDL}$ , for a pair of smectite layers in 0.1 M 52 NaCl, separated by 4 nm, is shown in Figure 4. At 53 equilibrium,  $P_{\rm vdW}$  and  $P_{\rm EDL}$  should be equal and 54 opposite. This equality is approximately satisfied if the 55 0.1 M NaCl shields half of the vdW contribution as 56 indicated by the results of Marra (1985). 57

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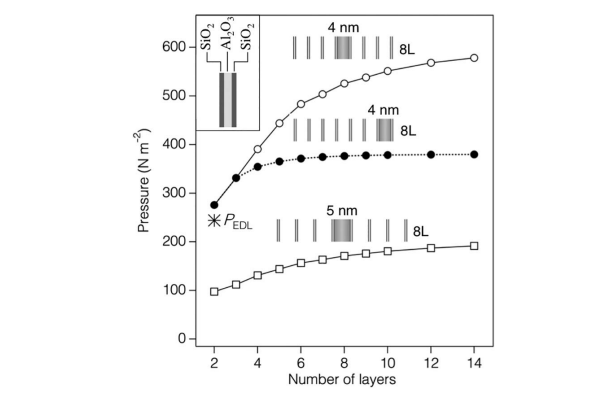


Figure 4. The calculated van der Waals force per unit area between one pair of smectite layers in water as a function of the total number of layers in a stack. A single 2:1 layer (composed of tetrahedral-octahedral-tetrahedral sheets), was represented by a silica-corundum-silica multilayer. (inset). Calculations were performed for the centermost (hollow symbols) and outermost (solid symbols) pair of layers and for interlayer spacings of 4 nm (circles) and 5 nm (squares). Illustrations of the geometry are depicted for 8-layer (8L) stacks, with the selected interlayer shaded. Also shown is the electrostatic repulsion pressure ( $P_{EDL}$ ) calculated for a single pair of 2:1 layers with a surface potential of 250 mV at a separation of 4 nm. 

The XRD data acquired from a suspension of treated SWy-2 in 0.1 M NaCl in a capillary and from a portion of the same suspension under filtration are compared in Figure 5. Under fully hydrated conditions, interlayer diffraction peaks with spacings up to 4 nm were not detectable (see figure inset). Thus, although clay particles with spacings in this range were observed at the cryo-TEM, the observed disorder broadened and weakened the scattering. Following filtration and drying, however, the clay exhibited strong (001) and (004) spacings, consistent with one water layer crystalline hydrate containing Na (Wilson et al., 2004). 

#### DISCUSSION

#### 48 Smectite structure in aqueous solution

The cryo-TEM observations of lattice-fringe spacings for the hydrated smectite structure are in fair agreement with previous XRD data, supporting the present conclu-sion that the flash-freezing approach did not introduce artifacts. Specifically, the measured interlayer spacings, which range from 2.6 to 5.4 nm, are close to the values reported for hydrated Na-montmorillonite by Zhang et al. (1995) but are lower than values reported by Norrish (1954) and Amorim et al. (2007). The observation of 

expanded smectite with a range of interlayer spacings is 33 in agreement with prior inferences based on XRD (Pons 34 *et al.*, 1981; Amorim *et al.*, 2007). The imaging results 35 add considerable new insight into the structure and 36 disorder in smectite osmotic hydrates, however. 37

Montmorillonite possesses a fixed negative charge in 38 the octahedral sheet and reactive oxygen sites at layer 39 edges that are expected to be protonated and positively 40 charged in circumneutral solutions (Bailey, 1988; 41 Tournassat *et al.*, 2004). These features have motivated 42 predictions of 'house-of-cards' or 'overlapping-coins' 43 structures (van Olphen, 1977; Jönsson *et al.*, 2008; 44 Delhorme *et al.*, 2011). Despite the considerable disorder, 45 no evidence was observed for such arrangements. 46

