Article

Comparison of microstructural features of three compacted and water-saturated swelling clays: MX-80 bentonite and Na- and Ca-purified bentonite

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Abstract

The planned final disposal repositories of spent nuclear fuel in several countries, including Finland, pose significant scientific challenges due to their extremely long lifespan. One of the key materials proposed for use in Posiva Oy’s repository in Finland is MX-80 bentonite in a compacted, water-saturated state. Border cases of calcium and sodium forms of purified bentonite were included in this study. The MX-80 in the repository is expected to undergo cation exchange due to the composition of the groundwater. The clays were studied at different dry densities between 0.7 and 1.6 g cm$^{-3}$. The microstructure of the water-saturated, compacted clays was investigated using small-angle X-ray scattering, nuclear magnetic resonance and transmission electron microscopy. Additionally, atomic force microscopy was used to characterize the shape and size of the fine-fraction clay platelets. As expected, the average shape of the fine fractions was smaller than the bulk material, but a more elongated shape was present in the purified material. Mainly due to sample density, the pore structure was noticeably different for the Na form of purified bentonite at 0.7 g cm$^{-3}$ density, but at higher degrees of compaction, no significant differences were noted between the samples. The laboratory results obtained in this study could be useful for safety and performance analysis of this high-level waste repository where sodium bentonite is used and is expected to change its charge-compensating cation composition during the repository’s lifetime. Microstructural data may be used in modelling of diffusion and sorption by surface complexation modelling, for example, or as a basis for mechanical and water transport models.

**Keywords:** MX-80, bentonite, montmorillonite, porosity, microstructure, HLRW repository, swelling clay

MX-80 bentonite is proposed for use in high-level radioactive waste (HLRW) repositories in several countries. The clay buffer in the KBS-3 method applied in Finland and Sweden (SKB, 2011; Posiva, 2012) is placed between a copper canister containing the waste and the granitic host rock. Therefore, among other functions, the buffer material should have very low hydraulic conductivity in order to limit the mass transport to diffusion only, but it also should be ductile enough to dampen the rock movements (Juvankoski, 2013). As the lifespan of the repository will be extremely long, the estimations of its performance are based on computer modelling. Currently, no consistent models explaining all of the macroscopic properties of bentonite exist (Savage, 2012). Therefore, a better understanding of the material is needed in order to form a solid scientific basis for the models used in the safety assessment of HLRW repositories.

One of the key factors influencing bentonite behaviour is its microstructure. Numerous articles presenting the microstructure of swelling clays exist in the literature, covering a variety of approaches and techniques. In various cases, either commercial bentonites or purified, homo-ionic materials were analysed. Investigation of the size and shape of individual montmorillonite platelets by means of atomic force microscopy (AFM) is usually done on Na- or Ca-montmorillonite (Cadène et al., 2005; Ploehn & Liu, 2006; Cao et al., 2010; Pettersson et al., 2010) or on purified material if various bentonites are compared (Delaverne et al., 2015). Electron microscopy is also used to visualize clay systems. Scanning electron microscopy (SEM) may be used to characterize the microstructural features of clays in terms of being either humid, dried, frozen or embedded in a polymer matrix (Montes-Hernandez, 2005; Katti & Katti, 2006; Cuisinier et al., 2014; Kaufhold et al., 2016). Similarly, transmission electron microscopy (TEM) samples are either embedded or frozen (Pusch, 1999; Carlsson et al., 2012; Segad et al., 2012). The methods using electron microscopy and ion milling are undergoing intense development. Combining advanced sample preparation (freezing the porewater to vitreous ice and embedding it in polymer resins) and precise 3D imaging of the samples...
means such methods give excellent insight into the clay microstructure (Holzer et al., 2010; Desbois et al., 2013; Houben et al., 2013; Keller et al., 2014). Another approach giving 3D information on the clay microstructure, albeit with a lower resolution, is X-ray tomography. Its advantage is the possibility of monitoring processes such as wetting in situ (Harjupatana et al., 2015) and of being combined with other methods like small-angle X-ray scattering (SAXS) or SEM (Suuronen et al., 2014; Kaufhold et al., 2016). X-ray diffraction and scattering methods are very common ways of studying the microstructure of swelling clays, providing information about the basal spacing – the periodic distances between stacked clay platelets. Most often, water-saturated, compacted samples are measured outside confined conditions (Muurinen, 2009; Holmboe et al., 2012; Cuisinier et al., 2014; Villar et al., 2016), but also in situ measurements in confined conditions have been made (Devineau et al., 2006; Warr & Berger, 2007; Perdrial & Warr, 2011). Mercury intrusion porosimetry (MIP) is the most commonly used method to determine pore-size distribution (Delage, 2007; Ye et al., 2009; Lubelli et al., 2013). It is also applied to swelling clays; however, only dry systems may be analysed, and the minimum pore aperture that may be resolved with MIP is ~5 nm (Marcial et al., 2006; Lloret & Villar, 2007).

As the montmorillonite platelets are the main structural components of bentonite, the porosity of bentonite is complex, and it varies depending on conditions such as the density or the level of hydration (Holmboe et al., 2012; Villar et al., 2012; Muurinen et al., 2013). The definition of the various groups of pores in bentonite is widely accepted. Interlamellar (interlayer, interlaminar) pores – slit-like pores with an aperture <2 nm – are formed between stacked montmorillonite platelets. All remaining pores (non-interlamellar) are sometimes divided into intra-aggregate and inter-aggregate porosities (Pusch et al., 1990; Villar, 2002). Consequently, many authors suggest the existence of waters of various properties in bentonite (Bourg et al., 2003; Bradbury & Baeyens, 2003; Fernández et al., 2004; Wersin et al., 2004; Kozaki et al., 2008); therefore, some microstructural information may be deduced by looking at the water that is present in pores. In line with this suggestion, anion exclusion from the pore space may be used to draw conclusions on the microstructure (Muurinen et al., 2004; Van Loon et al., 2007). Nuclear magnetic resonance (NMR) relaxation techniques are sometimes used to obtain pore-size distributions (Serland et al., 2007), and these techniques have been used to obtain information about water in swelling clays (Ohkubo et al., 2008; Dvinskikh & Furó, 2009; Matusewicz et al., 2013). Among the cited works, some use only purified clay (Ploehn & Liu, 2006; Kozaki et al., 2008; Pettersson et al., 2010; Segad et al., 2012; Matusewicz et al., 2013; Delavenne et al., 2015; Harjupatana et al., 2015) and a few use both purified and original clay (Montes-Hernandez, 2005; Devineau et al., 2006; Holmboe et al., 2012), while most focus on the original bentonites.

In this work, MX-80 bentonite is studied along with sodium and calcium montmorillonite obtained by purification of MX-80. It has to be kept in mind that the purification of MX-80 bentonite does not result in pure montmorillonite; rather, it is most accurate to use the term ‘fine fraction’ of MX-80 bentonite. However, as montmorillonite is the dominant clay mineral in the fine fraction of MX-80 bentonite, the authors use the simplified terms ‘Na-montmorillonite’ and ‘Ca-montmorillonite’ in this text. Each data point presented corresponds to one sample. The purified clay is intended to serve as a simplified system that might be used to better understand the influence of chemical conditions and density on the compacted, water-saturated clay microstructure. This research is focused on characterizing the changes in the clay microstructure induced by the purification process and on characterizing the pore structure of compacted, water-saturated clay.

**Materials**

Calcium and sodium montmorillonite were obtained by purification of MX-80 bentonite (CETCO, UK) used in experiments. The purification method included removal of coarse particles by centrifugation, of carbonates by acid treatment, of iron oxides by citrate buffer treatment and of organic matter by H2O2 treatment (Tributh & Lagaly, 1986). After drying, the purified clay was ground to the same grain size as MX-80 using a vibrating-disc mill. Cylindrical clay samples (1 cm high, 2 cm in diameter) were prepared at various densities by uniaxial compaction, water saturated and left to equilibrate with 0.1 M NaClO4 (Ca(ClO4)2 for Ca-montmorillonite) solution for a minimum of 1 year.