In the present study, smectite 2:1 layers, either in free 47 solution or stacked in particles, typically exhibited less 48 overall curvature (on the 100-nm and longer scale) than 49 observed for expanded smectite imaged in epoxy (*e.g.* 50 Schumann *et al.*, 2014). Only one image provided 51 evidence of curvature in isolated layers (Figure S6). 52 Flash freezing is expected to introduce fewer structural 53 artifacts than epoxy-based sample preparations; the 54 present observations are assumed, therefor, to depict 55 accurately the real planarity of hydrated smectite layers. 56 Close inspection of the images, as well as quantitative 57

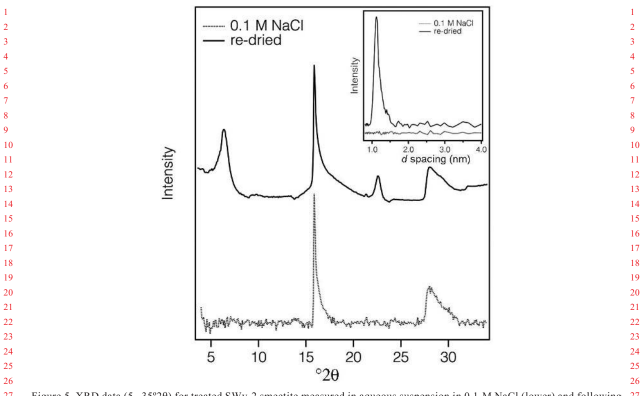
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27Figure 5. XRD data (5–35°2θ) for treated SWy-2 smectite measured in aqueous suspension in 0.1 M NaCl (lower) and following2728drying (upper). Inset: low-angle scattering regions (1.8–8°2θ) plotted vs. d spacing. X-ray wavelength was 1.2389 Å.28

measurements of the parallelism of the ordered stacking
arrangements (Figure S8), reveal fine-scale undulations
in most layers. These undulations could be a consequence of metal substitutions within the layers, but
higher-resolution studies will be required to explore this
possibility fully.

In many images, one or more layers were observed 36 close to, but not parallel to, layers in an ordered osmotic 37 hydrate particle. Brownian motion is suggested here to 38 result in osmotic hydrates that are in a state of dynamic 39 40 formation and disassembly. This cannot be tested by cryo-TEM, which captures snapshots of instantaneous struc-41 ture, or by conventional SAXS methods which provide 42 measurements averaged over many particles and over a 43 significant acquisition time. Fluid-cell TEM might allow 44 the dynamic structure of osmotic hydrates to be observed. 45 The present authors predict that the structural configura-46 tions observed by cryo-TEM will enable new models for 47 48 scattering from disordered lamellar systems to be devel-49 oped that will allow SAXS analysis of bulk solutions to 50 provide richer and statistically more significant analysis 51 of smectite suspensions.

## Long-range attractive interactions spanning multiple smectite layers

The present calculations predict that vdW interactions spanning multiple 2:1 and water layers is a nonnegligible contribution to the cohesion of expanded smectite particles. The uncertainties in the magnitudes of 30 the vdW forces described above do not affect this 31 important calculation-based prediction. Long-range vdW 32 interactions must be important in osmotic hydrates 33 because structures with large (i.e. >10 nm) layer 34 distances are stabilized. To the best of the authors' 35 knowledge, however, no prior experimental or theore- 36 tical study of osmotic hydrates has considered the role of 37 long-range, next-nearest-neighbor vdW interactions. 38 This prediction may be important because it implies 39 that the overall cohesion energy of expanded clay 40 particles is a function of the number of layers in the 41 particle and that the interlayer spacing will be smaller 42 for particles with a larger number of layers. Thus, 43 models of clay swelling based on a single pair of layers 44 (e.g. Liu, 2013), or considering only next-layer interac- 45 tions (e.g. Kleijn & Oster, 1982), will underestimate the 46 layer-normalized cohesion energy of larger particles, 47 and overestimate the layer distances. 48

#### Layer re-association and constraints for attachmentbased crystal growth

The XRD data in Figure 5 show clearly that 52 disordered, expanded clay layers, which exhibit no 53 detectable 00l scattering, re-aggregate readily when the 54 water activity is reduced by drying to form ordered 55 crystalline hydrates with strong diffraction perpendicular 56 to the layer plane. Because transformation of osmotic to 57