**Methodology**

Transmission electron microscopy and AFM have been used to investigate the changes induced in clay by the purification process. Transmission electron microscopy images give qualitative information about the microstructure of the samples. The samples were high-pressure frozen with a Leica EM Pact machine (Studer et al., 2001), embedded in EPI-ON™ epoxy resin, sliced with a microtome and imaged in an FEI Tecnai 12 microscope at a working voltage of 120 kV. Atomic force microscopy imaging was used to calculate the particle size and shape distributions of the clay particles. The clay suspensions were treated with ultrasound and deposited on a freshly cleaved mica surface. Imaging was then carried out using the scan assist mode of the Nanoscope MultiMode scanning probe microscope (Bruker Corporation, MA, USA). The image analysis was done using SPIP software (Image Metrology, Denmark), in which parameters of platelets were marked manually. The shapes collected in this way were used to calculate the equivalent diameter (defined as the diameter of a circle having an area equal to the area enclosed by the particle’s perimeter) and the aspect ratio (defined as the ratio of the particle length at the broadest point to its width in the perpendicular direction). Those simple geometrical parameters were chosen to determine whether the average size and shape of clay platelets changed during the purification process.

The porosity investigation was based on three different methods: (1) SAXS; (2) anion exclusion; and (3) proton-relaxation NMR. The SAXS patterns were obtained using an X-ray tube with a Cu anode (PANalytica, The Netherlands). The X-ray beam was collimated and monochromated to Cu-Kα radiation (wavelength 1.54 Å) using a Montel-multilayer mirror. The scattered intensity was collected using a Bruker Hi-Star area detector. The 0.3 mm section of compacted, water-saturated sample was confined with a thin polypropylene film to prevent drying during the measurement. The SAXS patterns were used to calculate the average distance between adjacent montmorillonite layers (d_{IL}), which, together with the specific surface area (SSA), were applied as in equation 1 (Matusewicz et al., 2017):

\[
V_{IL} = 0.5 \cdot d_{IL} \cdot SSA \cdot \frac{n - 1}{n}
\]
where \( n \) is the average number of layers per stack to estimate the volume of the interlamellar pores (\( V_{IL} \)). It was assumed that the edge surface area of montmorillonite was small enough to be disregarded in this calculation and the total SSA was used. The SSA value taken from the literature was \( 624 \pm 7 \text{ m}^2 \text{ g}^{-1} \) and was obtained using the ethylene glycol monoethyl ether (EGME) adsorption method (Kiviranta & Kumpulainen, 2011). The average height of the pore multiplied by the specific surface area of the slit-like pore forming montmorillonite particles gives the interlamellar pore volume. A correction for finite size of the stack needs to be included in the calculation. The ratio of interlamellar pore volume and the total sample volume is called ‘interlamellar porosity’ in the following text.

Anion exclusion was estimated by measuring the number of perchlorate ions that diffused into the sample during the sample equilibration. The anions were leached out of the clay, and their concentration in the leachate was measured using an ion-selective electrode (Nico2000 Ltd, UK). To calculate the anion-accessible porosity, it was assumed that the equilibration time was long enough to achieve the same perchlorate concentration in the accessible pore space as in the equilibrating solution and so no more anions would enter the sample. The anion-accessible volume is defined as the number of anions in the sample divided by the concentration of anions. The ratio of the anion-accessible volume to the total sample volume is referred to as the anion (in this work, perchlorate)-accessible porosity.

The NMR technique used was the \(^1\text{H} \) NMR Carr–Purcell–Meiboom–Gill (CPMG) echo-type experiment (Santry et al., 1988), which is a spin-locking pulse sequence with \( \tau = 22 \text{ ms} \). The measurements were carried out with a high-field Chemagnetics CMX Infinity 270 MHz NMR spectrometer. Relaxation curves were analysed using XPfit software (SoftScientific, Israel). The hydrogen nuclei in the sample during the experiment were brought to an excited state, with the recorded signal being the relaxation of the hydrogen nuclei of water molecules. As the relaxation time depends on the average distance of the water molecule from the pore wall, a cut-off time (Ohkubo et al., 2008; Matusewicz et al., 2013) was used to represent the total water population as a function of the pore size. The cut-off time was chosen to correspond to the relaxation time of water protons in a nanometre-sized pore. This means that the water population was confined to very fine pores, including interlamellar pores, as the relaxation time is shorter than the chosen cut-off time. The remaining water was found in the larger-pore structures.

More detailed descriptions of the sample preparation and the methods applied can be found in our previous work (Carlsson et al., 2012; Matusewicz et al., 2013).

Results and discussion

Transmission electron micrographs provide distinct images of MX-80 bentonite and the purified samples. In the case of MX-80, wavy, complex aggregates are seen (Fig. 1, right panel), whereas in purified samples, well-oriented layers of platelets dominate (Fig. 1, left and centre panels). The purification process, where clay is treated mostly in suspensions, destroys aggregates present in MX-80, and drying of the suspension towards the end of the process results in relatively organized deposition of fine-fraction platelets. No accessory minerals were visible in purified samples, whereas they were easily found in the original MX-80.

Atomic force micrographs show a micrometre-sized area with a nanometre-sized vertical resolution (Fig. 2). This allows measuring of the lateral size of individual clay platelets. If enough platelets are measured, statistical analysis can be performed. The sample preparation procedure applied did not break down all of the aggregates, so only delaminated, clearly separated platelets were taken into account.

Histograms of particle-size distribution show slightly different distributions for MX-80 and the purified material (Fig. 3). In both samples, most of the particles have a diameter in the range of 50–250 nm, yet within that range, the distributions vary. In the case of MX-80, the most populated region of the size distribution is between 100 and 150 nm, whereas in the case of the purified clay, it is between 50 and 100 nm. However, the purified distribution is bimodal, having a second peak with a value <200 nm. This range of sizes agrees well with the data previously reported in the literature (Ploehn & Liu, 2006; Delaverhe et al., 2015).

Few larger particles of equivalent diameter >600 nm and a log-normal distribution over the whole distribution were observed for MX-80, whereas purified clay particles >600 nm were not detected and the size distribution appears to be a superposition of two log-normal distributions. In another study reporting the platelet-size distribution in purified clay, sizes of >500 nm also were not observed (Delaverhe et al., 2015). However, it may be that the observation of very few particles >600 nm is linked to the larger population of the measured samples of MX-80 compared to the purified clay or to slight differences in the sample preparations. Nevertheless, the number of measured particles in the purified samples is large enough for the statistical analysis.

Some differences between fine-fraction platelets of MX-80 and purified clay are visible when analysing the particle aspect ratio. Note that in this research we focused on the aspect ratio between lateral platelet dimensions, whereas often in nanoscience the aspect ratio between the height and width of the platelets is of interest (Ploehn & Liu, 2006; Cao et al., 2010). Comparing the results presented in Fig. 4, a higher number of long, narrow particles (length to width ratio >3) is present in the purified clay. At the same time, the fraction of particles with an aspect ratio <1.5 is higher in purified clay samples. The presence of long and narrow particles could be a direct consequence of grinding at the end of the purification process, where dried clay was ground to grains of the same diameter as MX-80. Despite some variation in the fraction of the particles with the highest and lowest aspect ratios, the vast majority of platelets have an aspect ratio <2.5 for both samples.

In all of the approaches to characterizing the porosity of bentonite, the most important factor controlling the material microstructure is density (Sato et al., 1992; Marcial et al., 2006; Muurinen et al., 2013). It is expected that the degree of compaction affects the pore structure, but the following results show the finer influence of purification and chemical conditions observed with various methods.

Perchlorate-accessible porosity (Fig. 5) is clearly lower than the total porosity of the samples, which was expected due to the negative charge of montmorillonite platelets. The effect of the ionic strength of the solution, influencing the anion exclusion (Van Loon et al., 2007; Tournassat & Appelo, 2011), has to be considered here. The electrical double layer is thinner in solutions with a higher ionic strength; therefore, with a higher ionic strength, the exclusion effects decrease. The three analysed materials have different ionic strengths: Na-montmorillonite has the lowest strength (1:1 electrolyte, pure material), while MX-80 (1:1 electrolyte, accessory minerals dissolving in the pore space) and Ca-montmorillonite have higher strengths (pure material, but saturated with 2:1 electrolyte).
MX-80 and Ca-montmorillonite samples behaved in a similar way throughout the whole density range analysed, whereas in Na-montmorillonite, for the densities <1.2 g cm$^{-3}$, the anion-accessible porosity was clearly lower. Higher values for the anion-accessible volume in Ca-montmorillonites have been reported before (Muurinen & Carlsson, 2013). For all of the samples measured, perchlorate-accessible porosity decreased with increasing sample density not only in absolute values, but also as a fraction of the total porosity. Interlamellar porosity calculated from X-ray scattering patterns (Fig. 6) showed a similar trend for all of the samples, with Ca-montmorillonite having slightly higher values than MX-80 and Na-montmorillonite having slightly lower values than MX-80. The reason for this is that Ca-montmorillonite tends to form larger stacks than Na-montmorillonite (Schramm & Kwak, 1982). With increasing density, the fraction of interlamellar porosity calculated from X-ray scattering patterns (Fig. 6) showed a similar trend for all of the samples, with Ca-montmorillonite having slightly higher values than MX-80 and Na-montmorillonite having slightly lower values than MX-80. The reason for this is that Ca-montmorillonite tends to form larger stacks than Na-montmorillonite (Schramm & Kwak, 1982). With increasing density, the fraction of interlamellar porosity decreased with increasing sample density not only in absolute values, but also as a fraction of the total porosity.