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crystalline hydrates involves assembly of individual 1 smectite 2:1 layers in solution, it can be classified as an 2 example of oriented attachment-based crystal growth 3 (OA) (Penn and Banfield, 1998). Such OA has been 4 reported as a crystal-growth mechanism for many poorly 5 soluble nanoparticulate minerals, including oxides and 6 sulfides (Penn et al., 2001; Huang et al., 2003), and has 7 been observed in suspensions of nanoparticles with a 8 9 range of morphologies (Yuwono et al., 2010; Frandsen et al., 2014). The case of assembly of 2:1 smectite layers 10 11 is distinct from other growth processes described as OA 12 because the attachment process is readily reversible due to the unusually low strength of layer-layer interactions 13 (because attachment does not involve the formation of 14 covalent or ionic bonds). The authors here do not mean 15 to imply that smectite particles form originally through 16 the stacking of individual 2:1 layers that nucleate 17 homogeneously in solution (although this sometimes 18 may be the case). Rather, the 2:1 layer assembly is 19 20 described as one of the steps in the dynamic process of swelling, disaggregation, and re-aggregation that prob-21 ably occurs innumerable times over the 'lifetime' of 22 smectite particles. 23

24 In research on OA-based crystal growth, the nature of nanoparticle interactions in aqueous solution and the 25 distances over which they interact are uncertain. In situ 26 fluid-cell TEM observations of OA-based crystal growth 27 28 of ferrihydrite indicate the operation of long-range particle-particle interactions (Li et al., 2012). Studies 29 30 of smectite swelling and restacking can provide insights 31 into interparticle forces involved in OA-based crystal 32 growth because the two-dimensional geometry enables interactions that are parallel and normal to the 2:1 layers 33 to be resolved. Observations of attractive forces in the 34 3 to >20 nm range in expanded clays indicate that 35 particle-particle interactions can occur over such 36 distances (although nanoparticle size and morphology 37 will affect interaction distances). 38

In OA-based growth, an orienting force is suspected 39 40 to align nanoparticles as they approach so that interface elimination can yield a single crystal (Li et al., 2012; 41 Zhang and Banfield, 2014). Cryo-TEM images show that 42 separated smectite layers in ordered particles are parallel 43 to each other. This geometry is explained conceptually 44 by DLVO theory in which the balance between attractive 45 and repulsive forces orients the surfaces. The cryo-TEM 46 data also show that layers tend to overlap each other. 47 48 This implies that lateral forces parallel to the 2:1 layers 49 in the osmotic hydrate are sufficiently strong to drive the 50 system to an energy minimum in which the layers 51 superimpose. Attractive layer-layer vdW forces could 52 provide the driving force for this alignment, although electrostatic forces must play a role once the layers are 53 54 close enough that the electrical double layers overlap. Cryo-TEM observations of smectite in aqua, thus, 55 provide insight into the nature of layer-layer interac-56 tions controlling clay expansion and restacking and can 57

inform an understanding of related colloidal phenomena 1 including OA-based crystal growth. 2

#### CONCLUSIONS

Cryo-TEM observations of flash-frozen suspensions 6 of smectite in 0.1 M NaCl solution revealed that osmotic 7 hydrates, composed of highly planar and close to parallel 8 2:1 layers, coexisted with 2:1 layers in a range of 9 disordered conformations, but without evidence of 10 alternative edge-to-sheet interactions. The large range 11 of interlayer spacings measured in the osmotic hydrates 12 suggested a role for long-range attractive van der Waals 13 interactions in determining equilibrium structure. 14 Calculations of van der Waals forces in a multi-layer 15 model of expanded smectite predicted that attractive 16 forces acting over at least four smectite interlayers will 17 influence osmotic hydrate cohesion energy. Due to the 18 disorder in the expanded state, (001) diffraction peaks 19 were not detectable in XRD data, but were observed 20 following restacking (through drying). Assembly of 21 individual smectite 2:1 layers in solution can be 22 classified as an example of oriented attachment-based 23 (OA) crystal growth. Further elucidation of the forces 24 that act to align smectite particles during stacking, aided 25 by cryo-TEM observations, will improve understanding 26 of both clay swelling and OA based growth. 27

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