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and with values reported in the literature (Muurinen et al., 2012). The estimated time of its protective functioning is at least tens of thousands of years. Therefore, a detailed understanding of bentonite is necessary. Changes to the exchangeable-cation composition in bentonite cause changes in properties – instead of studying the mixed-cation forms, it is common to study purified Na- or Ca-montmorillonite because homo-ionic materials provide results that are easier to interpret. As only few works have compared purified clays with ‘as-delivered’ bentonites, the results of this study will help us to understand the differences between those systems.

The main factor affecting the porosity of bentonites is the density caused by compaction. The effect of exchangeable cations is much smaller – differences in the pore structures of compacted samples are most visible at low densities, where the sodium form of montmorillonite has less porosity confined in nanometre-sized pores, including interlamellar spaces, than the calcium form of montmorillonite and MX-80 bentonite. At high compaction densities, therefore, under conditions similar to those that would be expected in the repository, the differences between the samples, as well as those caused by the compensating cation composition, seem to be less pronounced.

The obtained results also allow for the conclusion that the applied purification process clearly affected the clay microstructure. The montmorillonite platelet-size distribution shifted towards smaller particles for purified clays, with approximately 35% of particles having an equivalent diameter of between 40 and 120 nm. A change in shape was also noticed, with purified clay having a greater fraction of long, narrow particles than MX-80. The different size and shape distributions of the montmorillonite platelets in MX-80 clay and clay obtained by purification may be attributed to additional grinding at the end of purification (separately from chemical treatment and multiple washing cycles). Naturally, various purification procedures may affect variably the montmorillonite platelets in the purified material.

The long-term performance analysis of bentonite for safe disposal of spent nuclear fuel is based on modelling of both chemical and mechanical behaviour. Microstructural data are used, for example, in modelling diffusion and sorption by surface complexation modelling or as a basis for water transport and mechanical models. Calculating the diffusion of anions requires data on the porosity that is available for them, and diffusion is applied to the transport of corroding agents and radionuclides. Another possible application is extending surface complexation modelling from dilute to compacted systems and the estimation of the extension of diffuse double layers in the pore system. Swelling and swelling pressure formation are connected to the distribution of water between larger pores and the interlamellar space, and the data regarding microstructure also provide important guidance for mechanical modelling. The development of advanced structural models (dual porosity or more complicated assumptions) of compacted swelling clays may make use of porosity values and the differences between the methods. However, porosity should be measured with various methods in many different conditions (salinity, pH, temperature and pressure). This will provide experimental results for testing the developed models under various conditions.

Conclusions

Bentonite clay is applied as a buffer material between bedrock and copper canisters containing HLRW. The long-term performance analysis of bentonite for safe disposal of spent nuclear fuel is based on modelling of both chemical and mechanical behaviour. Microstructural data are used, for example, in modelling diffusion and sorption by surface complexation modelling or as a basis for water transport and mechanical models. Calculating the diffusion of anions requires data on the porosity that is available for them, and diffusion is applied to the transport of corroding agents and radionuclides. Another possible application is extending surface complexation modelling from dilute to compacted systems and the estimation of the extension of diffuse double layers in the pore system. Swelling and swelling pressure formation are connected to the distribution of water between larger pores and the interlamellar space, and the data regarding microstructure also provide important guidance for mechanical modelling. The development of advanced structural models (dual porosity or more complicated assumptions) of compacted swelling clays may make use of porosity values and the differences between the methods. However, porosity should be measured with various methods in many different conditions (salinity, pH, temperature and pressure). This will provide experimental results for testing the developed models under various conditions.

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References

Bourl I.C., Bourg A.C.M. & Sposito G. (2003) Modeling diffusion and adsorption in compacted bentonite: a critical review. Journal of Contaminant Hydrology, 61, 293–302.

Bradbury M.H. & Baeyens B. (2003) Porewater chemistry in compacted re-saturated MX-80 bentonite. Journal of Contaminant Hydrology, 61, 329–338.

Cadene A., Durand-Vidal S., Turq P. & Brendle J. (2005) Study of individual Na-montmorillonite particles size, morphology, and apparent charge. Journal of Colloid and Interface Science, 285, 719–730.

Cao T., Fasulo P.D. & Rodgers W.R. (2010) Investigation of the shear stress effect on montmorillonite platelet aspect ratio by atomic force microscopy. Applied Clay Science, 49, 21–28.

Carlsson T., Muurinen A., Matusewicz M. & Root A. (2012) Porewater in compacted water-saturated MX-80 bentonite. Pp. 397–402 in: MRS Proceedings (R.M. Carranza, G.S. Duffo & R.B. Rebak, editors). Cambridge University Press, New York, NY, USA.

Cuisinier O., Deneele D., Maisroui F., Abdallah A. & Conil N. (2014) Impact of high-pH fluid circulation on long term hydromechanical behaviour and microstructure of compacted clay from the laboratory of Meuse-Haute Marne (France). Applied Clay Science, 88-89, 1–9.

Delage P. (2007) Microstructure Features in the Behaviour of Engineered Barriers for Nuclear Waste Disposal. Springer, Berlin, Germany.

Delaverre L., Steudel A., Darbha G.K., Schäfer T., Schulmann R., Wöll C., Geckeis H., Emmerich K. (2015) Influence of mineralogical and morphological properties on the cation exchange behavior of diocathedial smectites. Colloids and Surfaces A: Physicochemical and Engineering Aspects, 481, 591–599.

Desbois G., Urai J.L., Pérez-Willard F., Radi Z., Ohkubo T., Kikuchi H. & Yamaguchi M. (2008) An approach of NMR relaxation and diffusion for understanding water in saturated compacted bentonite. Clays and Clay Minerals, 56, 276–277.

Montes-Hernandez G. (2005) Swelling-shrinkage measurements of bentonite using coupled environmental scanning electron microscopy and digital image analysis. Journal of Colloid and Interface Science, 284, 271–277.

Muurinen A. (2009) Studies on the Chemical Conditions and Microstructure in the Reference Bentonites of Alternative Buffer Materials Project (ABM) in Åspö. Posiva Oy, Eurajoki, Finland.

Muurinen A. & Carlsson T. (2013) Bentonite Port Structure Based on SAXS, Chloride Exclusion and NMR Studies. Working Report 2013–53. Posiva Oy, Eurajoki, Finland.

Muurinen A., Karland O. & Leikokinen J. (2004) Ion concentration caused by an external solution into the porewater of compacted bentonite. Physics and Chemistry of the Earth, Parts A/B/C, 29, 119–127.

Murphy M., Matusewicz M., Pirkkalainen K., Liljeström V., Suuronen J.–P., Root A., Muurinen A., Serinna R. & Olin M. (2013) Microstructural investigation of calcium montmorillonite. Clay Minerals, 48, 267–276.

Perdrial J.N. & Warr L.N. (2011) Hydration behavior of MX80 bentonite in a confined-volume system: implications for backfill design. Clays and Clay Minerals, 59, 640–653.

Pedrill J.N. & Warr L.N. (2011) Hydration behavior of MX80 bentonite in a confined-volume system: implications for backfill design. Clays and Clay Minerals, 59, 640–653.

Pettersson T., Thorinn E. & Claesson P. (2010) AFM Study of Montmorillonite. Svensk Kärnbränslehantering AB, Stockholm, Sweden.

Ploehn H.J. & Liu C. (2006) Quantitative analysis of montmorillonite platelet size by atomic force microscopy. Industrial & Engineering Chemistry Research, 45, 7025–7034.

Posiva (2012) YJH 2012, Nuclear Waste Management at Olkiluoto and Loviisa Power Plants: Review of Current Status and Future Plans for 2013–2015. Posiva Oy, Eurajoki, Finland.

Pusch R. (1999) Experience from preparation and investigation of clay microstructure. Engineering Geology, 54, 187–194.

Pusch R., Karland O. & Hökmark H. (1990) GMM – A General Microstructural Model for Qualitative and Quantitative Studies of Smeectic Clays. Technical Report 90–43. Svensk Kärnbränslehantering AB, Stockholm, Sweden.

Santry G.E., Henkelman R.M. & Bronskill M.J. (1988) Variation in measured transverse relaxation in tissue resulting from spin locking with the CPMG sequence. Journal of Magnetic Resonance (1969), 79, 28–44.

Sato H., Ashida T., Kohara Y., Uyi M. & Sasaki N. (1992) Effect of dry density on diffusion of some radionuclides in compacted sodium bentonite. Journal of Nuclear Science and Technology, 29, 873–882.

Savage D. (2012) Prospects for Coupled Modelling. STUK Radiation and Nuclear Safety Authority, Helsinki, Finland.
Schramm L.L. & Kwak J.C.T. (1982) Influence of exchangeable cation composition on the size and shape of montmorillonite particles in dilute suspension. Clays and Clay Minerals, 30, 40–48.

Segad M., Hanski S., Olsson U., Ruokolainen J., Åkesson T. & Jönsson B. (2012) Microstructural and swelling properties of Ca and Na montmorillonite: (in situ) observations with cryo-TEM and SAXS. The Journal of Physical Chemistry C, 116, 7596–7601.

SKB (2011) Long-Term Safety for the Final Repository for Spent Nuclear Fuel at Forsmark. Svensk Kärnbränslehantering AB, Stockholm, Sweden.

Sørland G.H., Djurhuus K., Widerøe H.C., Lien J.R. & Skauge A. (2007) Absolute pore size distribution from NMR. Diffusion Fundamentals, 5, 4.1–4.15.

Studer D., Graber W., Al-Amoudi A. & Eggi P. (2001) A new approach for cryofixation by high-pressure freezing. Journal of Microscopy, 203, 285–294.

Suuronen J.-P., Matusewicz M., Olin M. & Serimaa R. (2014) X-ray studies on the nano- and microscale anisotropy in compacted clays: comparison of bentonite and purified calcium montmorillonite. Applied Clay Science, 101, 401–408.

Tournassat C. & Appelo C.A.J. (2011) Modelling approaches for anion-exclusion in compacted Na-bentonite. Geochemical et Cosmochemica Acta, 75, 3698–3710.

Tributh H. & Lagaly G. (1986) Aufbereitung und Identifizierung von Boden- und Lagerstätten tonen. I. Aufbereitung der Proben im Labor. GIT – Fachzeitschrift für das Laboratorium, 30, 524–529.

Van Loon L.R., Glaus M.A. & Müller W. (2007) Anion exclusion effects in compacted bentonites: towards a better understanding of anion diffusion. Applied Geochemistry, 22, 2536–2552.

Villar M.V. (2002) Thermo-Hydro-Mechanical Characterisation of a Bentonite from Cabo de Gata. A Study Applied to the Use of Bentonite as Sealing Material in High Level Radioactive Waste Repositories. Publicación técnica, Empresa Nacional de Residuos Radiactivos, 4, 15–258.

Villar M.V., Gómez-Espina R., Campos R., Barrios I. & Gutiérrez L. (2012) Porosity changes due to hydration of compacted bentonite. Pp. 137–144 in: Unsaturated Soils: Research and Applications (C. Mancuso, C. Jommi & F. D’Onza, editors). Springer, Berlin, Germany.

Villar M.V., Gutiérrez-Rodrigo V., Iglesias R.J., Campos R. & Gutiérrez-Nebot L. (2016) Changes in the microstructure of compacted bentonite caused by heating and hydration. EJS Web of Conferences, 9, 18001.

Warr L. & Berger J. (2007) Hydration of bentonite in natural waters: application of ‘confined volume’ wet-cell X-ray diffractometry. Physics and Chemistry of the Earth, 32, 247–258.

Wersin P., Curti E. & Appelo C.A. (2004) Modelling bentonite–water interactions at high solid/liquid ratios: swelling and diffuse double layer effects. Applied Clay Science, 26, 249–257.

Ye W.M., Cui Y.J., Qian L.X. & Chen B. (2009) An experimental study of the water transfer through confined compacted GMZ bentonite. Engineering Geology, 108, 169–176.